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# **GENE CONSTRUCTION KIT<sup>®</sup> 3.0**

## **Tutorials & User Manual**

**from Textco BioSoftware, Inc.**

**June 2008, First Edition**

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# **Chapter 1**

## **Overview**

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### **Installing Gene Construction Kit®**

The initial **GCK** installation places all needed files in a single folder. The installed files include the most recent restriction enzyme database, tutorial, sample, and help files, and a vector database. All the files need to be installed on your hard disk from this CD.

#### **On the Macintosh**

Here is the complete step-by-step installation process:

1. Insert the **GCK** Installer CD and locate the GCK folder.
2. Drag this folder to your hard disk. Although not strictly necessary, it is best to drag it into the Applications folder.
3. With the CD still in the computer, start up the **GCK** application you just installed. Enter the personalization information. The first time you run the software GCK needs to verify itself by checking the CD. For subsequent launches, this is not necessary.
4. You are finished installing the software. You can now remove the CD from your computer. Please read the notes below and enjoy using Gene Construction Kit.

#### **On Windows**

1. Insert the CD into the drive of the computer on which you wish to run Gene Construction Kit.

2. From this installation CD, double-click on the application file titled **Setup.exe** to launch the installer program. Click “Yes” or “Next” to keep the defaults and install the application into the **Program Files** folder of your hard drive.

3. With the CD still in the computer, start up the GCK application by choosing Start > Program Files > Gene Construction Kit. Enter the personalization information. The first time you run the software GCK needs to verify itself by checking the CD. For subsequent launches, this is not necessary.

4. You are finished installing the software. You can now remove the CD from your computer. Please read the notes below and enjoy using Gene Construction Kit.

### **A few notes about the installation**

- On the Mac, it is important to drag the entire folder from the CD to ensure that the application will run properly. Dragging just the application from the CD to your hard disk will not work. If you have a previous version of **GCK** and have files that you would like to keep with the application, you can place them into the new **GCK** folder once you have dragged it from the CD.
- On both Mac and Windows, the first time you run the new **GCK**, you will be asked to insert the original CD (unless it is already in the CD drive). This is the only time you will need to do this unless you reformat your hard disk.

### **Updating Gene Construction Kit**

Updates are available for **GCK** from our web site as they are needed. You must have at least version 3.0.0 to update through the web. Check <<http://www.textco.com/updates.html>> to see if there is a newer version. To run the newer version of the application, download it from the web site, remove the previous version of the application, and put the new version into the **GCK** folder on your hard disk. After checking that it works, discard the older version of **GCK**. Up to date instructions are always available at the web site.

On the **GCK** CD, we have included a demo version of Gene Inspector®. This is Textco's DNA and protein analysis program that complements **GCK**. You can install the demo using the installer provided in the Gene Inspector Demo folder.

Please call us if you have any questions or problems.

We hope you enjoy your new software.

## **Overview**

The Gene Construction Kit® has been designed to facilitate handling DNA in an innovative and intuitive way. Over a decade of development by a team consisting of molecular biologists and programmers has lead to an interface and a set of functions in Gene Construction Kit (**GCK**) that will significantly enhance laboratory productivity and minimize experimental design errors.

The backbone of the program is the **construct**, which represents a DNA sequence and all associated sites, regions of interest, segments, comments, and history. The construct can be manipulated and analyzed using four types of windows. Each **GCK** window visible on the screen corresponds to a file on the disk. The name of the window is the name of the file. There are four kinds of **GCK** windows: Construct, List, Gel, and Illustration. These are described briefly below and in their own chapters in this manual.

All manipulations of the actual DNA are carried out on the construct within the **Construct Window**. Restriction enzyme and other sites can be marked, regions of interest can be defined, silent mutations generated, PCR products generated, and the appearance of the construct and its annotations can be edited within the Construct Window. The construct can be displayed as linear or circular and either graphically or as a formatted DNA and protein sequence listing. Construct Window details are discussed in Chapter 3, page 3-1.

The **List Window** is used to create, maintain and edit lists of sequences. The list may contain restriction enzyme recognition sequences, protein binding sites, linkers, promoters, etc. Cut sites can be defined for sequences in the

list and comments can be associated with any item in the list. When the list items are used to mark sites in the construct, the comments remain associated with each site that is marked on the construct. More information about the list window is given in Chapter 4, page 4-1.

The **Gel Window** is used to simulate gel electrophoresis patterns. Any number of digests can be displayed in the Gel Window. Partial digests can be viewed and lanes can be cut and pasted within and between gel windows. In addition to a graphical view of the gel, the digests can be viewed as tables of fragment size information. The Gel Window is discussed in Chapter 5, page 5-1.

The **Illustration Window** is used to create documents for presentation or publication, and can also be used to track complex construction projects. Graphics and text from Construct, List, and Gel Windows can be pasted into an Illustration Window and legends can be automatically generated. Graphics and text generated in other programs can be copied and then pasted into a **GCK** Illustration Window. Once pasted into an Illustration Window, constructs are separate from their original file, but can still be edited using all the tools normally available in the Construct Window. The Illustration Window also contains a comprehensive set of graphical drawing tools for enhancing illustrations. For more information on the Illustration Window see Chapter 6, page 6-1.

## **Running The Program**

**GCK** is a comprehensive program with many capabilities. The first time you run it, the possibilities might seem so extensive that you are not sure of what to do first. To provide you with an overview of the capabilities and to give you a feel for how the program actually works, you should go through the various tutorials in Chapter 2. **If you read nothing else in this manual, you should read and carry out the tutorials.** These tutorials are designed to help orient you and will greatly facilitate your understanding of the program and its features, while pointing out features you may not think to look for. The time you spend in the tutorials will be repaid many times over in the future as you use **GCK**.

**GCK** has a straightforward and intuitive interface with extensive “undo” capabilities. It has what is called *revealed complexity* - you can see as much detail as you choose. You should not hesitate to try things for yourself. Select menu options, tool palettes, etc. to see what they do. The Format Menu, in particular, is available from many places within the program and will be useful in ways you might not expect. Besides the tutorials, one of the best ways to learn the capabilities of the Gene Construction Kit is to try new menu options or tools for yourself.



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# **Chapter 2**

## **Tutorials**

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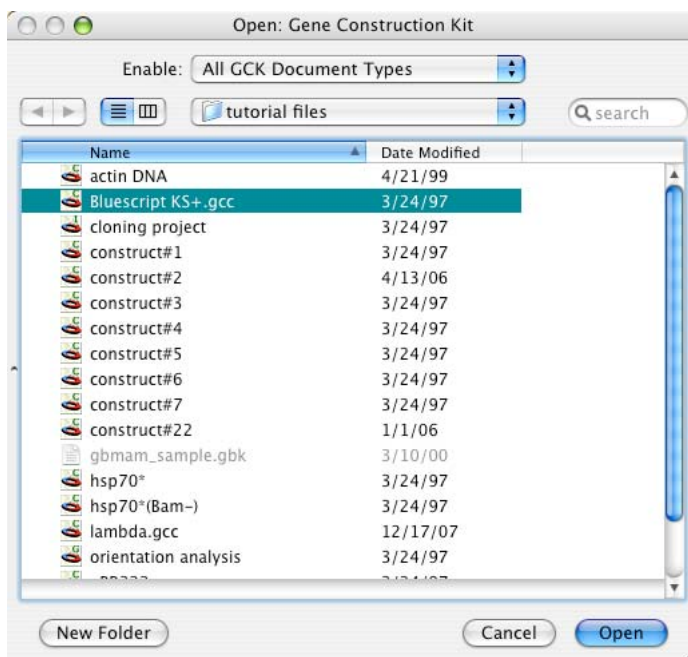
This chapter contains a number of tutorials that introduce you to the Gene Construction Kit®. You should do as many of the tutorials as you can because they provide an overview of how the program works. **Doing the tutorials now will save you many hours in the future.** The Gene Construction Kit has a number of unique features you might not have seen in any other application – the tutorials provide a way for you to learn about these unique capabilities.

In many locations in this chapter and throughout the manual you will be asked to select items in menus. To make your choices as clear as possible, all menu selections are indicated as hierarchical choices using this Menu font, such as Edit > Select All. This particular case means to locate the Edit menu and then choose Select All under the Edit menu.

## TUTORIAL 1: WORKING WITH CONSTRUCTS

Perhaps the most important window in GCK is the **Construct Window**. This is the place where you will do the majority of your cloning manipulations and can precisely define how your construct will look. In this tutorial you will learn how to modify the appearance of constructs. Tutorial 2: “Marking Sites”, page 2-8 shows how to mark restriction sites and other features in your construct. Tutorial 7: “Chronography – Tracking Cloning History”, page 2-30, shows how GCK can automatically keep track of construct history for you. Tutorial 4: “Viewing the Construct as a Sequence”, page 2-15 discusses the sequence view of a construct.

1. Start GCK.
2. Open the file called **Bluescript KS+** in the **tutorial files** folder by choosing File > Open.... This will bring up Figure 2.1. This folder should be in the GCK

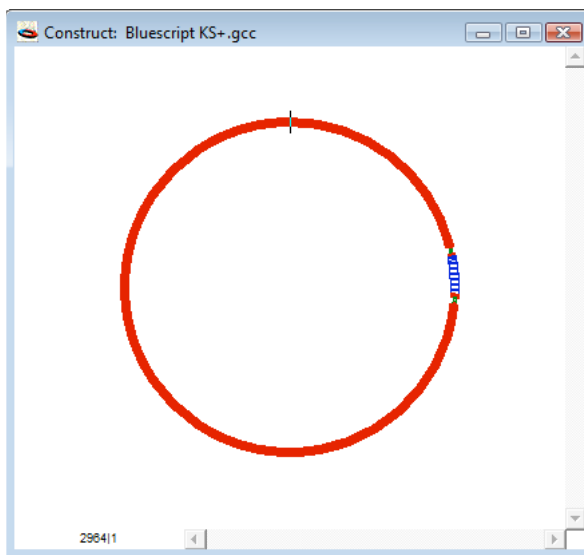


**Figure 2.1: Opening a file**

folder that was created when the program was installed. The buttons on the



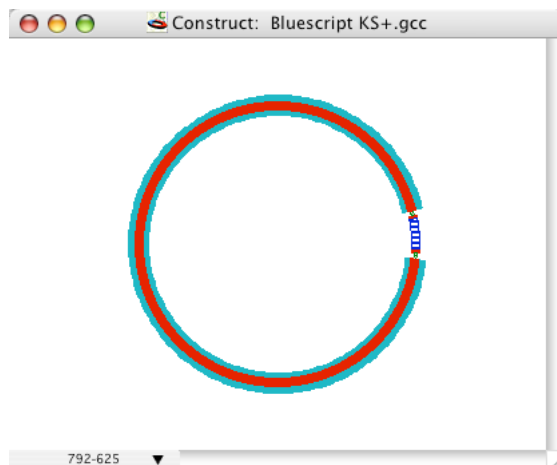
bottom let you choose to have listed different types of files in the list on the left. Since we are opening a construct file, leave the radio button set on **Construct**. When you open other kinds of GCK files, the buttons will be useful to you. Select the **Bluescript KS+** file and press the **Open** button. You will see Figure 2.2.



**Figure 2.2: Bluescript KS+**

3. This construct has a number of **segments**, or sections, of DNA indicated in different colors and patterns. The title of the file is indicated in the title bar at the top of the window. In the lower left corner of the Construct window is an indicator of the location of the current **insertion point**. The insertion point is where new sequence would be placed into the current construct by typing or pasting, between nucleotides 625 and 626 in this window (you might have a different location in your file). Clicking with the mouse anyplace along the DNA will place the insertion point at the closest boundary between two segments.

4. Double-click on the red segment of DNA to select it. This is similar to selections made in standard text editing programs, where double-clicking selects a word. This will highlight the red segment as shown in Figure 2.3, page 2-4. Note how the indicator in the bottom left corner now shows the



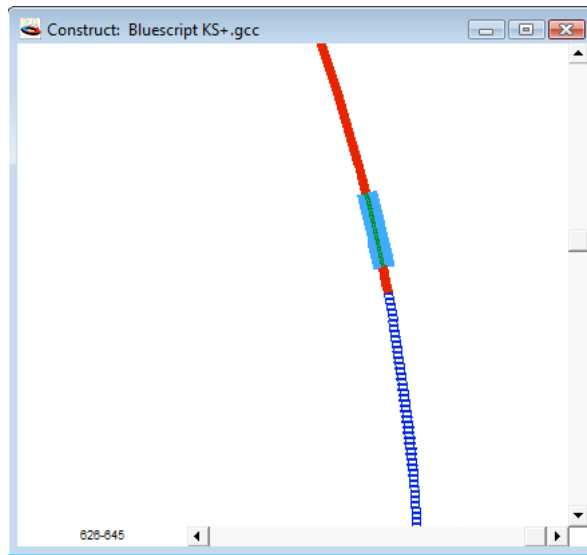
**Figure 2.3: Selected Segment**

range of nucleotides selected. The range is always shown in the clockwise orientation (or from left to right for linear constructs). You should see 792-625 selected. The origin (top of construct) is defined as position#1. Clicking on this popup menu will also allow you to show the position of the cursor as you move it over the DNA, or the size of the selected DNA segment.

5. Once an object (like a DNA segment) is selected, you can modify it. Choose **Format > Lines >** and choose the second line down to change the width of the selected DNA segment to 2 pixels. In the **Lines** menu, you can set the width using the line thicknesses shown in the menu, or you can choose **Format > Lines > Pick a Width...** to type in your own thickness. Using the **Format** menu, you can change the appearance of the selected object. For now, do not change anything else but feel free to explore the **Format** menu on your own when you finish this tutorial.

6. This vector has T3 and T7 promoters and a polycloning site that we want to indicate. The promoters are shown in green and the polycloning site is a striped blue. Let's add directionality to the promoters. Before you can add arrowheads to any segment of DNA it must first be selected. The promoter segments are rather small and might be difficult to select at this magnification. So click the mouse on one of the green segments (between the blue and red segments) to place the insertion point near what we are interested in and then choose **Construct > Magnification > Zoom In** to enlarge the view of the con-

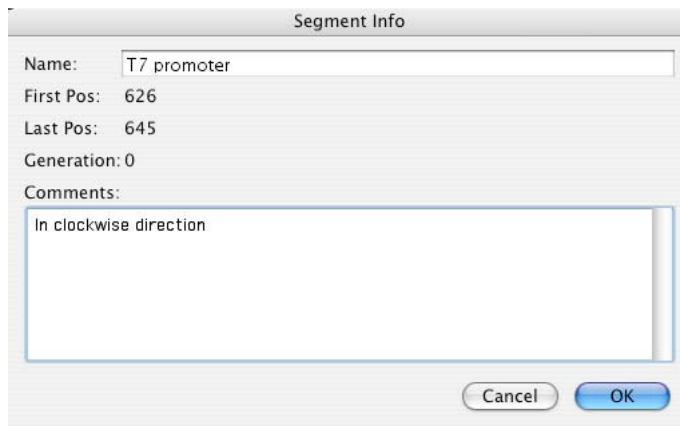
struct. Zoom In again and then select the top green segment by double-clicking. You will see Figure 2.4, page 2-5. Nucleotides 626-645 will be



**Figure 2.4: Magnified View**

selected.

7. Choose Construct > Get Info... to see the name of the segment and any comments associated with that segment. This is the T7 promoter as shown in Figure 2.5. Every segment of DNA can have a name and comments associated with it.

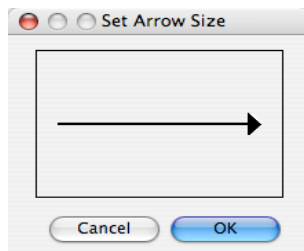


**Figure 2.5: Segment Get Info Box**

ated with it. These comments can be used to store important information about the construct and can be searched using the **Search Files** capability of **GCK** (Tutorial 8: “Finding Comments and File Searching”, page 2-38). We will talk more about this dialog box later (Tutorial 7: “Chronography – Tracking Cloning History”, page 2-30). For now just press the **Cancel** button – you don’t want to make any changes.

8. You should still have the T7 promoter selected as shown in Figure 2.4, page 2-5. If you do not, then select it again by double-clicking on the segment. First, let’s change the line thickness by choosing **Format > Lines >** and choose the third line down to change the width of the selected DNA segment to 3 pixels. Next add an arrowhead pointing in the clockwise direction by choosing **Format > Lines >** and choosing the rightward pointing arrow in the second section of the **Lines** menu.

9. The last step is to adjust the arrowhead size. This is done by choosing **Format > Lines > Size Arrowhead...** You will see Figure 2.6. Click and drag the



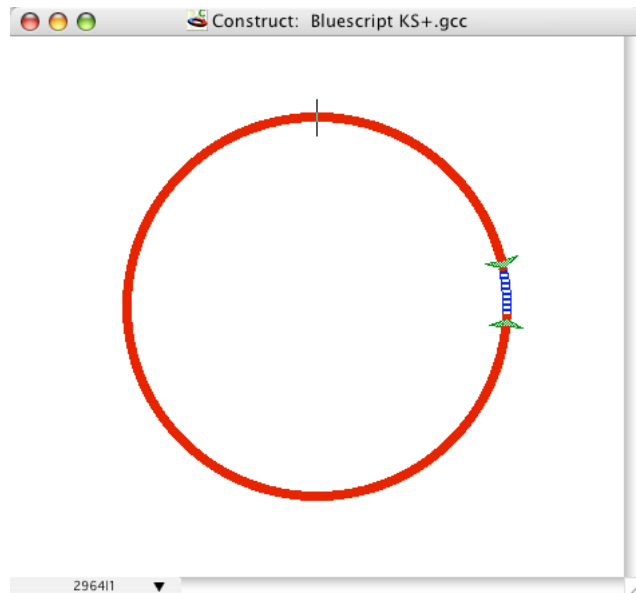
**Figure 2.6: Size Arrowhead**

mouse in the dialog box to resize the arrowhead to be approximately what is shown in this figure. Click **OK**.

10. We now need to adjust the T3 promoter segment to point in the counter-clockwise direction. This can be done by following the same steps we just did for the T7 promoter. Briefly, do the following: *Select* the lower green segment (the T3 promoter) by double-clicking on it. If you cannot see this segment in the window, use the scroll-bars to bring it into view. Choose **Format > Lines >** and select the third line down to change thickness, choose **Format > Lines >** and select the *leftward* pointing arrow. This will add an arrowhead that is the same size as the last arrowhead created so you should have symmetrical

arrowheads.

11. Finally, we want to see the whole construct again, so choose Construct > Magnification > Fit to Window. This will produce Figure 2.7, page 2-7. We have



**Figure 2.7: Modified Bluescript KS+**

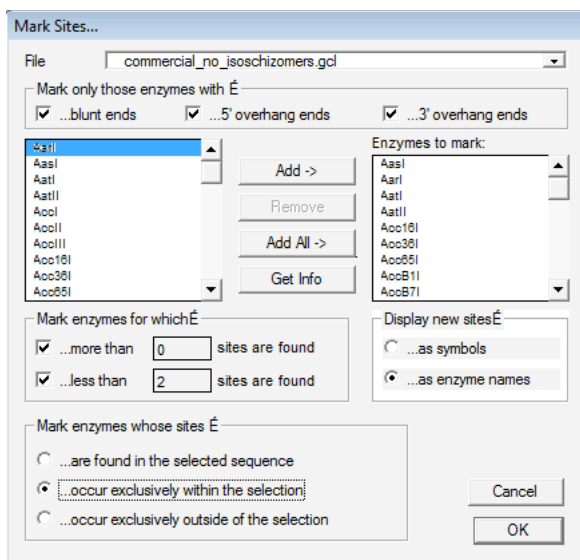
now formatted the segments of this construct to meet our needs.

12. You now need to save the construct under a new name. You might first want to create a folder on your desktop to hold the different tutorial files you will be creating in this and in subsequent tutorials. Having a single folder will make it easier to find the tutorial files you create as you need them in later tutorials. Choose File > Save As..., name the file **construct#1**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later on. Make sure you know where you are saving this file. You will need it again later.

You can quit now, or continue on to Tutorial 2: “Marking Sites”, page 2-8. If you plan on continuing, you can leave the file you just created, **construct#1**, open. If not, close this file.

## TUTORIAL 2: MARKING SITES

1. Start GCK and open the file **construct#1**, which you created in the first tutorial. If you do not have this file, you can start with **construct#1** from the **tutorial files** folder provided.
2. The next thing to do is to mark restriction enzyme sites that are located only in the polycloning region and nowhere else. To begin, double-click on the polycloning segment which has a blue lined pattern and is located at about 3 o'clock. You should see the segment become selected.
3. Choose Construct > Features > Mark Sites.... You will see Figure 2.8. This dialog



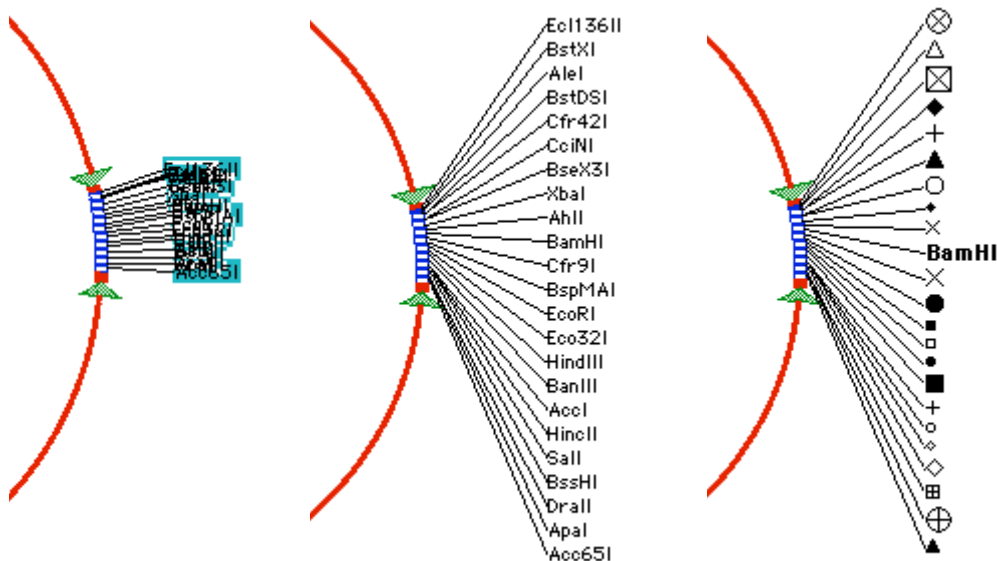
**Figure 2.8: Mark Restriction Sites**

is discussed more fully in “Marking Sites”, page 3-10. For now, make sure that the *commercial (no isoschiz)* list shows in the popup menu at the top of the dialog. A list of enzymes will appear in the left scrollable area. Press the **Add All ->** button to add all the enzymes in this list to the right side list of *Enzymes to mark*.

4. In the middle section on the left, set the numbers to match that in the figure so you will be specifying enzymes that cut more than 0 times but less than 2 times – i.e. unique cutters.

5. Finally, choose the middle radio button on the bottom left of the dialog box to specify that you want to see those enzymes which cut exclusively within the selected segment of DNA. Now press the **OK** button to mark the sites.

6. You will see the construct labeled as shown in Figure 2.9A. Note that all



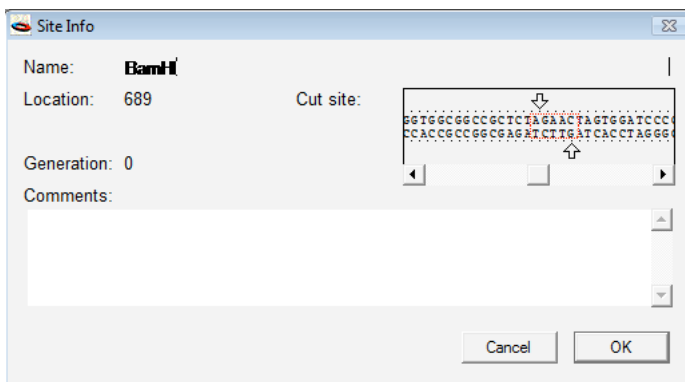
**Figure 2.9: Marked Sites**

the sites are crowded together. While all the newly marked sites are still selected choose **Format > Site Markers > Automatic Arrangement**. This will rearrange the site markers so that they do not overlap resulting in something that looks like Figure 2.9B.

7. As was the case for a selected segment of DNA, when a site marker is selected, you have access to items in the **Format** menu to change the display of the site marker. Now shift-click on the **BamHI** site to deselect it (all the other sites remain selected). Select the **Format > Site Markers > Show Site As Symbol** to change all of the sites to symbols except the **BamHI** site. Click on the **BamHI** marker and use the **Format > Style** menu to make the text bold; use the **Format > Size** menu to make the text 10 point. You should end up with some-

thing that looks like Figure 2.9C.

8. While the *Bam*HI site marker is still selected, choose Construct > Get Info... or press command-I (Mac) or ctrl-I (Windows). This will provide you with the information shown in Figure 2.10. If you wanted to, you could actually change the



**Figure 2.10: Site Marker Get Info Dialog**

name of the enzyme (not a good idea) or enter comments in the comments text box. The horizontally scrollable area in the lower part of the dialog box indicates the actual sequence at that location. The dotted red rectangle indicates the **recognition sequence** for the restriction enzyme and the two arrows indicate the actual **cut sites** for the enzyme. You can change the cut site by clicking on an arrow and dragging it. Press **Cancel** to close this dialog without saving any changes.

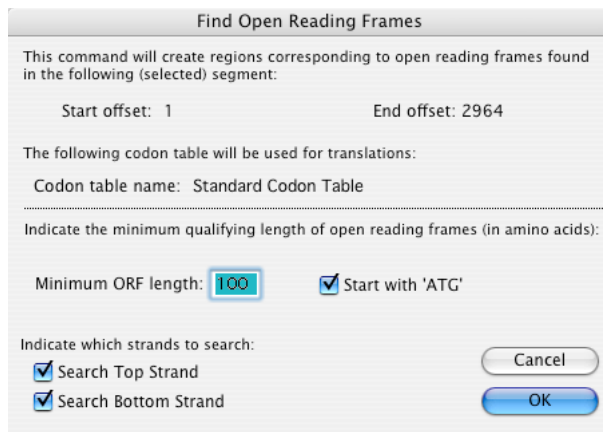
9. Choose File > Save As..., name the file **construct#2**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later. You will need it again in the next tutorial.

You can quit now, or continue on to Tutorial 3: “Marking Open Reading Frames”, page 2-11. If you plan on continuing, you can leave the file you just created, **construct#2**, open. If not, close this file.



## TUTORIAL 3: MARKING OPEN READING FRAMES

1. Start GCK and open the construct file you created in the last tutorial, **construct#2**. If you do not have this file, you can start with **construct#2** from the **tutorial files** folder provided.
2. You will finish annotating this construct in this tutorial and learn how to define **open reading frames (ORFs)**. First, choose **Construct > Display > Show Construct Title** which will place the title of the construct and the length of the construct into the center of the window.
3. Now choose **Construct > Features > Find Open Reading Frames...** You will see the dialog shown in Figure 2.11. This dialog gives you complete information about

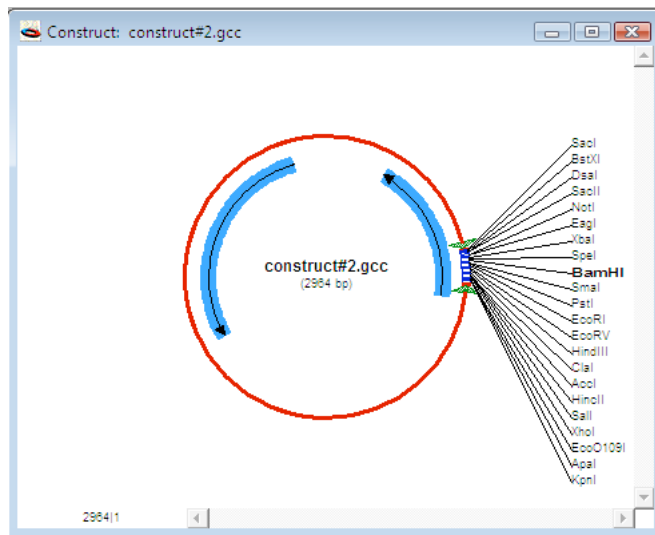


**Figure 2.11: Finding Open Reading Frames**

how the program will look for ORFs. The minimum ORF length text field tells the program to only display those ORFs containing 100 or more amino acids. As explained in “Finding Open Reading Frames”, page 3-36, you can select the codon table to be used and specify whether to search one or both strands of DNA. If you are looking for ORFs in exons you might want to turn off the **Start at ‘ATG’** check box - the program will then find any open reading frame (not just those starting with ATG). After you’ve adjusted the dialog to look like the one shown here, press the **OK** button to find the ORFs.

4. After running the ORF search you will see two new objects in the construct window: arrows representing the two different ORFs found by this

search, as shown in Figure 2.12. Both ORFs are highlighted. These ORFs

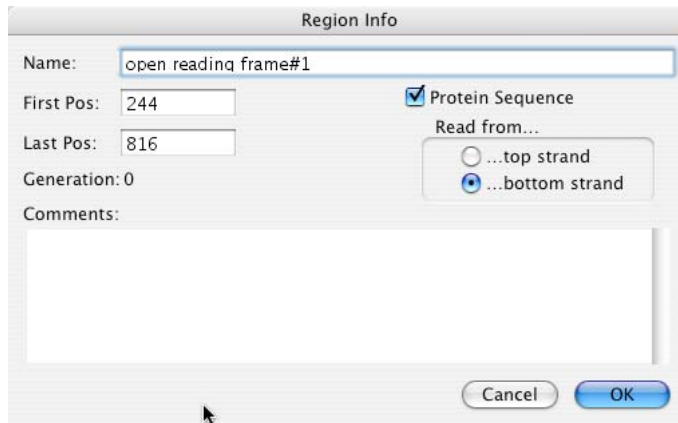


**Figure 2.12: ORFs are Marked**

represent **regions of interest**, or **regions**. In addition to site markers and DNA segments, regions are a third kind of object that can appear in the construct window. Regions can be formatted using the **Format > Regions >** submenu.

5. Click somewhere in the window to deselect the regions just created. Now click once on the region on the right side that overlaps the polycloning site. Now choose **Construct > Get Info...** to bring up Figure 2.13, page 2-13. This dialog contains information about the selected region. You can change the direction of the region by clicking the **Top Strand** or **Bottom Strand** radio buttons. Changing the numbers in the **First Nucleotide** or **Last Nucleotide** box will change the length of the region. The **Protein Sequence** check box determines if the region is to be translated or not when viewed as an actual sequence (see Tutorial 4: “Viewing the Construct as a Sequence”, page 2-15 for more information).

6. This region is the beta-galactosidase region, so type “beta-galactosidase” into the **Region Name** text box. You can also enter comments into the **Region Comments** text box. Both the name and the comments can be searched by **GCK** later (Tutorial 8: “Finding Comments and File Searching”, page 2-38). In this case it might be reasonable to enter some comments about this vector



**Figure 2.13: Region Get Info Dialog**

and how the polycloning site disrupts the  $\beta$ -gal coding region. Click **OK** to close this dialog box.

7. With the region still selected, choose Format > Regions > Show Region Names. This will display the region name in the construct window. You can select this name and alter the appearance of the text by using appropriate items under the Format menu. The region name text can be dragged around the construct window. When the region name text is selected, it is highlighted along with the region itself, making it clear which region is connected with the text.

8. Now select the left region by clicking on it once with the mouse. Choose Construct > Get Info... Change the name of this region to "ampicillin resistance" and put in comments indicating that the gene product is "beta-lactamase". Click **OK**.

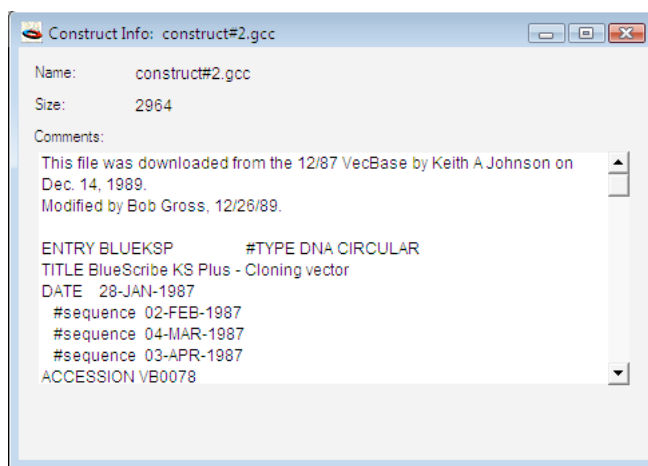
9. Note that when you have a region selected, you have access to many items in the Format menu. This will allow you to change the appearance of the arrow in the construct and even change directions by using the Format > Lines... menu. This would be equivalent to changing the protein direction in Figure 2.13.

10. Choose File > Save As..., name the file **construct#3**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later. You will need it again in the next tutorial.

You can quit now, or continue on to Tutorial 4: “Viewing the Construct as a Sequence”, page 2-15. If you plan on continuing, you can leave the file you just created, **construct#3**, open. If not, close this file.

## TUTORIAL 4: VIEWING THE CONSTRUCT AS A SEQUENCE

1. Up to now, we have been working with a graphical view of the construct, but underneath all of this is a real DNA sequence. Start **GCK** and open the file **construct#3** that you saved in the previous tutorial. If you do not have this file, you can start with **construct#3** from the tutorials folder provided.
2. Choose Construct > General Info... You will see something like Figure 2.14.

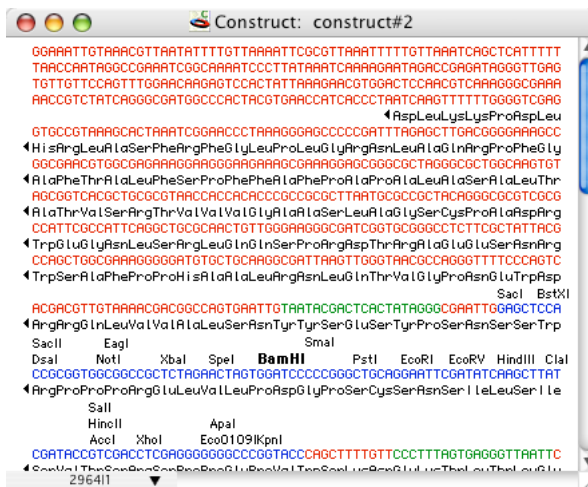


**Figure 2.14: General Info Dialog**

The General Info is stored with the construct as a whole and is not attached to any specific segment or region of the DNA. This means that it will not be lost if you reformat the construct or cut and paste segments. It is a useful place to store information such as storage location for the DNA and other information as shown in **construct#3**. In combination with the File Searching ability (Tutorial 8: "Finding Comments and File Searching", page 2-38), the comments can be used to create a database of information on all your clones.

3. Use the scroll bar until you see the information stating that the f1 DNA, which contains an origin of replication, is in the segment from 3-458. What we would like to do now is indicate the location of this particular segment of DNA, but how do we define the locations in our graphical construct which

does not have any markers at those locations? The answer is to view the construct as a sequence rather than graphically. Close the General Info window and then choose Construct > Display > Display Sequence. This will bring up Figure 2.15. If your display does not look like this figure, then scroll the



**Figure 2.15: Viewing the Construct as Sequence**

sequence all the way to the top of the window. Notice the correspondence between the graphical view and the sequence view. Colors are maintained, sites are marked, and regions are indicated. However, since we specified that the regions we marked are protein regions (Figure 2.13, page 2-13), the translated peptide is displayed below the actual DNA sequence. The small triangles adjacent to the amino acid line numbers indicate the direction of translation.

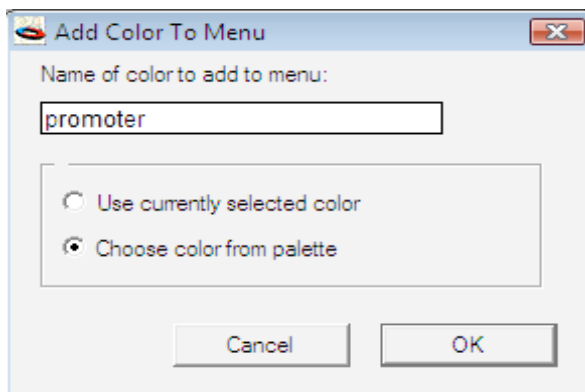
4. Before we can mark the f1 origin of replication, we need to be able to identify the nucleotides to be marked. Let's add position indicators by choosing Construct > Display > Show Positions. Remember how double-clicking on a segment in the graphical view of the construct selected that particular segment? This works the same way in the sequence view. So double-clicking anywhere in the DNA sequence will select that entire segment. Try this by double-clicking anywhere in the sequence to get a feel for how segments are defined. You will notice that segment boundaries can be site markers or changes in color or text characteristics.

5. With the insertion point (mouse cursor) anywhere in the DNA sequence, choose **Edit > Select All** (command-A, Mac or ctrl-A, Windows). This will select all of the DNA sequence. Now choose **Format > Grouping > Group by Tens**. This will arrange the sequence to be displayed in groups of tens. Note that there is an option in the **Format > Grouping** menu to define your own group size. This can be useful, for example, if you are working with something like an octamer repeat. You can define a groups size of 8 to display this repeat easily.

6. Grouping by ten is good for viewing the sequence and identifying specific locations, but does not look right for viewing those parts of the sequence that correspond to translated regions of the DNA. We need to group the translated regions into groups of three corresponding to the codons. This can be done in two ways. You could select the DNA corresponding to the peptide by clicking and dragging the mouse over the correct region of DNA. There is an easier way to do this, however, by just double-clicking on the *amino acid sequence*. This is a built-in feature of **GCK** to make it easier to select DNA segments that correspond to regions. Double-clicking on any region will select the corresponding segment of DNA. So double-click on the amino acid sequence corresponding to the  $\beta$ -galactosidase gene (it ranges from 244-846). Now choose **Format > Grouping > Group by Threes**.

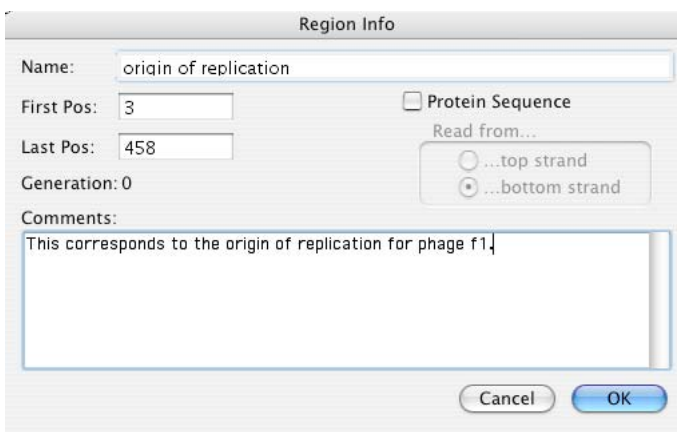
7. We now need to indicate our origin of replication. Identifying origins of replication is something that you might like to do in all of your constructs and you would like an easy visual method of identifying those regions from amongst the many that you might end up marking. You can do this by defining a specific color to represent an origin of replication. Choose **Format > Color > Add a Color...**. You will see Figure 2.16. Type in "origin of replication" for the name of the color and then press **OK**. Choose a color from your computer's standard Color Picker. Now you have defined a new color called *origin of replication*. This color should now appear in the **Format > Color** menu. The ability to define your own color and to name it provides an easy way for you to indicate features of interest such as promoters, enhancers, and other sites on DNA. If all your constructs are defined using your own "standard" colors, it will be easy to recognize specific functions of constructs just by observing the colors.

8. Let's take advantage of the new color we just created to identify the origin of replication for our construct. Use the mouse to select nucleotides 3



**Figure 2.16: Add a Color Dialog**

through 458. You can confirm that you have the correct nucleotides selected by looking in the lower left corner of the window. Choose Construct > Features > Make Region.... You will see the dialog shown in Figure 2.17. Make sure you



**Figure 2.17: Creating a New Region**

type in the correct name and put any comments you have into the comments area. This is NOT a protein sequence so do not check the **Protein sequence** checkbox. Confirm that the first and last nucleotides are 3 and 458, respectively. Press **OK**.

9. The newly created region will be selected<sup>a</sup> so you can now alter its

---

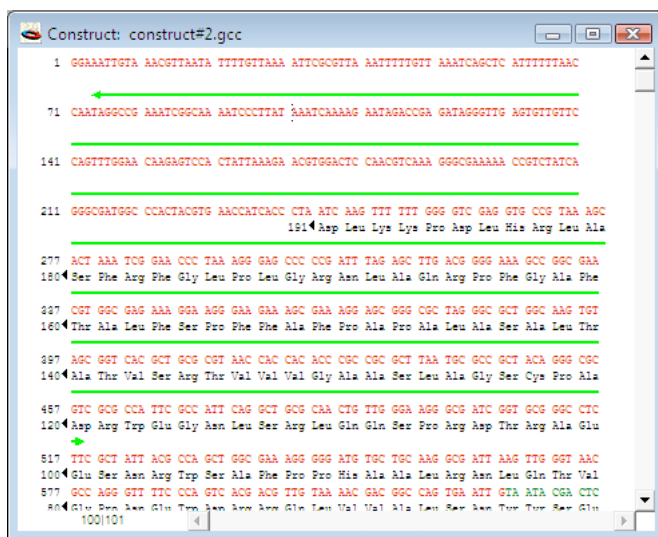
a. If it is not selected, then just click on it once to select it.



appearance. Choose Format > Lines and select the line having an arrowhead at both ends. This will change the region to have two arrowheads.

10. With the region still selected, choose Format > Color > origin of replication to change the color of the region to indicate its function.

11. You can also use the Format menu now to change the line thickness and arrowheads if you so desire. When you are done you should see something like Figure 2.18, page 2-19. You can now easily recognize the origin of repli-



**Figure 2.18: Origin of Replication Indicated**

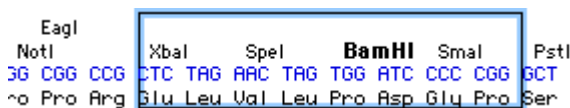
cation by color and can use that region to select the corresponding DNA by double-clicking on the region.

12. Choose File > Save As..., name the file **construct#4**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later. You will need this file in the next tutorial. That tutorial will explain how to enhance the appearance of your sequence in the construct window.

You can quit now, or continue on to Tutorial 5: “Modifying the Construct Appearance”, page 2-20. If you plan on continuing, you can leave the file you just created, **construct#4**, open. If not, close this file.

## TUTORIAL 5: MODIFYING THE CONSTRUCT APPEARANCE

1. Start **GCK** and open the file **construct#4** that you saved in the previous tutorial. If you do not have this file, you can start with **construct#4** from the tutorials folder we provided.
2. It is possible to modify the sequence appearance using many of the items in the **Format** menu. Select the **AT-rich** segment from nucleotides 10-33 by dragging with the mouse. Now Choose **Format > Font > Times** and notice that even though **Times** is not a monospaced font, **GCK** still spaces the characters so that each character occupies the same width along the line. Now choose **Format > Style > Bold**. The bold characters are wider than the non-bold characters so this part of the sequence becomes more “spread out”. You can adjust for that by choosing **Format > Style > Condense**, which condenses the spacing between characters. You can change colors, fonts, and styles for any segment of the DNA to help indicate certain features. Feel free to try some of these on this segment now.
3. Another method can be used to illustrate certain features. This is through the use of **Frames**. To see how frames work, select nucleotides 676-699 (they are in the polycloning region). Choose **Construct > Features > Make Frame**. This will create a frame around the segment of DNA you selected and will include the region (amino acid sequence) and site markers. It should look similar Figure 2.19. The frame is selected (highlighted) and encloses all infor-



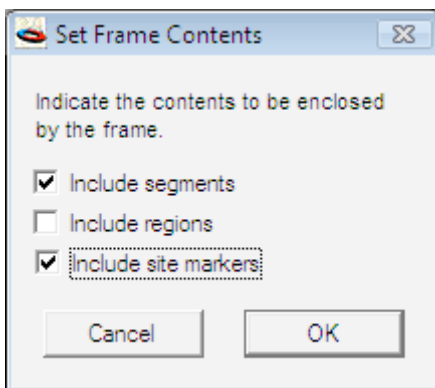
**Figure 2.19: A Framed Sequence**

mation related to that particular segment of DNA.

4. The appearance of the frame can be altered by using items in the **Format** menu, as usual. Note that a frame must be selected<sup>b</sup> (it must appear highlighted in the window) in order for items in the **Format** menu to have any effect on the frame itself. With the frame selected, choose **Format > Lines >** and choose

the second line thickness in the menu (it is two pixels wide). Choose `Format > Color > Yellow`, and then choose a fill pattern using `Format > Fill`. You can try different patterns and colors to see how they look. (If you plan to print the sequence, remember to use a light color for the fill pattern so that it does not obscure the text.)

5. Note that the frame encompasses the site markers, the DNA, and the region. You can limit what is enclosed in the frame by choosing `Construct > Features > Set Frame Contents...` as shown in Figure 2.20. This dialog allows you to



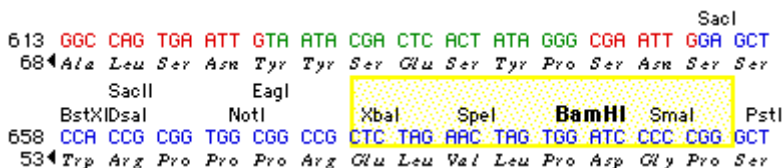
**Figure 2.20: Set Frame Contents**

define which items will be enclosed in the frame. Set your dialog box to look like the one shown here and then press **OK**.

6. It is somewhat difficult to distinguish between the DNA sequence and the protein sequence, so let's change the appearance of the protein sequence. Click once to select the protein sequence and then choose `Format > Font > Times`, then `Format > Style > Italic`. Your display should now resemble that shown in Figure 2.21, page 2-22. The ability to alter the display of the sequence in so many ways provides a great deal of flexibility in pointing out specific features of any sequence with which you are working.

7. Choose `Construct > Display > Display Graphics`. Note that the new region you

- b. A frame can be selected by option-clicking (on the Mac) on the frame or by choosing `Construct > Select Frame` and then clicking on the frame you want to select. For those of you who are "quick-fingered" you can also select a frame by triple-clicking on the DNA contained within a frame.



**Figure 2.21: A Modified Framed Sequence**

defined as the *origin of replication* also appears in the graphical display and is selected as it was in the sequence display. There is a direct correspondence between the two views of the construct. Changes you make in the graphical view appear in the sequence view and changes you make in the sequence view appear in the graphical view. The graphical view just represents an easier way to visualize your construct. For cloning fragments (Tutorial 6: “Cloning a DNA Segment and Silent Mutations”, page 2-23), this is essential to maintaining an easy to use interface.

8. Click on the *Bam*HI site marker name and then select all site markers by pressing command-A (Mac)/ctrl-A (Windows) or by choosing Edit > Select All. We want to change the appearance of all the site markers *except* for the *Bam*HI site so we need to deselect the *Bam*HI site while leaving the others selected. This is done in the standard way of holding down the shift key and clicking on the *Bam*HI site (this process is called shift-clicking)<sup>c</sup>. Make sure that you have all the sites selected with the exception of the *Bam*HI site. Now choose Format > Site Markers > Show Site As Symbol. You have something resembling Figure 2.9, page 2-9C. Note that you might not have the same set of symbols as shown in this figure.

9. Choose File > Save As..., name the file **construct#5**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later. You will need it again in the next tutorial. That tutorial will explain how to use your new vector as the recipient of foreign DNA.

You can quit now, or continue on to Tutorial 6: “Cloning a DNA Segment and Silent Mutations”, page 2-23. If you plan on continuing, you can leave the file you just created, **construct#5**, open. If not, close this file.

c. Note that you could accomplish the same result by clicking on a single site marker and then holding down the shift key and clicking on every other site marker except the *Bam*HI site.

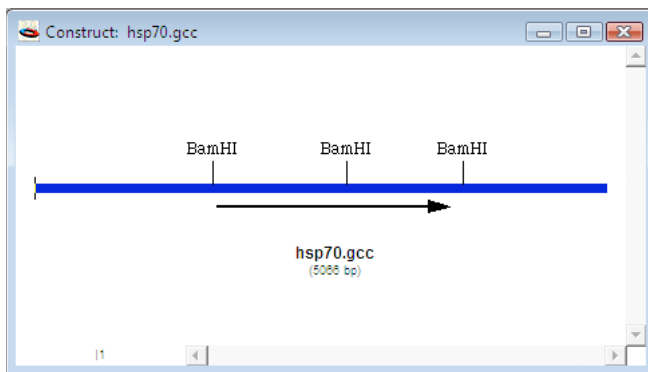
## TUTORIAL 6: CLONING A DNA SEGMENT AND SILENT MUTATIONS

Our goal in this tutorial will be to take the construct we have been working with in previous tutorials and use it to receive a new segment of DNA. In doing this you will learn about copying and pasting segments of DNA and how to keep track of the history of your construct.

1. Start **GCK** and open the file **construct#5** that you saved in the previous tutorial. If you do not have this file, you can start with **construct#5** from the tutorials folder provided.
2. Open the file called **hsp70** from the tutorials folder. This file contains the gene<sup>d</sup> we will be using as the source of the fragment being cloned. Notice that there is a single region indicating the open reading frame. We will be cloning a segment containing this region into the *Bam*HI site on **construct#5**.
3. The first thing to do is mark the *Bam*HI sites in **hsp70**. Make sure that the window with **hsp70** is in front and then choose Construct > Features > Mark Sites [or press command-M (Mac)/ctrl-M (Windows)]. Select *Bam*HI in the left hand list and press the **Add ->** button to add this enzyme to the *Enzymes to mark* list on the right (see Figure 2.8, page 2-8). Select to display new sites **as enzyme names**. Press **OK** to mark the sites. You will see Figure 2.22.
4. To cut out the segment we are interested in we need to address the problem of the middle *Bam*HI site. In the lab, you could do partial enzyme digests and obtain the fragment containing the entire coding region by elution of the fragment from a gel. You can also select this entire region in **GCK** by dragging with the mouse from the first (left) *Bam*HI site in the DNA to the third *Bam*HI site. This fragment can be copied and pasted into the vector. An alternative is to eliminate the middle *Bam*HI site through a **silent mutation**. This mutation will eliminate the *Bam*HI site without interfering with the coding capacity of the DNA. This is the approach we will take in this tutorial.

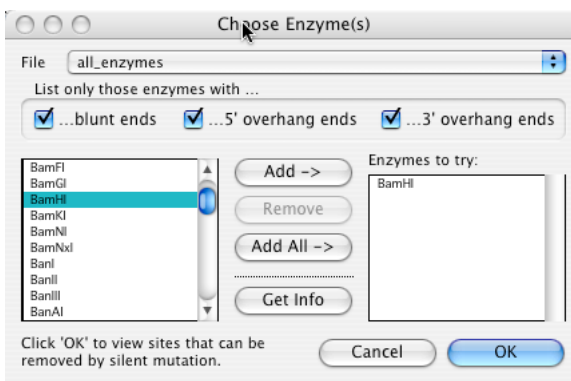
---

d. Note that this sequence has been derived from an hsp70 gene from *Drosophila*. The original sequence has been modified for use in this tutorial. Do not use this sequence as a real hsp70 sequence.



**Figure 2.22: BamHI sites marked on hsp70**

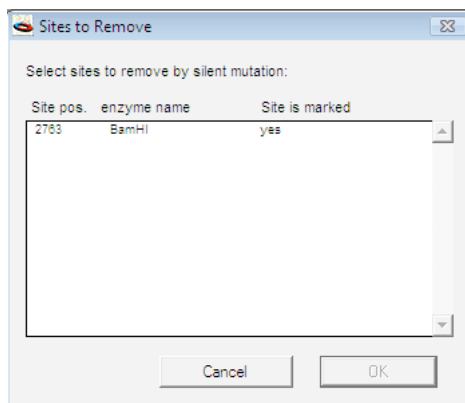
5. To make a silent mutation you first need to specify the reading frame to which you are referring. To do this, select the single region in the **hsp70** construct by clicking on it once. Now choose Construct > Features > Remove Sites by Silent Mutation. You will see the familiar dialog shown in Figure 2.23. Select the



**Figure 2.23: Select Enzyme Sites to be Removed**

*BamHI* enzyme in the list on the left and press **Add ->** to bring it to the right list of *Enzymes to try*. Click **OK**. This will bring up Figure 2.24. Click on the *BamHI* name in the list to select it (if there were more than one *BamHI* site, you could choose any one or more of them). Now press **OK**.

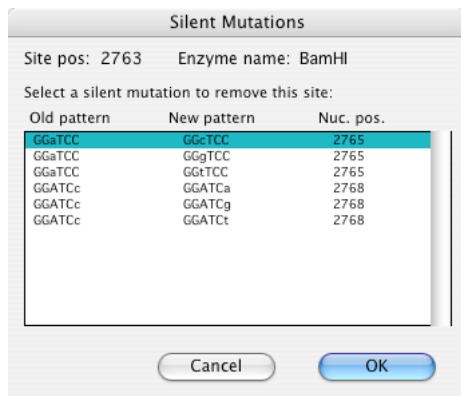
6. This will bring up another dialog containing a list of the different possibilities that could be used to remove the *BamHI* site without altering the coding information in the DNA. This is shown in Figure 2.25. The original sequence is shown in the first column and the possible changes are shown in the sec-



**Figure 2.24: Selecting the Site To Remove**

ond column. The actual changed nucleotide is shown in lower case. Select the first item in the list (as shown) and then press the **OK** button.

7. You should now see the **hsp70** construct with the middle *BamHI* site missing. The actual nucleotide sequence has been changed as specified in



**Figure 2.25: Selecting a Silent Mutation**

Figure 2.25 with the changed nucleotide being indicated in lower case in the actual sequence. Choose File > Save As..., name the file **hsp70\_Bam-**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later. You will need this file again in Tutorial 10: “Making Illustrations”, page 2-50.

8. You are now ready to proceed with the cloning. Click in the DNA seg-

ment at the left *Bam*HI site and drag the mouse cursor to the right until you have selected all the DNA between the two *Bam*HI sites. The entire segment should be highlighted<sup>e</sup>.

9. Choose Edit > Copy [or press command-C (Mac)/ctrl-C (Windows)] to copy this segment to the clipboard. Make the **construct#5** window the active window either by clicking in the window with the mouse or by selecting it from the Window menu.

10. You now need to tell GCK where it is that you would like to paste the *Bam*HI fragment that is now in the clipboard. To do this you must position the insertion point at the *Bam*HI site in **construct#5**. The easiest way to do this is to click on the site marker text, *Bam*HI. Do this now.

11. You are finally ready to do the actual cloning step. Choose Edit > Paste. You will be shown a dialog warning you<sup>f</sup> that the editing operation you are about to perform (pasting in a fragment) will disrupt a region in the current construct (the  $\beta$ -galactosidase segment that spans the polycloning site). Press **OK** to that dialog. You should see Figure 2.26.

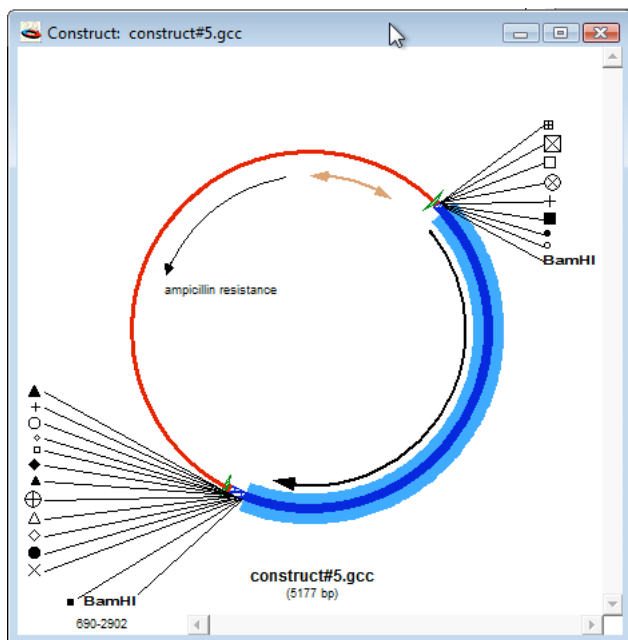
12. Notice in this figure that the coding region from **hsp70** is intact and displayed and that the size of the construct shown in the center of the window is adjusted to reflect the newly inserted fragment. Because a *Bam*HI fragment was pasted into a *Bam*HI site, both of the *Bam*HI markers have been restored.

13. Choose File > Save As..., name the file **construct#6**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later.

14. Make sure that the blue inserted DNA segment is still selected and then choose Edit > Special Paste. Since the **hsp70** fragment is still on the clipboard (in

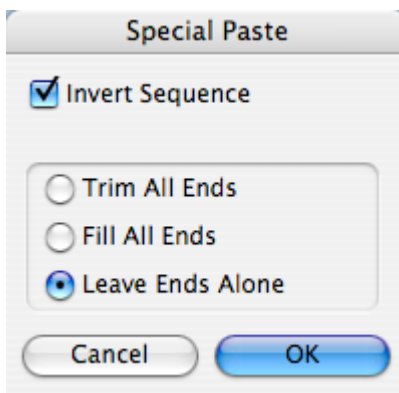
- 
- e. You might notice that the selection “pauses” at the location of the former middle *Bam*HI site. This is because the changed nucleotide has a different color and represents a “boundary” in the sequence. Selections in the graphical view occur between boundaries.
  - f. This alert is shown to warn you of editing operations that you might not be aware of. If you are pasting into a site that is part of a “feature” in the construct, the pasting operation will disrupt that feature. Therefore it needs to be removed to prevent any confusing or ambiguous annotations from remaining.





**Figure 2.26: Pasted hsp70 Fragment**

memory) and you have a segment of DNA selected, pasting will replace the current selection with what we paste into the construct from the clipboard. This is just like replacing selected text in a word processor with the contents of the clipboard. You will see Figure 2.27. Check the **Invert Sequence**



**Figure 2.27: Special Paste**

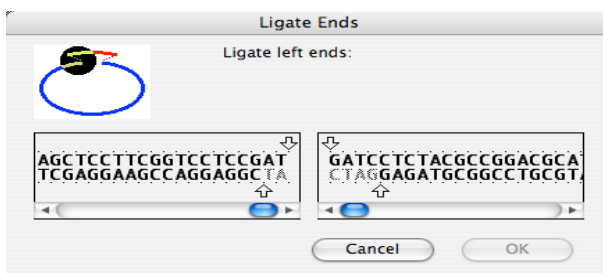
check box and select the **Leave Ends Alone** radio button. Press. **OK**. This

will replace the currently selected **hsp70** segment with an inverse version of the same DNA – i.e., it will be flipped over.

15. Choose File > Save As..., name the file **construct#7**, and save it in your own folder – *please do not replace the tutorial files provided* because others might want to use them later. Since cloning the **hsp70** fragment into the *Bam*HI site has an equally likely chance of occurring in either orientation, these constructs (**construct#6** and **construct#7**) represent two possible outcomes in a real experiment. You will be able to use these constructs and **GCK** to predict gel patterns as detailed in Tutorial 9: “Running Gels and Orientation Analysis”, page 2-45. This demonstrates how easy it is to replace DNAs and to generate constructs with inserts in either orientation.

16. Before ending this tutorial let's see how **GCK** behaves if you try to paste the *Bam*HI fragment into a site other than a *Bam*HI site. The fragment is still on the clipboard so all you need to do is define a different insertion point in the construct to be the target for the pasting operation. To do this, click on any symbol in the construct to select that site.

17. With the insertion point in the DNA at the selected site, choose Edit > Paste [command-V (Mac)/ctrl-V (Windows)]. This will bring up a **ligation dialog**, as shown in Figure 2.28. Because the fragment ends are incompatible (*the*



**Figure 2.28: Ligation Dialog**

*selected site* in construct, *Bam*HI in clipboard), **GCK** will not let you paste the fragments together (ligate them). The sequences shown in this dialog represent the junction shown in the top left corner – in this case, the left DNA sequence is the result of the *selected* cut in the vector and the right DNA segment is the result of the *Bam*HI cut fragment in the clipboard. The arrows indicate the actual cut sites and illustrate the staggered ends. Note that the **OK** button is disabled because the operation is not biologically possible. To

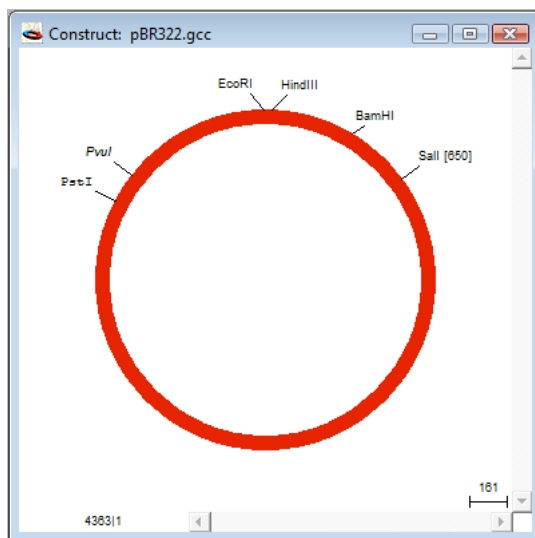
adjust the ends you can drag the arrows with the mouse until you have compatible ends. As soon as the ends are made compatible, the **OK** button will become enabled and you will be asked to deal with the “other” junction between the fragment being pasted and the target sequence. There is a shortcut to this process using `Edit > Special Paste`, which is discussed in detail in “Special Paste...”, page 7-14. For now, press **Cancel**.

18. Close **construct#6** and **hsp70\_Bam-** but do not save any changes. You may continue with the next tutorial now, or quit the program.

## TUTORIAL 7: CHRONOGRAPHY – TRACKING CLONING HISTORY

One of the most useful features of GCK is called **chronography**. This feature allows you to keep track of construct history as well as to store different views of the same sequence in a single file. It is important to realize that although you will be seeing different graphics on screen, there is only a single sequence in each construct file. Chronography allows you the flexibility of displaying the same sequence in a number of ways to bring out key features.

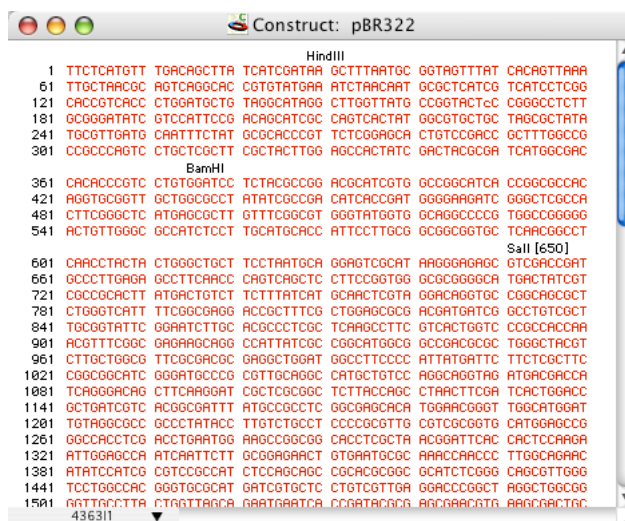
1. Start **GCK** and open the file called **pBR322** in the **tutorial files** folder. You will see Figure 2.29. This is a fairly standard view of a construct with a



**Figure 2.29: pBR322 as Graphic**

few restriction sites marked. Note that the *SalI* site shows both the name and position for the marker. This can be set in the **Format > Site Markers** menu (see “Site Markers”, page 7-23 for more detail). The bottom right corner of the window shows a scale marker indicating the scale of the drawing. The scale marker can be shown or hidden by using the **Construct > Display > Show/Hide Sites**.

2. Choose **Construct > Display > Display Sequence** to bring up Figure 2.30, page 2-31. As you will see, chronography can be used to store the appearance of a



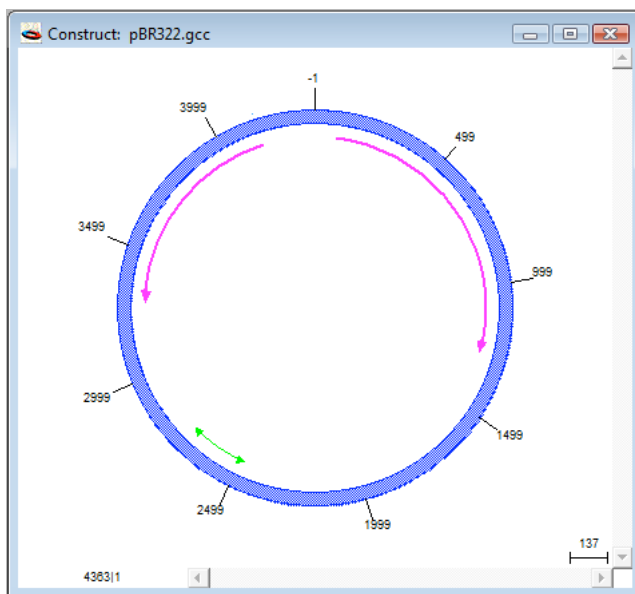
**Figure 2.30: pBR322 as Text**

construct sequence as well as the graphical construct appearance. This view of the sequence is a simple one, consisting just of the sequence grouped in tens and a few sites marked.

3. Choose **Construct > Display > Display Graphics** to return to the graphical view as in Figure 2.29. All the chronography features are accessible in the **Format > Chronography** submenu. Each different view of the construct is referred to as a different **generation**. Notice that at the bottom of the Chronography menu are three different named generations - restrictions sites, regions of interest, and DNA sources. These are the three different generations present in the pBR322 construct. The generation currently being shown is checked in the menu. Choose **Format > Chronography > Show Previous Generation**. This will bring up Figure 2.31, page 2-32, which is one generation in the past and is referred to as generation#1 (the most recent generation is referred to as generation#0). In this generation the coding regions are shown along with an origin of replication.

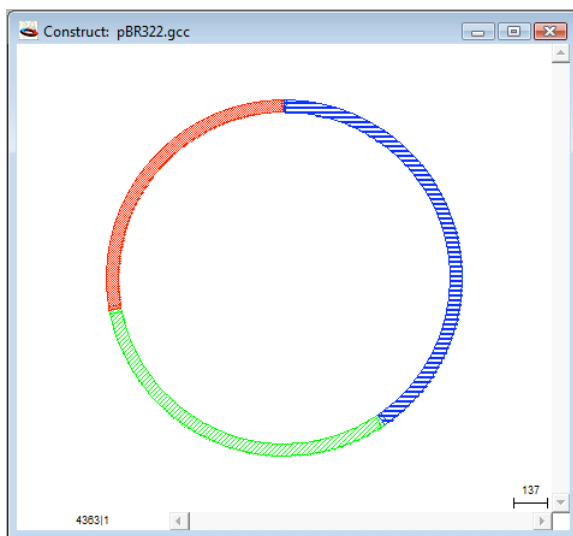
4. Choose **Construct > Display > Display Sequence**. You will see Figure 2.33 Note that the sequence is now grouped in threes and the translation of the tetracycline resistance gene is shown. *This is not a different sequence - it is just being displayed differently.*

5. Choose **Construct > Display > Display Graphics** and then choose **Format > Chronog-**



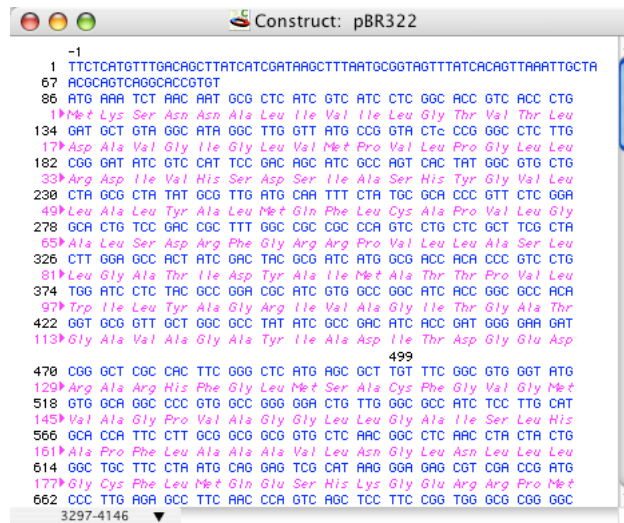
**Figure 2.31: pBR322 Generation#1**

raphy > Show Previous Generation again to see the most ancient generation for this construct. This is Figure 2.32, page 2-32. This generation shows the three



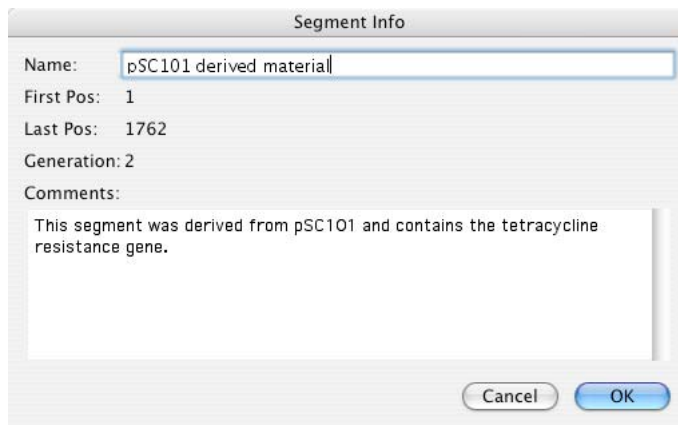
**Figure 2.32: pBR322 Generation#2**

different segments that were used to construct the original pBR322 plasmid..



**Figure 2.33: pBR322 Generation#1 Sequence**

6. Double-click on the blue segment from about 12 o'clock to 5 o'clock to select that segment. Now choose **Construct > Get Info...** [or press **command-I (Mac)/ctrl-I (Windows)**]. You will see Figure 2.34. This provides information



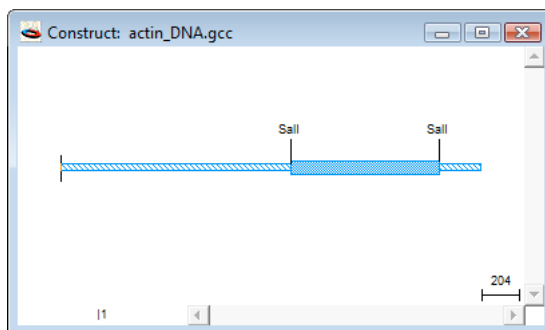
### Figure 2.34: Segment Info

about the source of each DNA segment in the construct. By carefully documenting the sources of each DNA segment, the entire history of a construct

can be maintained easily. All the comments entered can also be searched using the File Searching capabilities of **GCK** (see Tutorial 8: “Finding Comments and File Searching”, page 2-38). Press the **Cancel** button to return to the graphical window. Select the entire DNA of the construct by choosing Edit > Select All [or pressing command-A (Mac)/ctrl-A (Windows)].

7. Choose Format > Chronography > Restriction Sites to bring you back to the most recent generation of the construct. Remember, this is really just a different view of the same sequence you have been looking at throughout this tutorial.

8. Choose File > Open and open the file called **actin DNA**, which is in the **tutorial files** folder. Double-click on the *Sa*II fragment to select it (Figure 2.35), and then choose Edit > Copy. We will be cloning this fragment into



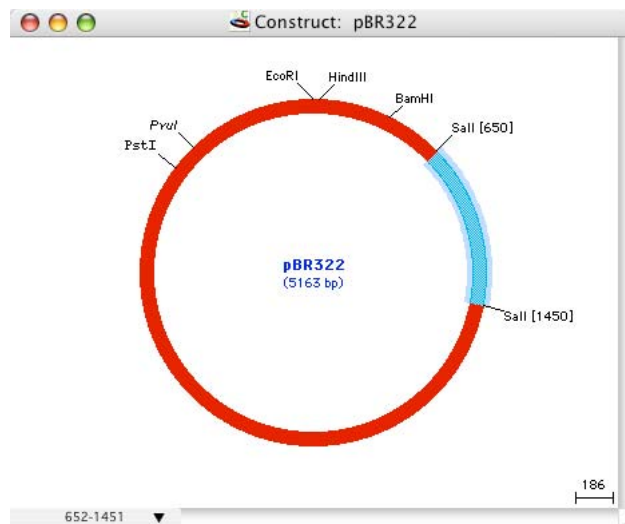
**Figure 2.35: *Sa*II Fragment of Actin DNA**

**pBR322.**

9. Click once in the **pBR322** window to make it the active window and then put the insertion point at the *Sa*II site on the **pBR322** DNA. You can do this either by clicking in the DNA at the location of the *Sa*II marker or by clicking on the *Sa*II marker itself. Once the insertion point is correctly located, choose Edit > Paste to paste the **actin DNA** *Sa*II fragment into the **pBR322** at the *Sa*II site. You will get a warning that you are about to delete a region (the tetracycline region in generation#1 is there, even though you cannot see it - click **OK**) and then you will see Figure 2.36, page 2-35. Note that the positions in the *Sa*II markers have been updated and the size of the construct has been updated in the center of the window.

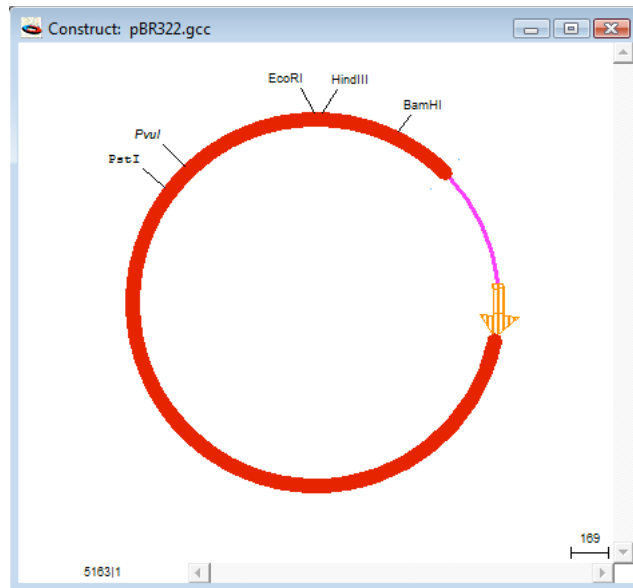
10. Choose Edit > Select All to select the entire construct and then choose Format





**Figure 2.36: Actin Fragment into pBR322**

> Chronography > Show Previous Generation. This will bring up Figure 2.37. What you

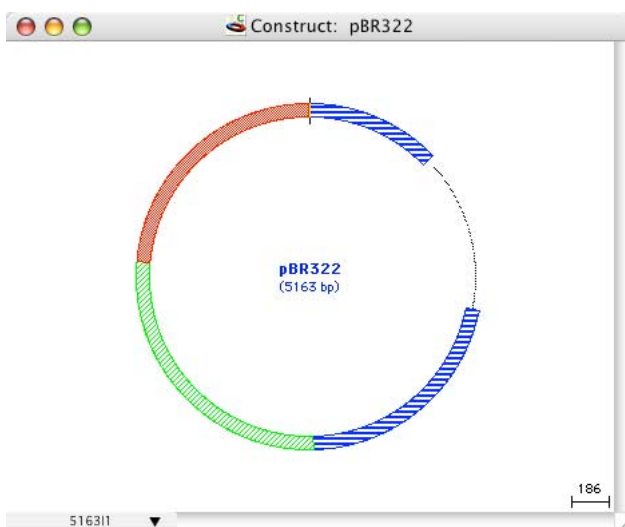


**Figure 2.37: Actin Fragment into pBR322, Generation#1**

see here is that the previous generation of the **actin** DNA was copied when the actin fragment was copied. *All generations of a fragment of DNA are cop-*

*ied when that fragment is copied.* This provides an easy way to keep track of where all the different pieces of DNA have come from when you assemble a construct of your own. If comments are entered with each segment, it is always possible to figure out the entire construct history by looking at different generations.

11. Again, choose **Format > Chronography > Show Previous Generation** again. This brings up Figure 2.38, page 2-36. Notice here that the actin fragment is just



**Figure 2.38: Actin Fragment into pBR322, Generation#1**

a plain gray line because there was no generation specified for the actin DNA that far into the past. Click once in the DNA at the beginning of the actin fragment to deselect the entire construct.

12. Choose **Construct > Display > Display Sequence** to view the construct as a sequence in generation#2. Notice how the gray from the actin fragment is also used to display the sequence itself. You can use different generations to keep different sequence appearances just as you can keep different graphic appearances.

13. You may find that in some complex construction projects, you might accumulate a large number of generations and it might be difficult to navigate from one generation to the next by only one generation at a time. In **GCK**, you can jump to any generation by using the **Format > Chronography > submenu**.

There will be a list of all generations at the bottom of this menu. Select the Regions of Interest **generation** to go to that generation.

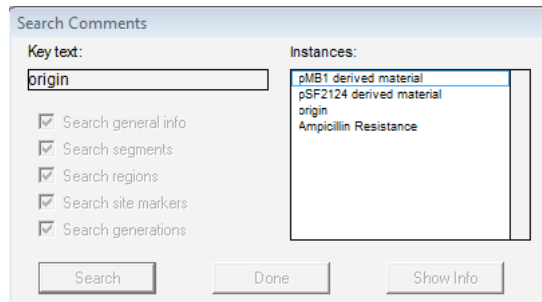
14. Explore the different formats for this generation. Change back to the graphical view now (Construct > Display > Display Graphics). Try selecting a segment and then changing the generation for that segment. Note how it is possible to show different generations for different segments at the same time. This provides a great deal of flexibility for displaying sequences and graphics in exactly the way you want.

This concludes the tutorial. You will not need the construct you have just made so close any open files and **do not save any changes**.

## TUTORIAL 8: FINDING COMMENTS AND FILE SEARCHING

You have seen in previous tutorials how you can enter comments in association with site markers, segments of DNA, chronographic generations, and regions of interest. Given that you may have many generations in a construct, it might sometimes be difficult not only to remember where in a construct you have stored information, but in which file the information was stored. In addition, if you store information such as storage locations for a given construct as part of the general information associated with each construct, it is useful to be able to search quickly through all the files you have to obtain a list, for example, of all the constructs in the freezer in room 211. This is done using the **File Searching** capability of **GCK**.

1. Start **GCK** and open the file **pBR322** in the **tutorial files** folder. You might recall that there is an origin of replication in this vector but it is not visible in the view you see, nor can you remember with which generation it is associated. You can identify the location of this feature by using the Search Comments function.
2. Choose **Construct > Search Comments...**, which will bring up a dialog box. Type the word “origin” into the Key Text field and press **Search**. This will bring up Figure 2.39. This search examines all comments and feature names associ-

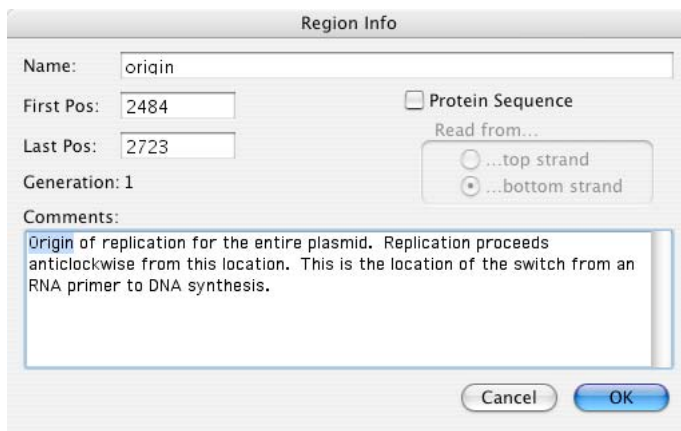


**Figure 2.39: Searching Comments**

ated with all chronography, segments, regions, site markers, or part of general info for the key work “origin”. You can limit your search to just specific

features of the construct by using the checkboxes on the left of the dialog box.

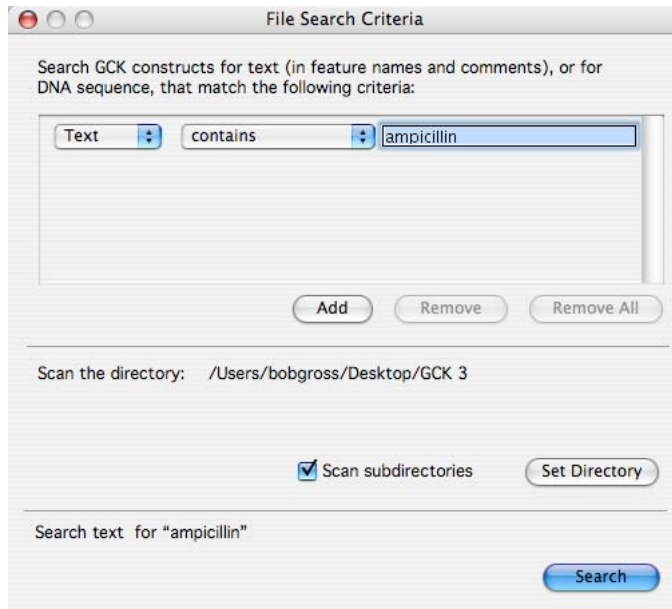
3. The list that appears in the right of Figure 2.39 contains all those features which have the word “origin” in their names or in their comments. Click on the word “origin” in the right side list and press **Show Info** to see the context of the match. This will bring up Figure 2.40. This is the standard



**Figure 2.40: Origin Found**

region “get info” box. Note that the word “origin” is highlighted in the *Region Comments* field so that you can easily see where it is located. From this dialog, you know that the origin of replication is from nucleotides 2484 to 2723 and can be found as a region in generation#1 (as indicated in the bottom left corner of the dialog).

4. Searching comments is very useful for looking just within a single construct, but what about looking through all the files on a disk or in a specific folder or even in a folder on a common file server? This is where the file searching capability comes in. **File Searching** allows you to set up sophisticated search criteria and find files on your disk. Choose Tools > Search Files... to bring up Figure 2.41, page 2-40. In this dialog you enter your search criteria in the lines at the top of the dialog. For now, we will do a simple search. Just type in “ampicillin” in the top line. Note that you could just as easily search for something like “room 211-3B” to find all the constructs stored in the freezer in room 211, Box 3B - IF you had entered this in the comments. The goal here is to find all of the files that have comments indicating that they



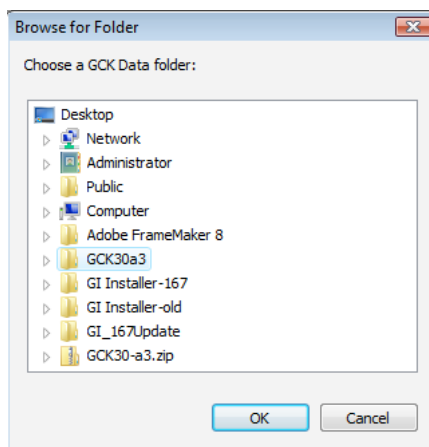
**Figure 2.41: Searching Files**

contain the word “ampicillin” in their comments - this is not the same as searching for the DNA sequence. If you are careful in entering comments for each of your files, File Searching can work as a searching tool for your database of constructs.

5. Note that the search query box will display the text you are searching for. In the *Find Match In* section of the dialog, you can specify if you want to search DNA sequences or comments (and titles). Click the radio button that says **Any Comments**. You next need to specify the *Search Directory*. This is the folder that you wish to search to find matches satisfying the search criteria. If you choose a folder that has other folders inside it, and you want to search the contents of those folders too, you should check the **Check Sub-directories** checkbox at the bottom of the dialog.

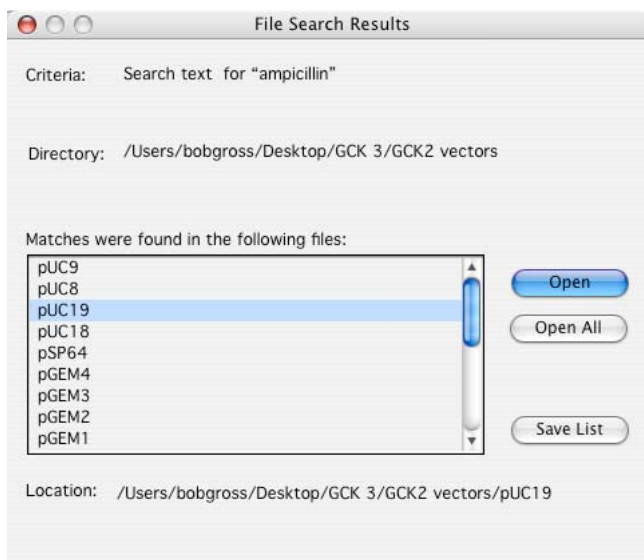
6. Press the **Set Directory** button to define the directory (folder) you wish to search. You will see Figure 2.42. Select the directory containing the files you wish to search. In this case, just select the **GCK3 folder**.

7. Now you are ready to do the search. Press the **Search** button as shown



**Figure 2.42: Set Directory**

in Figure 2.41, page 2-40. You will see a progress indicator informing you of what is going on and then you will see Figure 2.43. This window shows the

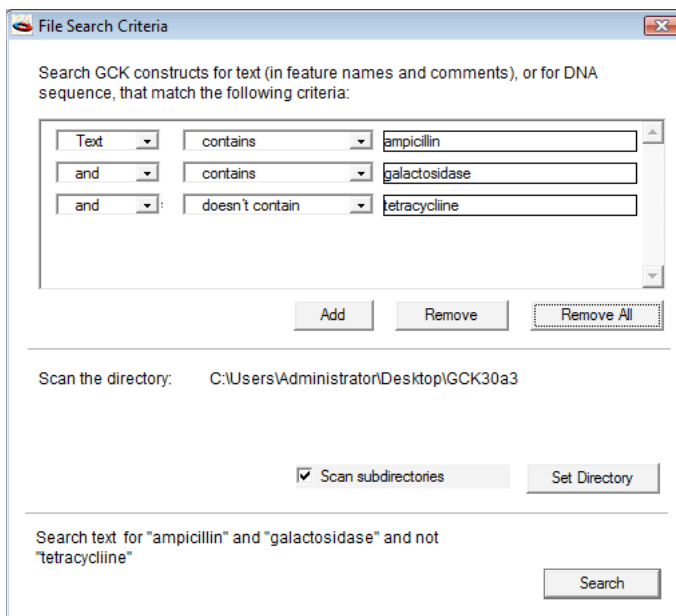


**Figure 2.43: Search Results (#1)**

Search Query and then lists all of the files that matched the criteria set in the search. Clicking on a file in the list will allow you to open it. You can also press the **Save List** button to create a text file containing all the matches with your query. This might be useful for generating a list, for example, of all

the constructs containing an ampicillin resistance gene (or those stored in room 211 freezer box 3B).

8. Let's do another search. Choose File > Search Files... again and fill in the query to match Figure 2.44, page 2-42. In this case, the search is more



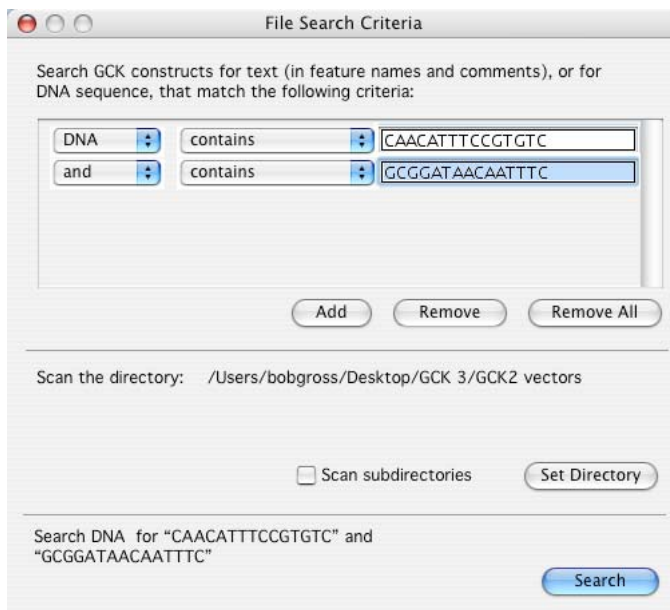
**Figure 2.44: Searching Files (#2)**

complex. We are looking for files that contain ampicillin **AND** galactosidase but do **NOT** contain tetracycline. You can try different combinations of the **and**, **or**, and **contains/doesn't contain** popup menus (these are sometimes referred to as Boolean operators) to see how they work by looking at the search query box at the bottom of the window. Set the query to match the figure and then press **Search**.

9. You will see another list similar to the one shown in Figure 2.43, page 2-41. But what if someone did not enter all the comments that should have been entered? or what if the files were simply imported sequences that had not been annotated (contained no comments)? If comments were in fact incomplete, the search results would not be that meaningful (especially if you are looking for the **absence** of certain text items like tetracycline). To address this issue, **GCK** allows you to search using DNA sequence information



instead of comments. Choose File > Search Files... again. Now select the radio



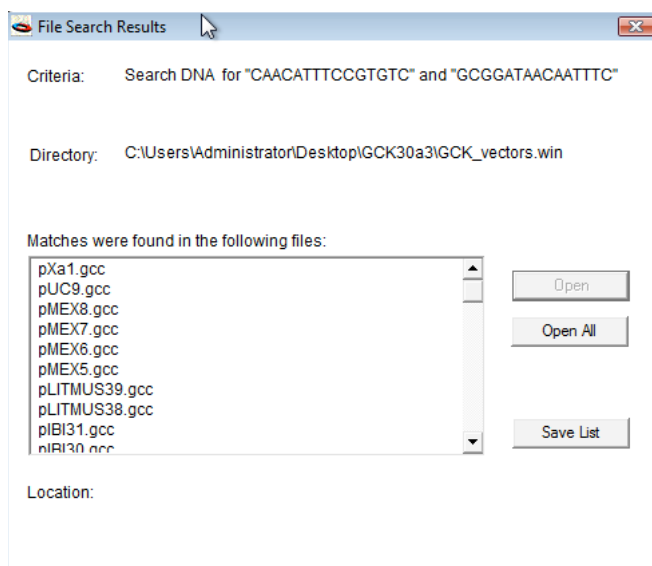
**Figure 2.45: Searching Files (#3)**

button that says **DNA Sequence**. We want to find constructs that contain an ampicillin gene and also contain the  $\beta$ -galactosidase gene. You can use a 15 nucleotide sequence to represent each gene. For ampicillin resistance ( $\beta$ -lactamase gene), type in CAACATTTCCGTGTC (you can use lower case too); for  $\beta$ -galactosidase, type in GCGGATAACAATTTC. Your screen should match Figure 2.45, page 2-43. Press **Search** when you are ready.

10. The search results you obtain will be accurate because every file has a DNA sequence. Thus, if a file has both query sequences, it will appear in the list. Your search results should be similar to Figure 2.46. In this case **pGFP** is selected so the pathname to get to the **pGFP** construct is shown in the Search Directory area of this window.

11. This concludes this tutorial. You should close all open windows at this time (do not save changes to **pBR322**).

File searching is a very powerful feature. You might elect to have all the constructs at your organization stored on a file server. If users are conscientious



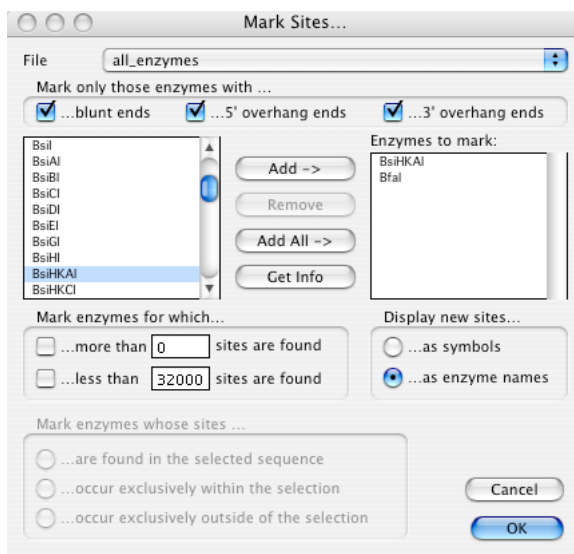
**Figure 2.46: Search Results (#3)**

in entering comments, anyone will be able to search the comments of all the files at the organization and by looking at the comments know who to contact about obtaining the construct.

## TUTORIAL 9: RUNNING GELS AND ORIENTATION ANALYSIS

Often in a cloning project there is a need to assay the success of each cloning step. Typically, this is done by gel electrophoresis. This tutorial will teach you how to run gels in GCK.

1. Start GCK and open the files **construct#6** and **construct#7** which you created in Tutorial 6: “Cloning a DNA Segment and Silent Mutations”, page 2-23. If you don’t have these files available, you can use the files that were installed in your **tutorial files** folder by the same names. Recall that these files

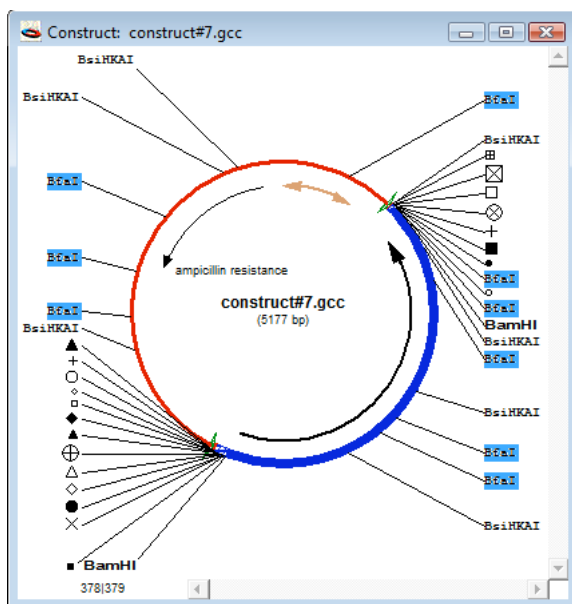


**Figure 2.47: Marking sites for orientation analysis**

represent an **hsp70** fragment cloned into a vector in two different orientations. We will differentiate between the two orientations by gel electrophoresis in this tutorial.

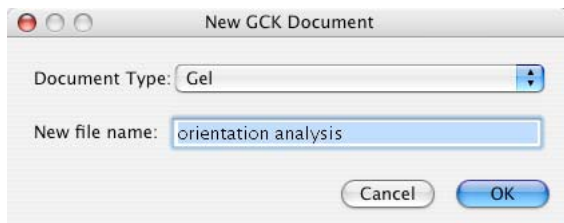
2. Bring the **construct#6** window to the front by clicking on it or by choosing it from the Window menu. Make sure nothing is selected in the DNA by clicking once in the DNA – you should see a blinking insertion point. Choose Construct > Features > Mark Sites [command-M (Mac)/ctrl-M (Windows)]. Find *BfaI* in the list on the left and double-click on it to move it to the list on the right.

Now find *BsiHKAI* and move it to the list on the right. Your dialog should look like Figure 2.47. Press **OK** once you have set up the search.



**Figure 2.48: BfaI sites marked in construct#6**

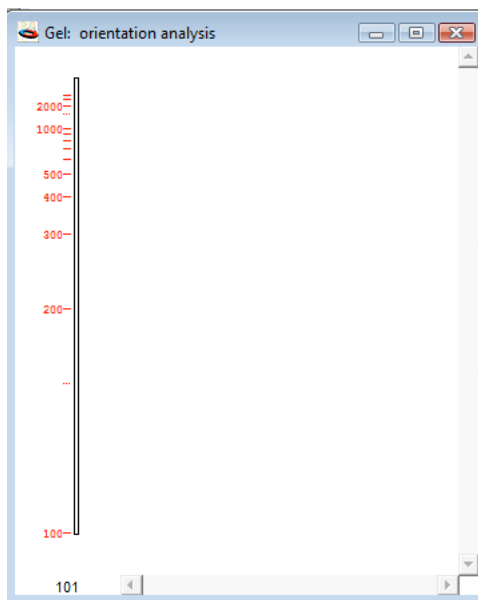
3. You will see a number of sites marked in **construct#6**. Click someplace in the window background to deselect the sites that were just marked. Double-click on one of the *BfaI* sites. Double-clicking a site name will select all other sites having the same name. Your window should look like Figure 2.48..



**Figure 2.49: Creating a new Gel window**

4. Choose **Edit > Copy** to copy the selected sites to the clipboard. Notice that you are not copying a DNA sequence, but rather a set of markers.

5. Choose File > New..., which will bring up Figure 2.49. Select the **Gel** popup menu to specify that you want to create a new **Gel window** and then type in the name **orientation analysis** for the new file name. Press the **OK** button to create the new Gel window. You will see Figure 2.50. Along the left side are



**Figure 2.50: An empty gel window**

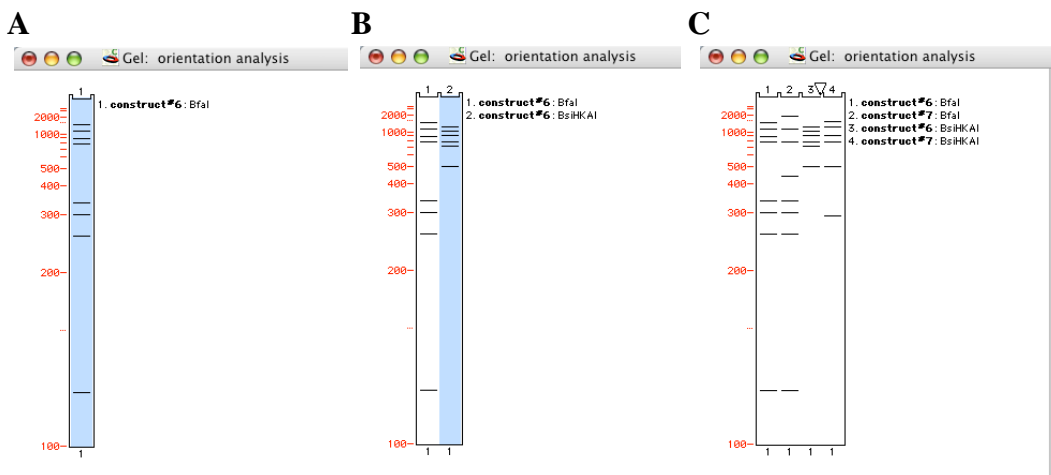
size markers. The blinking triangle at the top of the window indicates the insertion point – where the next lane will be created.

6. Choose Edit > Paste to paste the site markers<sup>g</sup> into lane 1. You will see Figure 2.51A (page 48). This shows the predicted migration of the restriction fragments on a gel. For more details on features of the Gel window see Chapter 5, “Gel Window”. Click the mouse in this window near the top right corner of the gel, just above the gel “tab” that is sticking up. This will place the blinking triangle insertion point at that location.

7. Now bring the **construct#6** window to the front again and double-click on a *BsiHKAI* site to select all of these sites. Choose Edit > Copy to copy the

---

g. In actuality, what is on the clipboard is not just site markers. Rather, it is the construct DNA along with the positions of the marked sites. This can be converted to fragments in the Gel window.



**Figure 2.51: Orientation analysis gels**

selected sites to the clipboard.

8. Return to the Gel window, make sure the triangle is on the right of the gel and choose **Edit > Paste** to paste the site markers into lane 2. You should see Figure 2.51B. These are the digests you would get from **construct#6**, having the **hsp70** insert in the **clockwise** direction.

9. In order to compare these digests with ones from **construct#7**, which has the **hsp70** insert in the **counter-clockwise** direction, you need to do a similar series of steps on **construct#7**. So...

10. Repeat the digests with *Bfal* and *BsiHKA1* using **construct#7** DNA by marking sites (Construct > Features > Mark Sites — you should see something like Figure 2.47, page 2-45 again).

11. In the **construct#7** window double-click on a *Bfal* site to select all of those sites. Choose **Edit > Copy** to copy the selected sites to the clipboard.

12. Go to the **orientation analysis** gel window (use the Window menu if you cannot find it) and click the mouse at the top of the gel between lanes one and two to place the blinking triangle insertion point at that location. Choose **Edit > Paste** to paste the site markers into the new lane 2 — note that the **construct#6/ BsiHKA1** digest is now lane 3.

13. Finally, go back to the **construct#7** window, select all the *BsiHKA1* sites,

copy them and paste them into the right edge of the gel (this becomes lane 4). You should have Figure 2.51C.

14. If you would like to add a lane of standards, you can open the file called **standards gel** in the **tutorial files** folder to see several standard marker digests. Click on the lane that is of interest to you to select it (it will become highlighted), copy it, and paste it into your **orientation analysis** gel, just as you did for the digests. If you would like to see what this looks like, open the file called **orientation analysis** in the **tutorial files** folder<sup>h</sup>.

15. If you do the actual cloning experiment and run a gel, you should be able to distinguish the orientation of the insert by comparing the real gel to the predicted gel pattern shown in Figure 2.51C. Open the *Illustration*<sup>i</sup> called **real gel analysis** in the **Sample Files** folder to see a real example of this. You will learn about Illustrations in Tutorial 10: "Making Illustrations", page 2-50.

16. This concludes this (lengthy) tutorial. Thanks for sticking with it. Close all the open files now but do not save any changes.

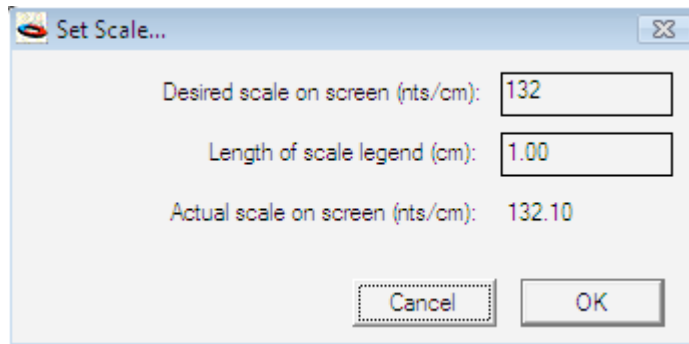
- 
- h. Remember that this file is a **Gel** file, so you need to press the **Gel** button in the open dialog in order to be able to select the file to open it. See Figure 2.1, page 2-2.
- i. Remember that this file is an **Illustration** file, so you need to press the **Illustration** button in the open dialog in order to be able to select the file to open it. See Figure 2.1, page 2-2.

## TUTORIAL 10: MAKING ILLUSTRATIONS

In the previous tutorials you worked with one file type at a time – either a Construct window or a Gel window. However, when you are designing a cloning strategy it is often desirable to include many different constructs and gels in the same document along with descriptive text. This is the function of the Illustration window. More details about the specific functions of the Illustration window can be found in Chapter 6: “The Illustration Window”. This tutorial is designed to give you a general idea of what the Illustration window can be used for.

1. Start **GCK** and open the file **construct#5**, which is in the **tutorial files** folder. This is the vector we will use in our cloning project.
2. Click in the construct and choose **Edit > Select All** and then choose **Edit > Copy** to copy the entire construct to the clipboard.
3. Choose **File > New...** and choose to create a new **Illustration** window by clicking the **Illustration** radio button (see the dialog in Figure 2.49, page 2-46). Name the new window “**cloning project**” and press **OK**. A new Illustration will be created.
4. In the new window you should see a blinking cursor that represents the top left corner of whatever you choose to paste into the window. Click the mouse about 3 cm from the top and 3 cm from the left and then choose **Edit > Paste**. This will paste **construct#5** into the Illustration window. You might want to make the window larger at this time to be able to see what is going on.
5. Notice that the pasted construct now has 8 **handles** (little square filled boxes) around its edges, indicating that you can drag it around the window and resize it. Also notice, there is now an Illustration menu in the menu bar because an Illustration window is the active window. Choose **Illustration > Set Construct Scale...** and enter 300 into the dialog as shown in Figure 2.52, page 2-51. Press the **OK** button. The set scale option in the Illustration window allows you to make sure that all of the constructs in any given Illustration are all drawn to the exact scale you desire. This operation will make your construct look a little ugly – the title will be too big for the construct and the arrowheads and line thickness for the orange region in the middle will be too





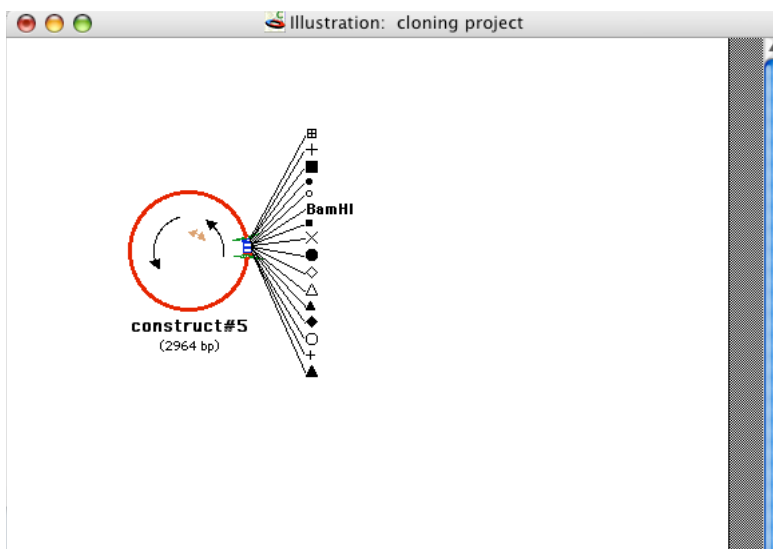
**Figure 2.52: Set Construct Scale...**

large. They need to be fixed.

6. One of the most powerful features of the Illustration window is that objects can be edited as if they were in their own windows. To access the editing, simply double-click on the **construct#5** - this is called making the object a **target**, or **targeting** an object. Target **construct#5** now. Note that there is now a Construct menu in the menu bar in addition to the Illustration menu. Click on the construct title and drag it below the circle. Adjust the line thickness and arrowhead size for the orange region by using Format > Lines >. If region names are shown, hide them using Format > Regions > Hide Region Names. You should try to get something that looks like Figure 2.53, page 2-52.

7. Now let's put a title in the illustration. Click on the "T" icon in the lower left corner of the Illustration window to activate the **Text tool**. Click the mouse near the top left corner of the Illustration window. You will see a blinking insertion point. *Before you type anything you need to set the font and font size.* Using the Format menu, choose **Arial** as the font and set the font size to **24**. Now type "An Important Cloning Project". Don't worry about the appearance of the text for now – we will fix it in the next step.

8. What you will see is large letters of text overwriting the construct. To fix this you need to adjust the size of the **text object** you just created. You are currently editing the text within the text object (it is targeted) but you want to be able to resize the text object as a whole. This is similar to being in the mode of editing a construct (as you just were) and wanting to change the size of the construct. What you need to do is **deselect** the text object by clicking in a blank area of the Illustration window, and then select the text



**Figure 2.53: Illustration#1**

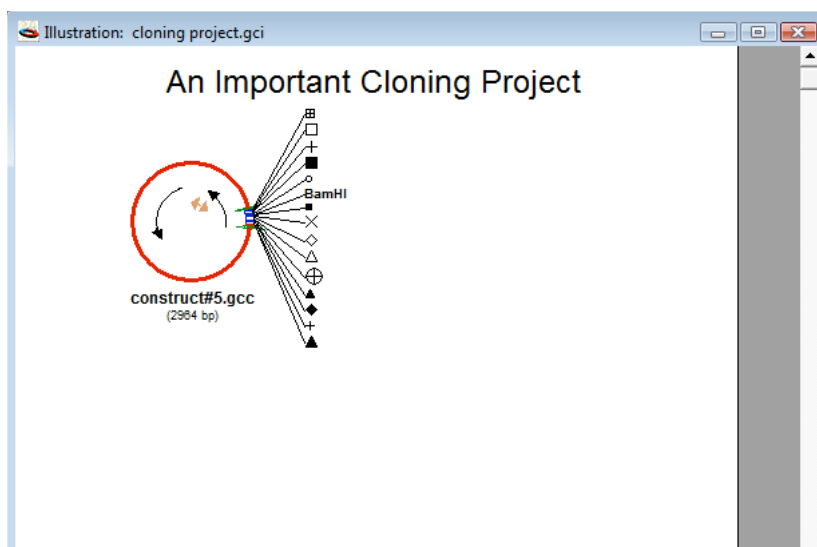
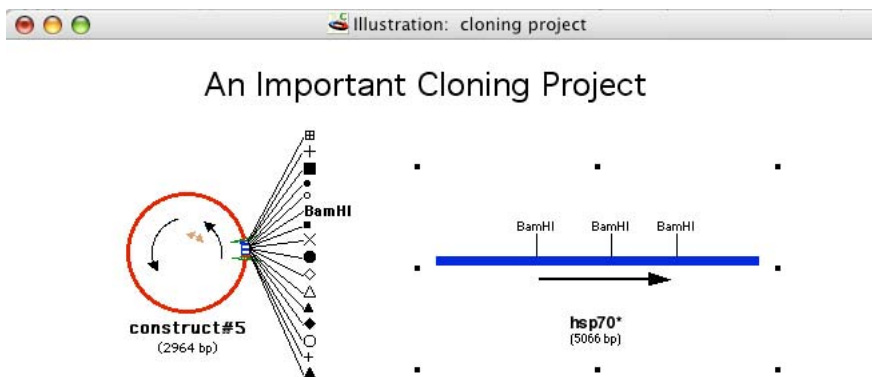
object by clicking once on it. This will give you the eight handles around the object. Use the top right handle to drag the right edge of the text object towards the right edge of the window. You will end up with something like Figure 2.54.

9. Open the file **hsp70**, which is in the **tutorial files** folder. Mark all the *Bam**HI* sites as you did in Tutorial 2: “Marking Sites”, page 2-8 (there should be three sites).

10. Click once in the **hsp70** DNA and then press command-A/ctrl-A (or choose Edit > Select All) to select the entire construct. Copy this construct to the clipboard by pressing command-C/ctrl-C (or choose Edit > Copy).

11. Bring the Illustration window to the front by clicking on it or by using the Window menu. Click the mouse to the right of **construct#5** and paste (Edit > Paste). This construct is much too large to fit on this page, so choose Illustration > Set Construct Scale... and set the scale to **800 nts/cm**. You should have a figure that looks like Figure 2.55.

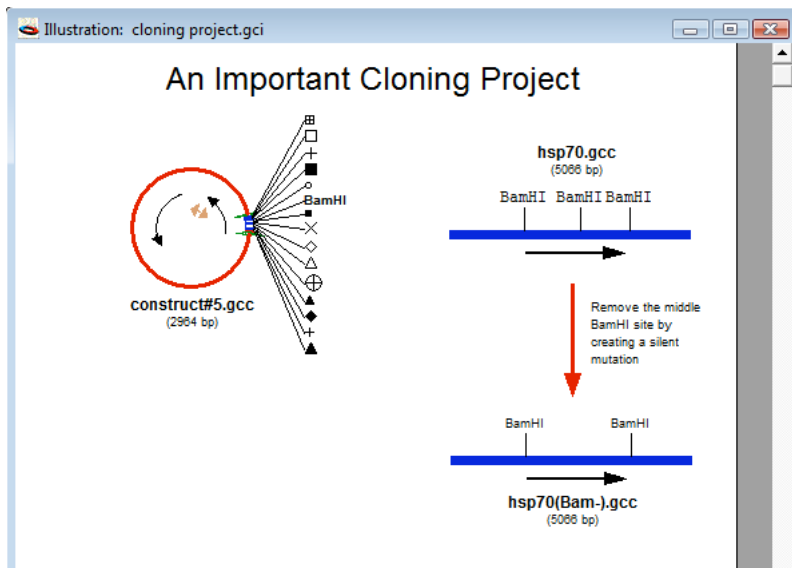
12. Now open the file **hsp70(Bam-)**. Select the entire construct, copy it, and paste it below the **hsp70** construct in the Illustration window. As you did in the last step, set the scale of this construct to 800 nts/cm.

**Figure 2.54: Illustration#2****Figure 2.55: Illustration#3**

13. We want to indicate in the Illustration what was done to get from **hsp70** to **hsp70(Bam-)**. To do this, we need to type in some descriptive text between

these two constructs. To make room for this text, you first need to move the construct title for **hsp70** to the top of the construct itself (we are creating Figure 2.56 if you want to look ahead). Double-click on the **hsp70** construct to make it the target and then drag the construct title above the DNA.

14. Now click on the “T” tool again in the bottom left corner of the window so that you can enter your description of the procedure. Click between the two linear constructs (you will get a blinking insertion point) and set the font to Arial 9 point plain text using the Format menu. Type in a description of the



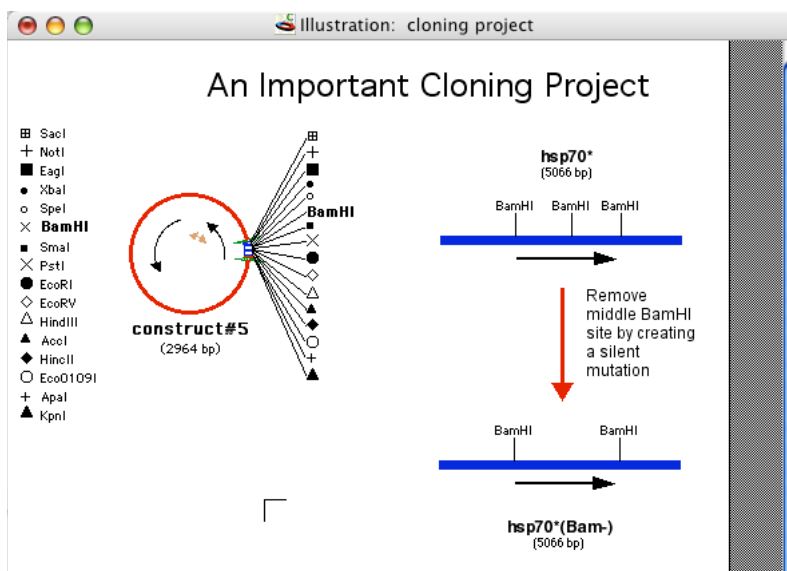
**Figure 2.56: Illustration#4**

step used to modify **hsp70**.

15. Click on the **arrow tool** (the third tool from the left) in the bottom left corner of the Illustration window and draw a vertical arrow from top to bottom as shown in Figure 2.56.

16. We now need to create a legend for the site marker symbols in **construct#5**. Double-click on **construct#5** to make it the target. Now click on one of the site markers and press command-A/ctrl-A (Edit > Select All) to select them all. Hold down the shift key and click on the *BamHI* marker to deselect it. Choose command-C/ctrl-C (Edit > Copy) to copy all the sites to the clipboard.

17. Click in the Illustration window under **construct#5** and paste the site markers into the window (command-V/ctrl-V). Drag the legend you just created to the left of **construct#5** making it look like Figure 2.57.



**Figure 2.57: Illustration#5**

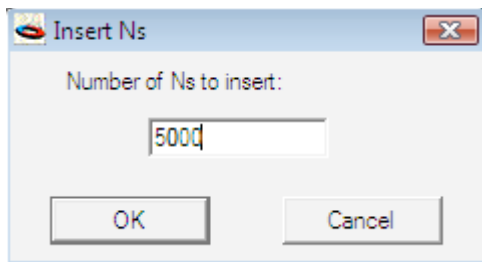
18. There are many more steps to complete this project but you should get the flavor of the Illustration window by this time. You can open the file **cloning project** in the **tutorial files** folder to see what the final illustration might look like. Feel free to explore the completed illustration to learn some other tricks. Notice also that you can copy the gel lanes and paste them into the Illustration window to complement the other information in that window. The **Sample Files** folder also contains some interesting examples.

This concludes this tutorial. Close all the open windows, but please don't save changes to any of the tutorial files provided with the program so that others can do the tutorials in the future.

## TUTORIAL 11: WORKING WITH GENERIC CONSTRUCTS

There are times, especially when you are just beginning to work with a piece of DNA, when you might know the restriction map of a cloned fragment of DNA but you might not yet know the sequence. Such generic constructs can be created in **GCK** and the segments manipulated just like any piece of DNA whose sequence is known.

1. Start **GCK**. You should see an untitled Construct window as the active window. To create a generic construct of 5000 nucleotides, choose Construct > Insert Ns... (command-H/ctrl-N). This will bring up Figure 2.58. Enter 5000 and



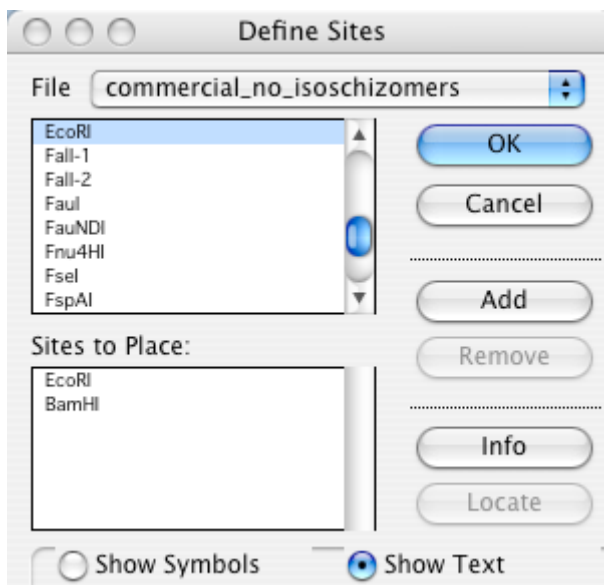
**Figure 2.58: Inserting Ns**

press the **OK** button.

2. You should now have a linear construct containing 5000 Ns. In the graphical view, it will simply appear as a horizontal line. Choose Construct > Display > Display Sequence to view the sequence as a series of Ns. Now you need to place restriction sites at specific locations in the construct.

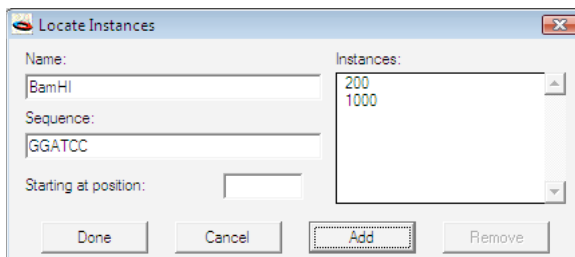
3. Choose Construct > Features > Place Sites... (command-J/ctrl-J). You will see Figure 2.59, page 2-57. This is very similar to the mark sites dialog you saw previously (Figure 2.8, page 2-8). You select enzymes in the top list and press the **Add** button to move them to the bottom, *Sites to Place*, list. For this tutorial, choose *BamHI* and *EcoRI*.

4. The next step is to define the locations of each of the enzyme sites you will be placing into the construct. This is done by first selecting an enzyme in



**Figure 2.59: Place Sites Dialog**

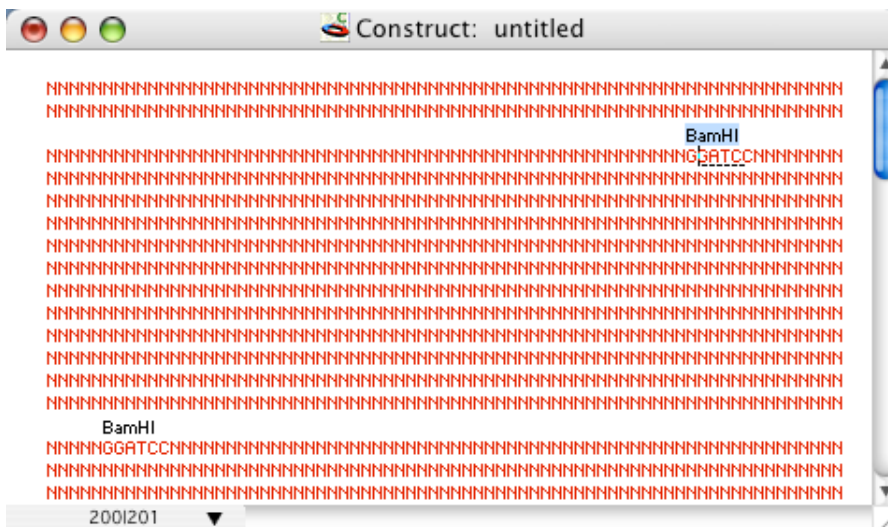
the bottom list and then defining the locations of the sites for that chosen enzyme. Select *BamHI* in the bottom list and then press the **Locate** button on the right. This will bring up the locate sites dialog shown in Figure 2.60.



**Figure 2.60: Locate Sites Dialog**

This dialog gives information about the selected enzyme and then lets you define the site locations. Type in 200 in the *Starting at position* text box and press the **Add** button to add this location to the list of sites to place – it will appear in the list of “instances” at the right. Next type in the position 1000 and add it to the list, and then add 3456. You should have these three locations in the list on the right. Once you have done this, press the **Done** button because you are done entering locations for *BamHI* sites.

5. Select *EcoRI* in the list (Figure 2.59, page 2-57) and then press the **Locate** button. In the locate sites dialog that appears (Figure 2.60, page 2-57), enter 2000 and 4000 as the *EcoRI* site locations. You have now defined the enzymes and their sites so press the **OK** button to actually place those sites.



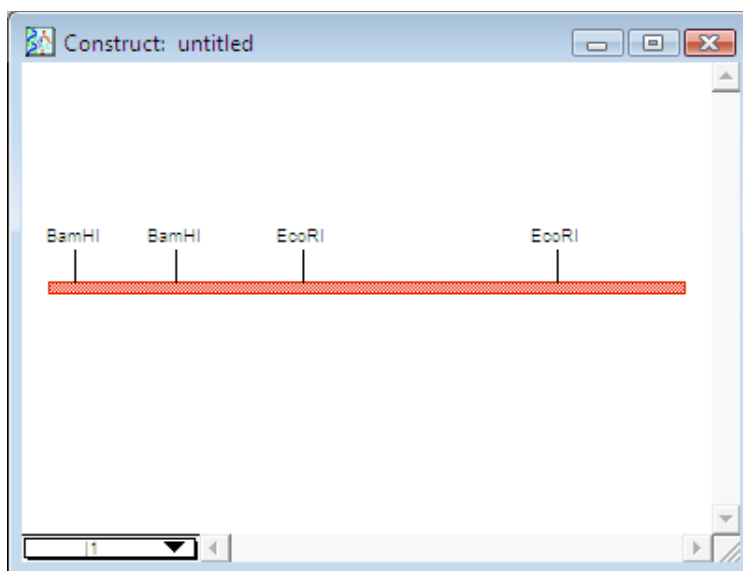
**Figure 2.61: A Placed Site (Sequence View)**

6. In Figure 2.61 you can see that the *BamHI* recognition sequence has actually replaced some of the Ns. This is therefore a “real” site that can be cut with the enzyme and used to produce fragments from this DNA.

7. Choose Construct > Display > Display Graphics. This will bring up Figure 2.62, page 2-59. You can now select fragments and use them in cloning steps as was done for fully sequenced DNAs in previous tutorials. As more sequence information becomes known for this construct, you can use the real sequence to replace the Ns in this generic construct.

This concludes this tutorial. You can close the open window (don’t save changes – you will not need this file any more).



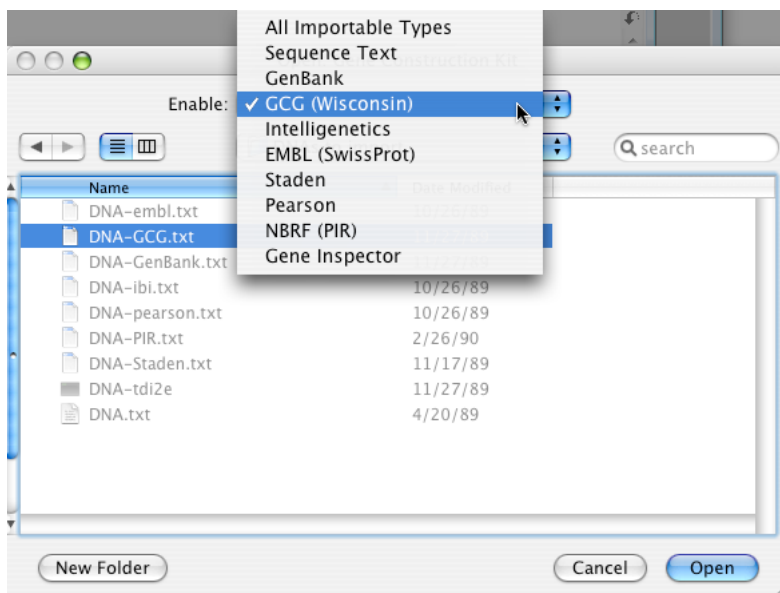


**Figure 2.62: Placed Sites (Graphic View)**

## TUTORIAL 12: IMPORTING AND EXPORTING SEQUENCES AND OTHER INFORMATION

One often obtains sequences in other formats and would like to work with them in **GCK**. This can be done by importing the sequences into **GCK**. Most other formats really are just plain text files with the sequence information organized in a specific format. Perhaps the most common of these formats is GenBank or GCG format. Just a plain sequence file (nothing but sequence) is also fairly common and is often the form of the output you might get from a sequencing facility.

1. Start **GCK** and choose File > Import to begin the importing process. You will see something like Figure 2.63. Use the popup menu above the list of files to



**Figure 2.63: Import Sequence Dialog**

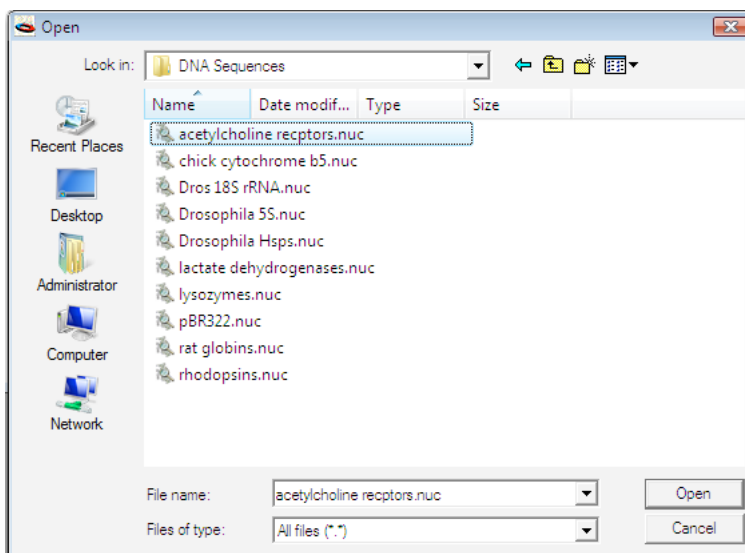
navigate to the **DNAs to import** folder which was installed when **GCK** was installed. This folder holds DNAs in a number of different file formats for you to test importing. The radio buttons in this dialog allow you to specify in which file format you believe the file to be. Clicking on the radio button might also change the list of files that are visible. Gene Inspector and DNA Inspector

files are not plain text files so clicking on one of those buttons will limit the list of displayed files to just those specific types of (non-text) files.

2. Click on the **GCG** button and select the file **DNA.GCG** to be imported as shown in Figure 2.63, page 2-60. Press **Import**.

3. The sequence will now be imported and displayed as a sequence in a new Construct window. Any comments associated with the text file in the specified format will be imported and placed in the General Info for the construct. Choose Construct > General Info... to see the comments that were part of this particular GCG formatted file.

4. Gene Inspector® files might contain multiple sequences, so you will be presented with a dialog like Figure 2.64. This shows the dialog presented

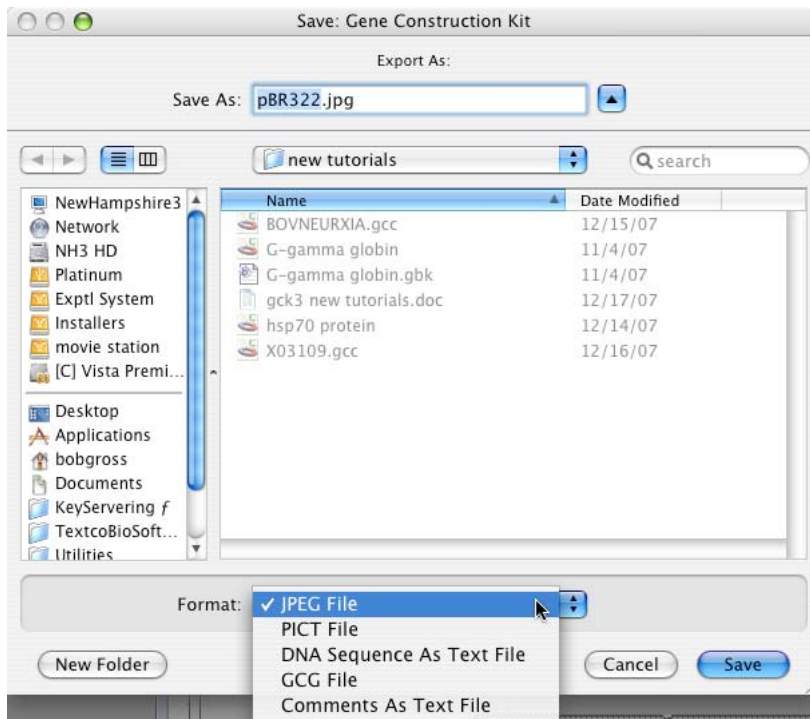


**Figure 2.64: Importing Sequences from Gene Inspector**

when a file containing multiple sequences is selected. You must choose one of the sequences and then press **Import**.

5. **Exporting** file information is another way GCK can interact with other applications. Since GCK can display information in graphical as well as sequence formats, both formats are available to export. Choose File > Open... and open **pBR322** in the **tutorial files** folder. This will open a graphical view.

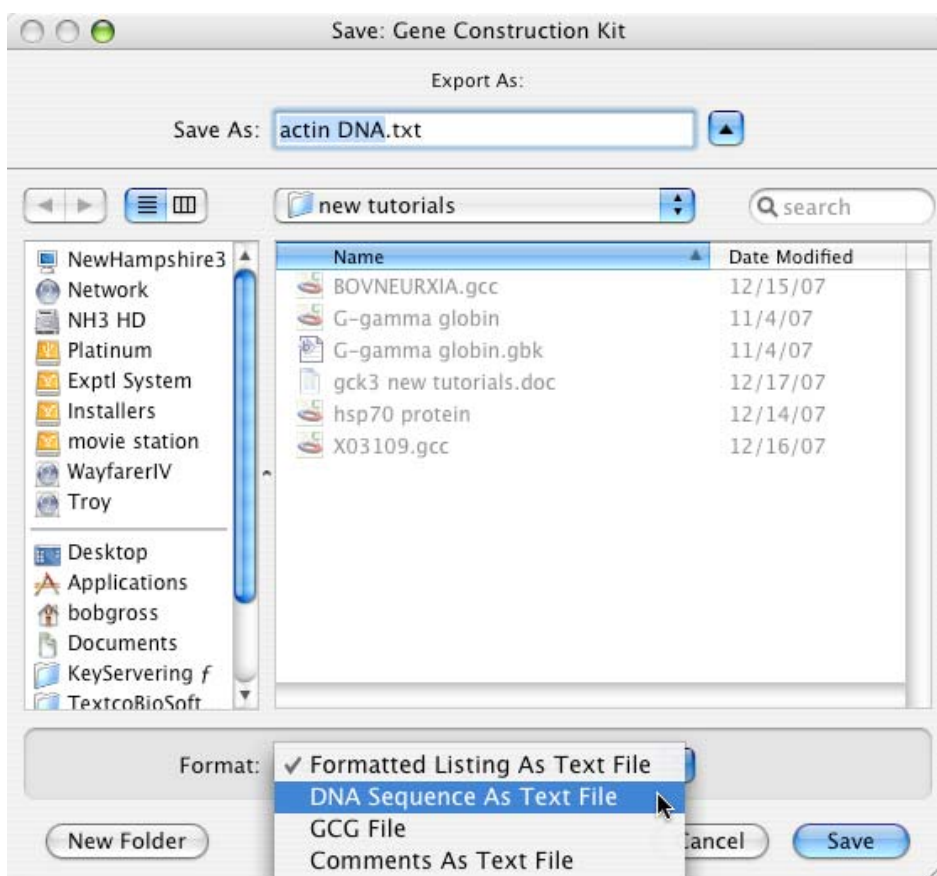
6. Choose File > Export... and you will see Figure 2.65. Choosing the **JPEG**



**Figure 2.65: Export (Graphics)**

**file** popup menu as shown here will create a file that can be imported into most graphics programs. The **DNA Sequence as TEXT file** button will create a text only file of the DNA sequence which can be opened by any word processing program as well as almost any other sequence analysis package. Click **Cancel** to dismiss this dialog unless you are really interested in playing with the output files in other programs.

7. With the **pBR322** window active, choose Construct > Display > Display Sequence to change the construct view to a sequence listing. Now, choosing File > Export... will bring up a slightly different dialog, Figure 2.66. In this case, there is an extra option called **Formatted Listing as TEXT file**. This option will produce a text file containing all the line numbers, restriction sites, spacing, and amino acid sequences. Note that in order to view these output files correctly formatted in another application, you will need to use a monospaced font like Monaco or Courier.



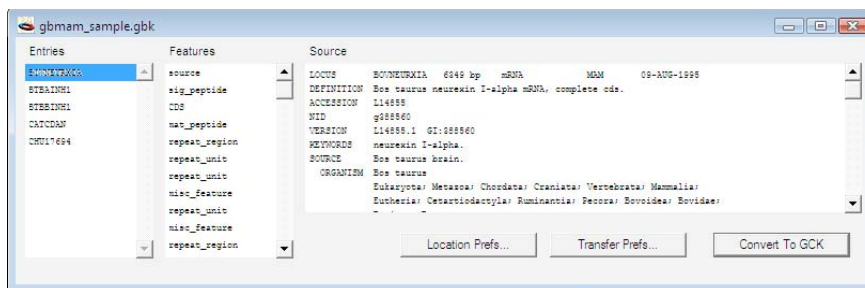
**Figure 2.66: Export (Sequence)**

This concludes this tutorial. You can close the open window (don't save changes – you will not need this file any more).

## TUTORIAL 13: IMPORTING GENBANK SEQUENCE FILES USING DELUXE IMPORTING

Through the Deluxe Import feature, Gene Construction Kit contains comprehensive features to allow GenBank searching and file retrieval over the Internet with importing directly into GCK. All GenBank features are converted one-to-one to GCK features using a conversion scheme you can customize. The following tutorial illustrates the basics of these features. Images alternate between Macintosh and Windows screenshots. The program works the same way on each platform.

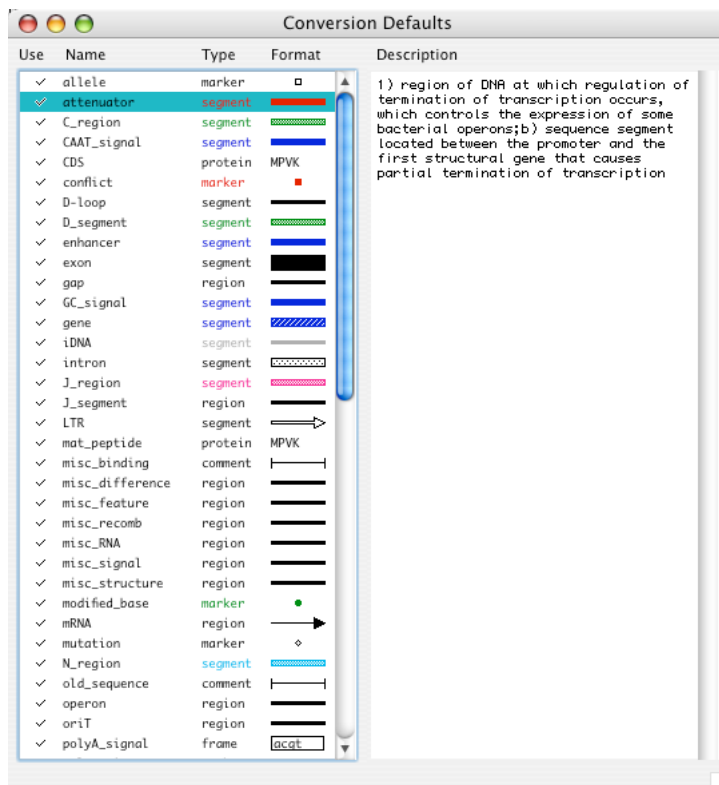
1. Select Tools > Deluxe Import > Open Sequences File. Choose the tutorial file called **gbmam\_sample.gbk** for this tutorial. You will see Figure 2.67.



**Figure 2.67: Deluxe import initial window**

2. Note in this figure that there are actually 5 different genes in the single file we are examining. These are listed along the left side of the window in the **Entries** list. The first entry, *BOVNEURXIA*, is selected in the figure. The second list, **Features**, shows all of the features of this particular entry. These are all taken from the GenBank features table. The third column (**Source**) shows the GenBank text that is associated with whatever is selected in the features column. If nothing is selected, as shown here, the **Source** field shows everything in the GenBank comments. Try clicking on a few Features to see what the associated **Source** text is.
3. The **Location Prefs...** button allows you to define how different indicators of position in GenBank files are translated into GCK positions. The

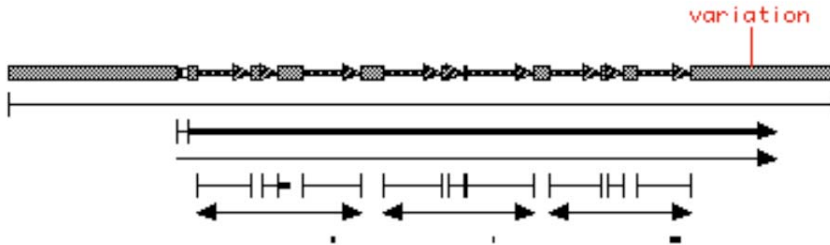
**Transfer Prefs...** button allows you to tell GCK how to handle the transfer of information into GCK. You can take a look at these choices, but for now do not change any of them. Press the **Convert To GCK** button. This will bring up the conversion default as shown in Figure 2.68. The **Use** column allows



**Figure 2.68: Deluxe Import conversion defaults**

you to toggle between a check mark and an x by clicking with the mouse. Only those features that are checked will be transferred to GCK. The second column shows the GenBank feature **Name**, while the third column (**Type**) shows how that feature will be indicated within GCK. You can change how each feature will be treated by selecting the feature and then using the Feature menu and the Format menu to change the appearance of the feature in GCK. Under Tools > Deluxe Import > you can Save or restore you choices of conversion defaults.

4. After pressing the **Convert to GCK** button you will end up with a new construct like that shown in Figure 2.69. This figure shows all of the Gen-



**Figure 2.69: Imported construct**

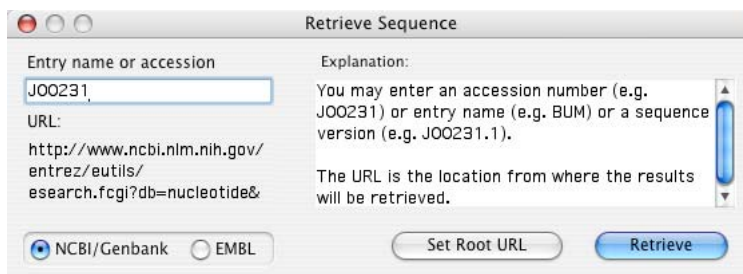
Bank features that were selected to be imported in the original GenBank file. You can click on each feature and choose [Construct > Get Info...](#) to see what that feature is.



## TUTORIAL 14: SEARCHING AND RETRIEVING SEQUENCE FILES FROM GENBANK

The following steps will demonstrate how to search and retrieve sequences from GenBank and have them automatically imported into GCK.

1. To perform the following, you need to be connected to the Internet. If you are not already on line, please connect now. Choose Tools > Deluxe Import > Retrieve Sequence to start the process. You will see Figure 2.70. Enter

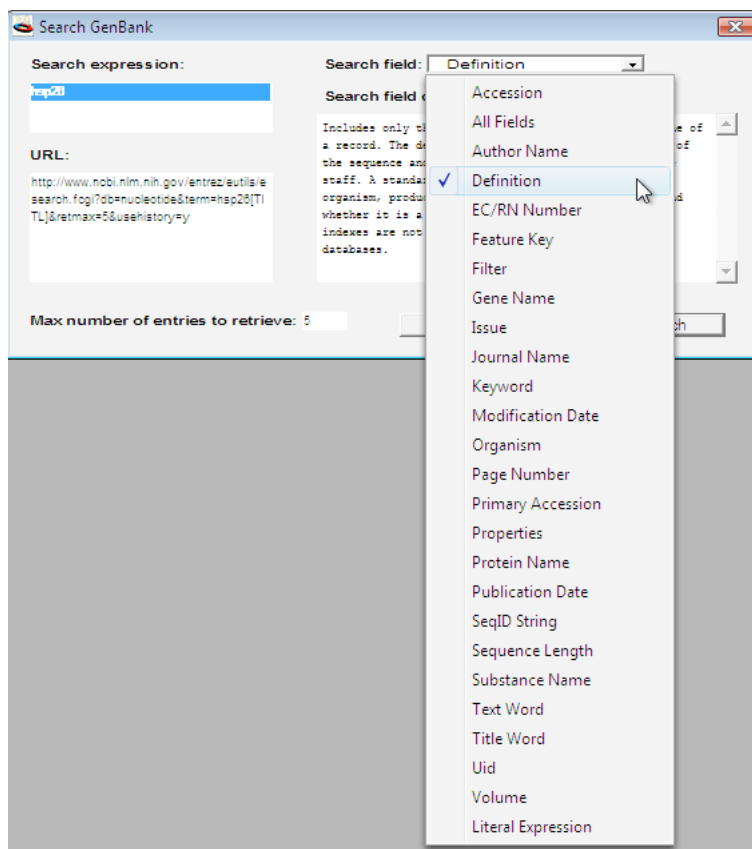


**Figure 2.70: Retrieve Sequence Dialog**

"J00231" (or any other accession number or sequence name you might be interested in) into the box in the top left corner. As you enter the accession number or sequence name into this box, you can see a URL being constructed in the bottom left box (this is not editable, but is shown for you to examine what GCK is actually doing for you). This URL will be sent to the server that you have designated. **GenBank** and **EMBL** are two default options, but you can also specify a local server by pressing the **Set Root URL...** button.

2. Pressing **Retrieve** will contact your target database and retrieve the sequence for you. Once the sequence is retrieved, you will see a window very much like that shown in Figure 2.67, page 2-64. You can proceed from that point to finish importing the sequence into GCK using the same steps described in the previous tutorial ("Cloning a DNA Segment and Silent Mutations", page 2-23).

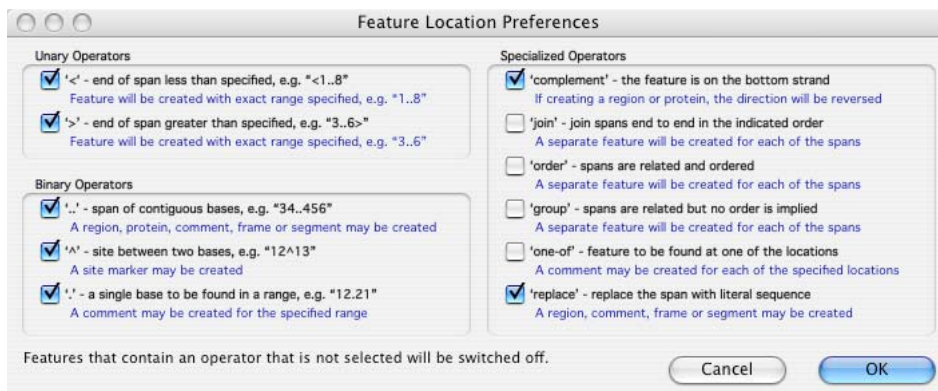
3. You can also search GenBank using GCK. Selecting File > Deluxe Import > Search GenBank will bring up Figure 2.71, which shows the search dialog for defining how you want to search GenBank. Note that the popup menu allows



**Figure 2.71: GenBank Searching Options**

you to define the database fields to be searched by your query. Type in the search expression you wish to look for and then press the **Search** button. A progress dialog informs you of progress, and then you will get a list of items matching your query. The list will be similar to see Figure 2.68, page 2-65. The sequences you are interested in are now on your hard disk and can be imported directly into **GCK**.

4. If your DNA source file has entries in the features table that might be ambiguous, you will see a button that says **Location Prefs...**. Pressing this button will bring up Figure 2.72, allowing you to define how you want the feature locations to be interpreted. This window lists the possible ways in which various features are defined by NCBI for use in GenBank files. Some of these are not directly translatable into **GCK** features, so you can define how you



**Figure 2.72: Location Preferences Dialog**

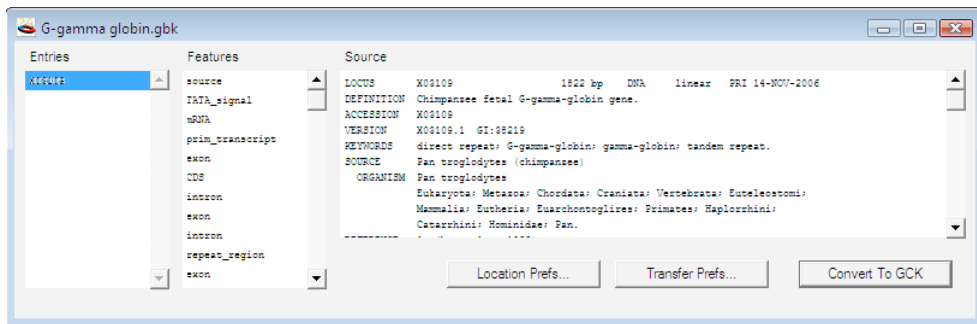
want to handle those situations in this window.

Feel free to try different operations using the Deluxe Importer. The searching works the same way as an Entrez search would work at the NCBI web site.

## TUTORIAL 15: TRANSLATING ACROSS INTRONS

When importing genomic sequences, there is typically a need to deal with exons and introns in order to determine protein sequences. This can be done easily in GCK3. This tutorial illustrates how the process works.

1. Start the application and choose Tools > Deluxe Import > Open Sequences File and then select the file called **G-gamma globin.gbk** and open the file. This file contains GenBank entry X03109, which is the fetal G-gamma globin from Chimpanzee. You could also open this file in any text editor to see what it contains. You will see Figure 2.73. For more details on Deluxe Importing see



**Figure 2.73: Deluxe importing of G-gamma globin**

Tutorial 13: "Importing GenBank Sequence Files Using Deluxe Importing", page 2-64 and "Deluxe Import", page 7-35.

2. The middle section of this window contains a list of features in this DNA. Exons, introns and CDS (coding sequences) are of interest to us in this tutorial. Clicking on a feature will provide more information about that feature in the right hand text box. Try exploring to see what is in this file. Click on the CDS feature and note that the first line starts with "join" and then lists three pairs of numbers (108..199, 322..544, and 1438..1566). This means that the coding sequence is comprised of nucleotides from 108-199 and 322-544 and 1438-1566. Note also that the exons also have ranges of nucleotides associated with them. Click the button **Convert To GCK**.

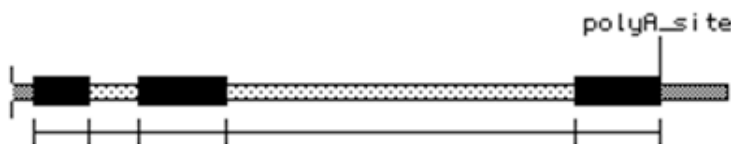
3. You will see the dialog window shown in Figure 2.74. This shows how each GenBank feature is converted into a GCK feature. You want to have all



**Figure 2.74: G-Gamma globin features**

of the exons and introns “checked” along with the “polyA\_site.” Uncheck all the other items because they will just make the display confusing at this point. When everything is set, press the **Save Construct** button.

4. A new window will open up that shows you the construct you just created. It will look similar to Figure 2.75 but may not be identical because the



**Figure 2.75: G-Gamma globin imported**

conversion choices you have might be different from the ones used here. Note that the three exons are shown as thick black lines, while the introns show a pattern of black dots. The thin horizontal lines below the construct indicate the presence of comments. You can turn them off by choosing Construct > Display > Hide Comments.

5. Double click on the first intron (200-321) and assign it a color of green (Format > Color > Green). Do the same thing for the second intron (545-1437). Note that you can see the range of nucleotides selected in the very bottom left of the construct window. This will make it easier to identify introns and exons when we switch to sequence view.

6. Choose Construct > Display > Display Sequence to view the construct as a sequence. Let's make it a little easier to read by first making the window

larger (whatever is comfortable on your screen). Choose **Construct > Display > Show Positions** to place position numbers at the beginning of each line.

7. Click somewhere in the sequence and then choose **Edit > Select All**. Choose **Format > Grouping > Group by Tens**. If you can adjust the width of your window to contain 100 characters per line, it will be easier to make selections in subsequent steps.

8. We need to now define introns. Double-click on the first intron (it is green). You should see 200-321 in the lower left corner of the construct window. Choose **Construct > Features > Define Intron**. This will “invert” the selection so that the letters appear white on a green background. Repeat this process for the second intron.

9. The next step is to define what we want to translate. From Step 2 (above) we know that the coding region starts at 108 and ends at 1566. Select this range of nucleotides - confirm that you have selected the correct range by looking in the bottom left of the construct window.

10. . Now choose **Construct > Features > Make Region**. This will allow you to translate the selected sequence. Enter a name in the dialog box and make sure to check the **Protein Sequence** checkbox. It should look like Figure 2.76. Once

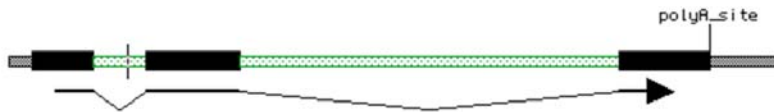


**Figure 2.76: Making a globing region**

you have this window appropriately set, press the **OK** button. You will now see the translated protein created by reading codons in the exons and skip-

ping over the sequences in the introns. Note that the codon spanning from exon 1 to exon 2 is actually broken into two pieces (ag then g to make up agg).

11. Switch back to a graphical view (Construct > Display > Display Graphics). You should see Figure 2.77. Notice that the introns are not displayed as part of



**Figure 2.77: G-Gamma globin final construct**

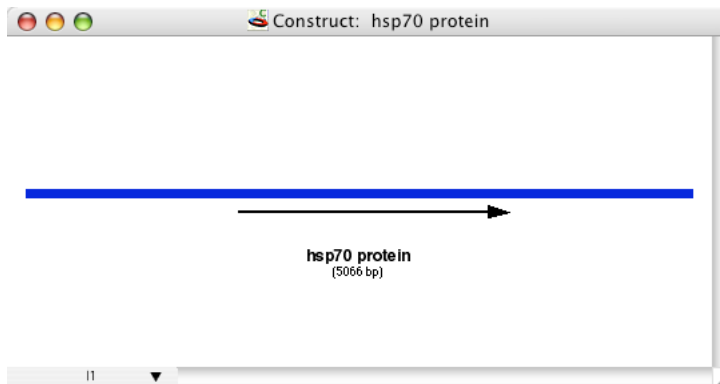
the coding sequence.

12. One final comment. In the sequence view, if you select an intron and then extend the selection (e.g. by shift-clicking) you can actually redefine the intron by choosing Construct > Features > Expand Intron. This will redefine the intron but it will not automatically update the translated protein. You have to do this manually since there might be times when you do not want to lose the protein that already exists (e.g. to illustrate alternative splicing situations).

## TUTORIAL 16: PCR ANALYSIS

We will use the PCR analysis<sup>j</sup> code to amplify a promoter region of a *Drosophila* hsp70 gene.

1. Open the file Hsp70 protein found in the Tutorials folder. You will see



**Figure 2.78: hsp70 protein construct**

Figure 2.78.

2. The arrow below the DNA is the actual transcript for this *Drosophila* heat shock gene. We want to select the 300 nts upstream of this transcript to amplify, which we suspect has an important promoter element. Double-click on the arrow to select the corresponding DNA segment. This will select nucleotides 1617-3677, as indicated in the bottom left corner of the construct window.

3. Switch to sequence view (Construct > Display > Display Sequence). The same region is selected. Using your mouse, click and drag “upstream” starting at the first nucleotide before the coding sequence, 1616. Drag approximately to

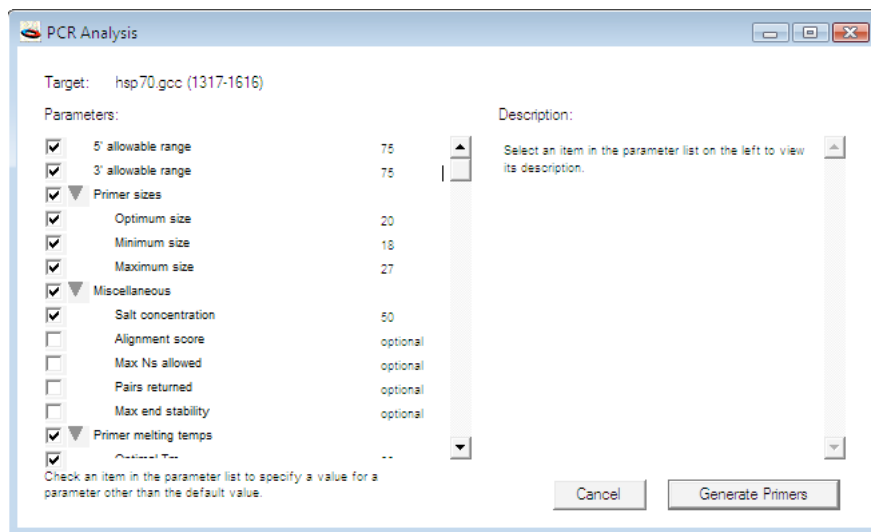
---

j. GCK uses Primer3 PCR code developed at the Whitehead Institute for Biomedical Research and used with permission. This code is Copyright (c) 1996, 1997, 1998, 1999, 2000, 2001, 2004 Whitehead Institute for Biomedical Research. All rights reserved. The algorithm for this software is published: Steve Rozen and Helen J. Skaletsky (2000) Primer3 on the WWW for general users and for biologist programmers. In: Krawetz S, Misener S (eds) Bioinformatics Methods and Protocols: Methods in Molecular Biology. Humana Press, Totowa, NJ, pp 365-386.



1317 and release the mouse button. You can fine tune the selection by holding down the shift key while selecting a new starting point in the sequence. Keep trying until you have selected nucleotides 1317-1616. Keep this selection and switch back to the graphics view (Construct > Display > Display Graphics).

4. Select Tools > PCR Analysis to bring up the PCR dialog. You will see something like Figure 2.79. If you click on an item in the *Parameters* list, you will

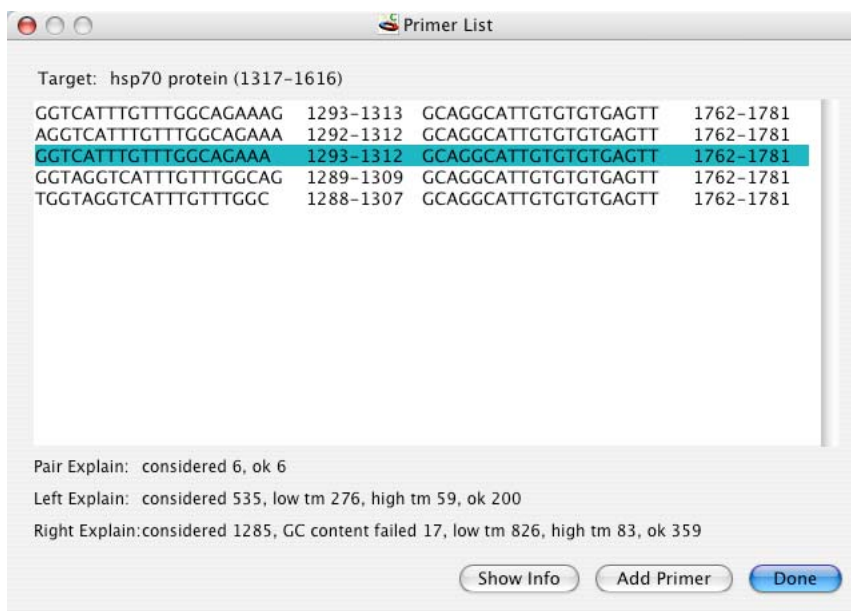


**Figure 2.79: PCR setup panel**

see a description of that item on the right side of the window. Feel free to explore the parameters that you are allowed to set. Those parameters that say “optional” can be left as they are, or you can fill in values for them. For now, set the 5' and 3' allowable ranges to 75 and press the **Generate Primers** button.

5. This will bring up Figure 2.80. There are 5 possible primers that meet the criteria set in the original parameters. The bottom of the window presents some statistics on how all possible primer pairs were filtered to generate the list shown in the window.

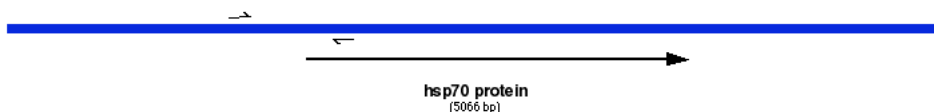
6. Select the middle primer pair (1293-1312, 1762-1781) and press the **Add Primer** button. If you want to, you can add multiple primers to the construct, but for now, just add the one. After you have added the primer, press



**Figure 2.80: Selecting a primer pair**

the **Done** button.

7. You will see Figure 2.81, which shows the two primers. Clicking on either primer will select the primer pair. Once you click on the pair, you can get information by pressing command-I (Mac) or control-I (Windows), or choosing the Construct > Get Info menu item. The results of the Get Info window are



**Figure 2.81: Marked primers**

shown in Figure 2.82. This provides all the details of how the primer was actually generated from the original sequence.

8. You can double click on the primer pair (either of the primers will work) and it will select the DNA amplified by the primer pair. You can now copy this sequence to another window (or another application) to work with it in more

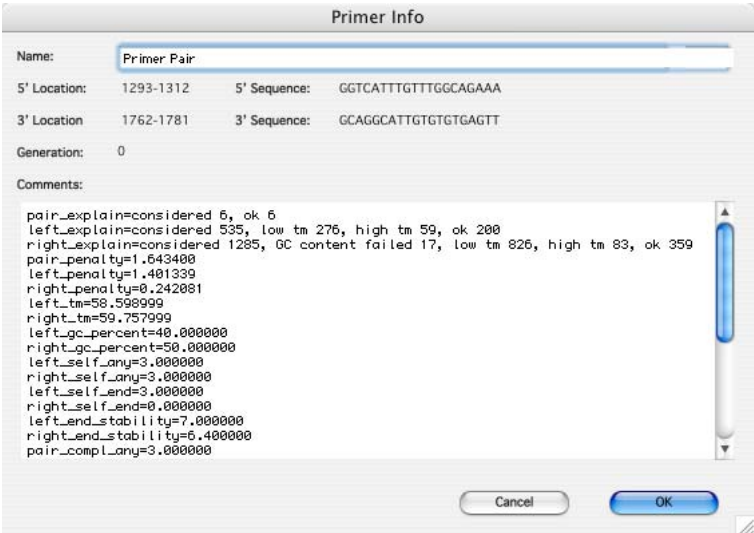


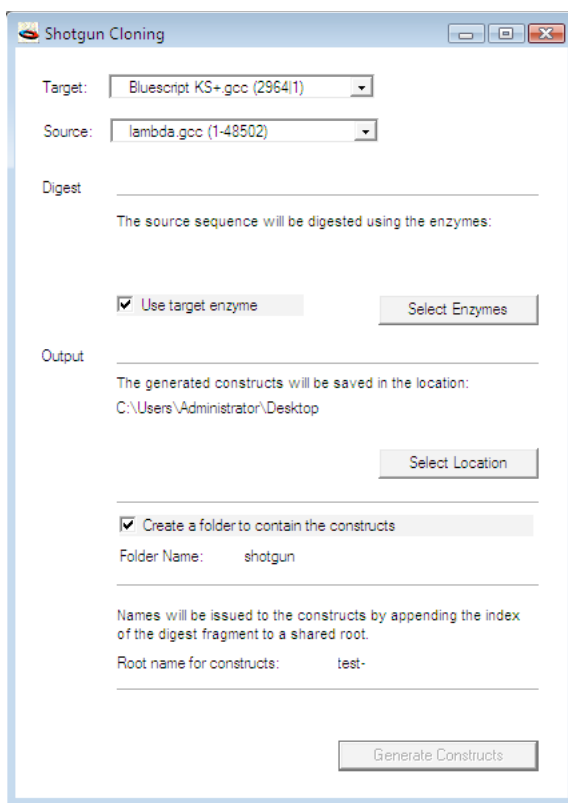
Figure 2.82: Primer information

detail.

## TUTORIAL 17: SHOTGUN CLONING

There is often a need to fragment a piece of DNA with a restriction enzyme and then place each of those fragments into a vector. This process, called shotgun cloning, can be conducted automatically for you by GCK3.

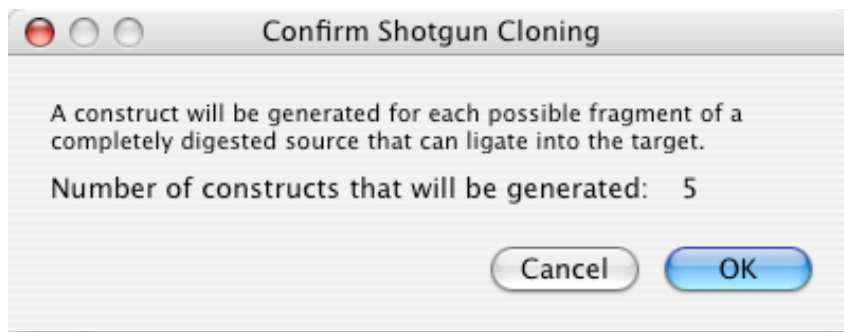
1. Open the file **lambda.gcc** and note that the HindIII (and EcoRI) sites are already marked.



**Figure 2.83: Shotgun cloning setup**

2. Open the construct **pBluescript SK+** and mark the HindIII site by using Construct > Features > Mark Sites. You should end up with a single site marked in the blue-striped region of the DNA.

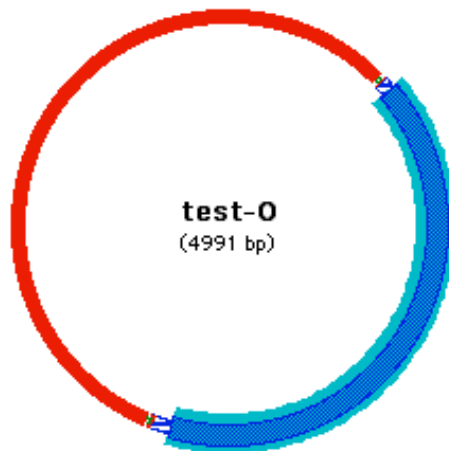
3. Click on the HindIII site marker text in pBluescript SK+ to select it. This also sets the cursor in the DNA to the cut site for HindIII.
4. Choose Tools > Shotgun Cloning.... We will be taking all of the lambda HindIII fragments and cloning them into the HindIII site of the pBluescript SK+ vector. In this case our target site is already selected in the pBluescript SK+ vector and the source of the DNA is lambda, as shown in Figure 2.83. Note that you can choose a different enzyme in which case GCK3 will find all of the sites for you. You can choose a folder for the output (as done here) or you can ask GCK3 to create one for you by checking the checkbox next to *Create a folder to contain the constructs*. You also need to provide a root name for the constructs - in this case the constructs will be named test-0, test-1, test-2 etc. Once your window looks like this figure, press the **Generate Constructs** button.
5. You will see a confirmation dialog Figure 2.84 telling you what is about



**Figure 2.84: Shotgun cloning confirmation dialog**

to happen. This dialog is provided so that if you accidentally are about to generate hundreds of constructs, you will not be surprised by all of the files generated in this process. Press the **OK** button.

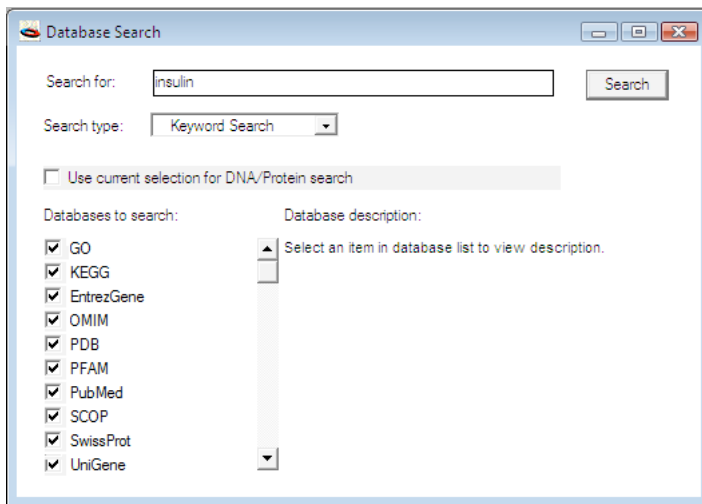
6. There are now 5 new constructs in the folder you designated, each containing a particular fragment of lambda DNA inserted into the HindIII site of the vector. The first such construct is shown in Figure 2.85. Notice that the inserted segment is already highlighted for you.



**Figure 2.85: A “shotgunned” construct**

## TUTORIAL 18: DATABASE SEARCHING

1. GCK facilitates your ability to search multiple web sites for matches with DNA sequences, protein sequences or key words through its database searching tool. To access the DB search, choose Tools > Database Search.... GCK will check with the database server that Textco BioSoftware has set up and will then show you a list of available databases that can be searched (Figure 2.86).



**Figure 2.86: Keyword databases available**

2. Choose Keyword Search from the popup menu. This will show you a list of the currently available web sites that can be searched for key words by GCK. As new sites become available and others go offline, Textco BioSoftware will update the database server to reflect those changes. You can choose which databases to search by using the checkboxes. Leave them all selected in this case.

3. Type in the keyword "insulin" and press the **Search** button. You will be taken to your web browser to see the result. You should see something resembling Figure 2.87. This shows all of the searches that have been initiated. As each search completes, its status will change from **Processing** to **Completed**. If you click on the **Completed** link you can see the results of your

The screenshot displays a web interface for database searching. It is divided into three main sections: Search Parameters, Search Results, and User Information.

**Search Parameters**

Search type:	Keywords
Criteria:	insulin
Search Requested:	2007-12-17 02:16:48
Expiration date:	2007-12-18 02:16:48
Current date:	2007-12-17 02:16:51

**Search Results**

Database Site	Search Status
GO	Processing
KEGG	Processing
OMIM	<a href="#">Completed</a>
PDB	<a href="#">Completed</a>
PubMed	<a href="#">Completed</a>
SCOP	<a href="#">Completed</a>
SwissProt	Processing

*click search status to see results*

**User Information**

Username:	Textco BioSoftware, Inc.
DB Search license expiration:	2007-12-31

**Figure 2.87: Database search in progress**

search displayed in your web browser.

4. Now let's do a protein search. Open the **HSP70 protein** file and select the protein sequence, which is the arrow below the construct - do this by clicking on the arrow once. If you view this as a sequence (Construct > Display > Display Sequence) you will see that it is a single letter representation of the HSP70 protein. Now copy the selection and choose Tools > Database Search. Make sure that you check the checkbox that says *Use current selection for DNA/protein search* and make sure that the text field at the top of the window is empty. Now press the **Search** button to see the results



Note how you have the ability to use whatever you have selected in GCK to initiate the search. This makes it very easy to find out more information about whatever sequence you might be examining.

This concludes the tutorials. For more details about specific features, please use the index to look elsewhere in the index. Enjoy using GCK!

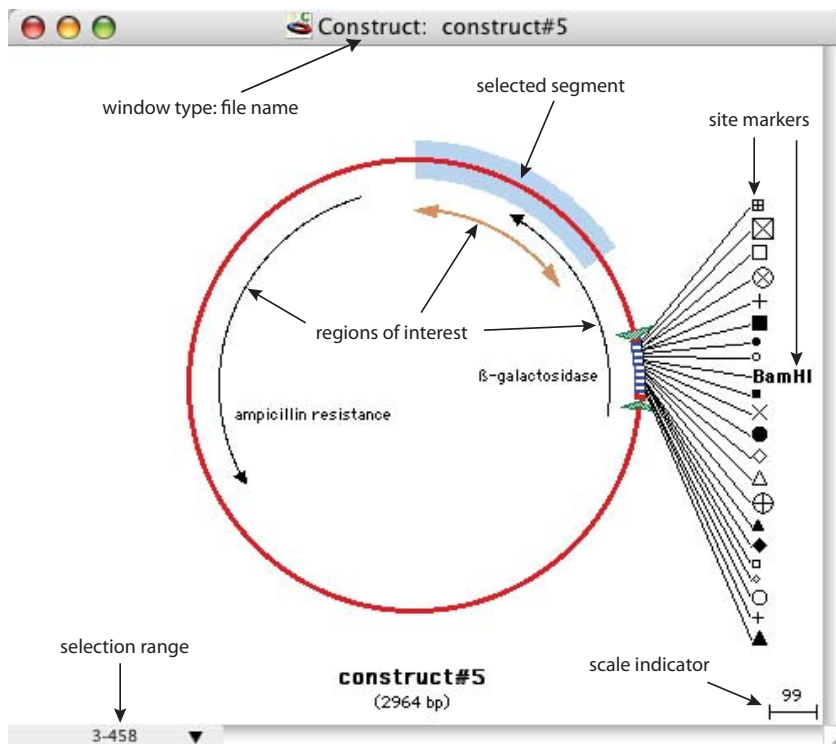


# Chapter 3

## Construct Window

### Overview

The Construct window is the heart of Gene Construction Kit®. The Construct window presents a graphical view of a DNA sequence allowing for direct visualization of sequence features and rapid manipulation of sequence segments.



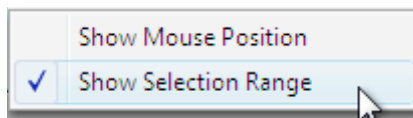
**Figure 3.1: Construct Window (Graphical View)**

It is used for generating figures for publication as well as for direct selection

and manipulation of DNA sequences. A typical Construct window is shown in Figure 3.1, page 3-1.

Many parts of this window are interactive and can be used to bring about changes in the display and to edit the DNA sequence of the construct. The window's title bar shows the type of window (Construct) and the name of the current construct, which also is the name of the corresponding file. **Site markers** can be shown as text or as symbols, **regions of interest** can be shown, and the **segments of DNA** can be defined to have specific appearances. The scale can be shown and set directly or the construct display can be allowed to fit to the size of the window, changing as the window grows and shrinks. The title and size of the construct can also be displayed in the center of the window.

The bottom left corner of the window shows the range of selected nucleotides. Clicking with the mouse on this object will bring up Figure 3.2. By



**Figure 3.2: Mouse Indicator**

using this popup menu, **GCK** can be made to show the position of the mouse as it moves over the construct instead of the range of selected nucleotides.

Parts of a construct can be selected and the appearance of that segment can be adjusted. Once selected a segment can be copied or cut and then pasted into other Construct windows or elsewhere in the same construct.

## **Some Important Definitions**

- **Segment:** A segment in a construct is a range of nucleotides that has a distinct appearance and spans between two borders. In the sequence view, all nucleotides in a segment must have the same font, style, size, and color. In the graphic view, all locations (nucleotides) in a segment must have the same color, fill pattern, and line thickness.
- **Region:** A region (or region of interest) is a feature that corresponds to a

range of nucleotides in the DNA. Regions appear inside circular DNAs or below linear DNAs in the graphical view. In the sequence view, regions always appear below the sequence itself. A region may have a direction because an arrowhead can be placed on one or both ends of the region indicator. Regions with arrowheads at both ends or at neither end do not have a direction. Regions may (optionally) be protein sequences, in which case they *must* have a direction. In the sequence view, “protein” regions are created by translating the corresponding DNA sequence in the direction indicated by the arrowhead. It is possible to have multiple (overlapping) regions corresponding to a given range of nucleotides, and regions do not have to correspond to segments of DNA.

- **Site Marker:** A site marker is a feature attached to the DNA at a specific location between two nucleotides. Site markers are placed outside circular constructs or above linear constructs to avoid overlapping regions. Site markers may represent restriction sites or other sites placed on the DNA by the user.

## **Selecting Segments**

Individual segments of DNA can be selected and modified. The first step to modifying a segment is to select the segment. There are several ways to select a segment. **Dragging** the mouse over the actual sequence in the sequence view of a construct will select characters one at a time just as would occur in a word processor. In the graphic view of a construct, dragging the mouse will select between borders on the construct display. The selection will actually extend from one border to the next as the mouse passes over the midway point between the two borders. **Double-clicking** in the DNA will select the segment that contains the location being double-clicked. Double-clicking will work in the graphical view as well as in the sequence view. Finally, if a **region** is shown, double-clicking on the region will select the corresponding DNA. This is a convenient way to select polycloning sites, coding regions, or origins of replication.

## **Changing the Construct Display**

The appearance of a selected region of the construct can be changed by using the choices under the Format menu. The items that are enabled under

the Format menu depend on what is actually selected in the Construct window. For example, if you have some nucleotides selected in the sequence view, the line thickness menu will not be available; similarly, if you have a graphical segment selected, the Font menu will not be available. Therefore, depending on what you have selected, the availability of items in the Format menu might change.

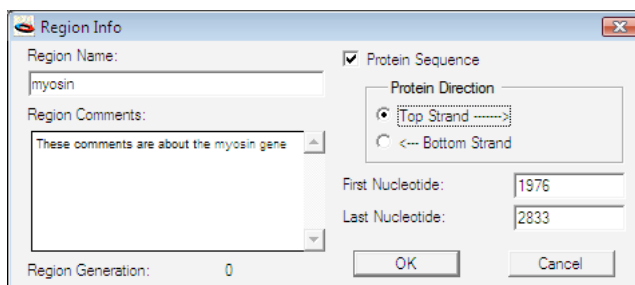
The Format menu options allow the line thickness (Format > Lines >, page 7-19), fill pattern (Format > Fill >, page 7-18), and color (Format > Color >, page 7-22) of the selected segment(s) to be altered. The Format > Lines > menu also allows placing an arrowhead at one or both ends of the segment, thus adding directionality to the segment itself. If the segment has one or two arrowheads, the appearance of the arrowhead can also be adjusted (Format > Lines > Size Arrowhead...). This is discussed in more detail in "Size Arrowhead...", page 7-19. Pre-defined line thicknesses can be chosen, or you can define your own line thickness. *Also see Tutorial 5: "Modifying the Construct Appearance", page 2-20.*

There are many features in Gene Construction Kit many kinds of information that can be displayed in the Construct Window. In order to prevent information overload, you have the option of customizing the display by turning on and off different elements. These options are available under the Format > Display menu (see page 7-58 for more discussion on these menu options). Show Double/Single Stranded, Show/Hide Positions (nucleotide and amino acid position numbers), Show/Hide Breaks (forced line breaks), and Set Line Spacing... (extra spacing between sequence lines) are available only when the construct is viewed as a sequence. Show/Hide Sites will either show or hide all the sites you have marked. Show/Hide Regions will either show or hide the Regions of Interest you have created. Show/Hide Comments will indicate (or not) those regions of the construct to which comments have been attached. Show/Hide Scale Legend will either show or hide the scale legend. Show/Hide Generations will either show or hide brackets that indicate any generation being displayed other than the current generation (see "Chronography - Tracking Construct History", page 3-32). Using the Site Markers submenu provides the ability to change any one or more marked site(s) to be shown as either text or symbol and to define how

the line from the construct will be attached to the site marker.

## **Regions of Interest and Translating DNA**

**Regions of interest** may be used to represent coding portions of the DNA, exons/introns, promoters, origins of replication, etc. A region of interest can be created by selecting the part of the construct that will correspond to the region and then choosing Construct > Features > Make Region... (command-R/ctrl-R). If no segment of DNA is selected and Make Region... is chosen, you can simply type in the first and last nucleotides to be included in the region. The create



**Figure 3.3: Creating a Region**

region dialog box is shown in Figure 3.3. You can provide a name for the region and can enter any comments that you would like to keep stored with the region. If the corresponding segment of DNA is copied and pasted into another construct, all the comments associated with the region will be carried to the new construct as well. By placing an “x” in the checkbox **Protein Sequence**, the region could also be defined as a protein region. In this case, the corresponding DNA will be translated and the protein sequence shown whenever the construct is viewed in a sequence view of the Construct window. This was illustrated in Tutorial 3: “Marking Open Reading Frames”, page 2-11.

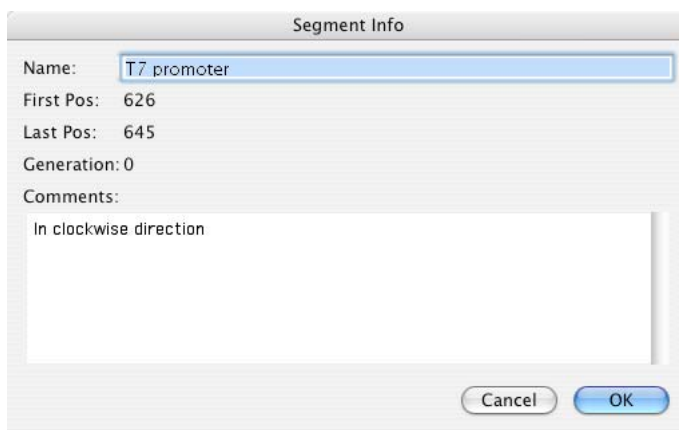
Overlapping regions can be defined by choosing appropriate segments on the DNA that correspond to the desired region and then selecting Construct > Features > Make Region.... Once a region is formed, it can be selected by clicking once with the mouse. The appearance of a selected region can be altered using many items listed in the Format menu, just as can be done for segments (see “Changing the Construct Display”, page 3-3).

The dialog box shown in Figure 3.3 can be viewed by selecting a region and then choosing Construct > Get Info... (command-I/ctrl-I). You can type in new values in the **First Nucleotide** and **Last Nucleotide** boxes to adjust the end points of the region. You may type up to 32,000 characters of comments into each Get Info dialog box. These comments can be searched for key words using Construct > Search Comments... (page 7-69) or by using File > Search Files... (page 7-35). See Tutorial 8: “Finding Comments and File Searching”, page 2-38) for a hands-on example of searching for comments. Also see “Searching Comments”, page 3-22.

Regions can also be rearranged by moving them to the outside or the inside (see “Move Region Outwards/Upwards”, page 7-27). If a region is selected and “moved to the top” GCK will move the selected region to the top of the stack of regions being displayed – it will never move the region to the outside of a circular construct or above a linear construct<sup>a</sup>.

## **Attaching Comments to Objects in Construct Window**

Comments may be attached to any segment of the construct which is selected by using Construct > Get Info.... These comments may be used to store



**Figure 3.4: A Segment Get Info... Dialog**

notes about where the particular piece of DNA came from, the functionality of the sequence, or any other information you wish to have associated with the

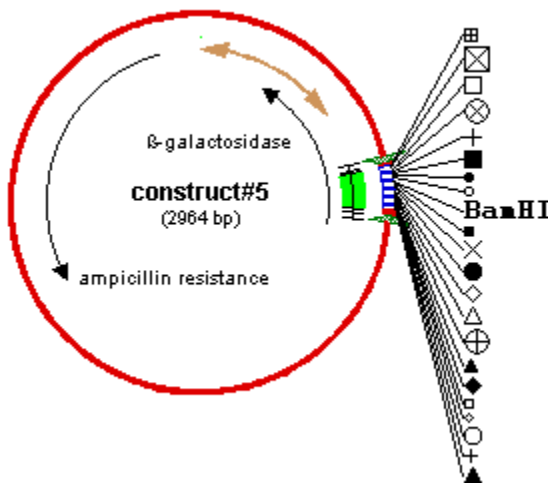
---

a. This prevents regions from unintentionally overwriting site markers.



particular DNA segment (see Figure 3.4, page 3-6). Comments can be reviewed and edited at any time using Construct > Get Info..., and can be used as an electronic notebook to document the various steps used in generating the construct. When a segment is pasted from one location to another (even into a different construct), the comments are pasted along with the DNA sequence itself.

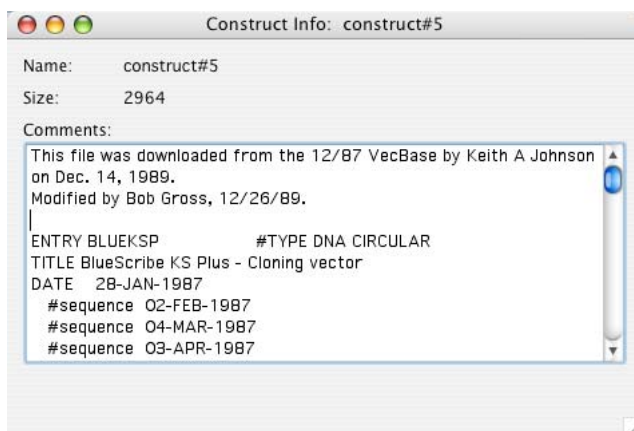
Because it is not obvious from looking directly at a construct which parts of the construct have comments associated with them, **GCK** can display an indicator of the location of comments by choosing Construct > Display > Show Comments. This will produce a display like Figure 3.5, page 3-7. The selected object is a comment indicator and indicates that there are comments associated with that particular segment of DNA. You can view the comments by clicking once on the segment indicator to select it and then choosing Construct > Get Info.... Alternatively, you can double-click on the comment indicator to



**Figure 3.5: A Comment Indicator**

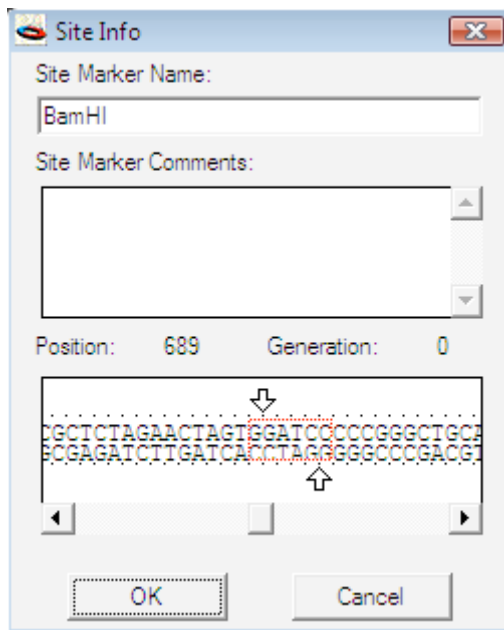
select the corresponding segment of DNA and then view the comments by choosing Construct > Get Info.... In operational terms, these comment indicators behave very similarly to regions. However, comment indicators and regions can be shown and hidden separately, regions do not have to have comments, and regions can represent protein sequences.

Construct > General Info... is another kind of information that is attached to a construct. A dialog showing general information is shown in Figure 3.6, page 3-8. This information is actually attached to the file (whole construct) and not an individual object in the Construct Window, as are all of the other Get Info... comments. General Info may be used to store literature references or information about the location of the construct in the lab (e.g. freezer in 318, shelf 4). If you select the entire construct, copy it and paste it into an empty Construct Window, the General Info will be preserved as General Info. On the other hand, if you paste into an existing construct, the General Info will be converted to information attached to the resulting segment.



**Figure 3.6: General Info Dialog**

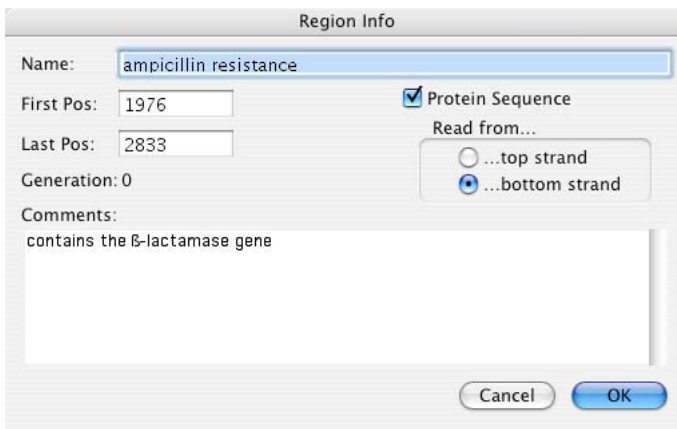
Comments can be associated with different objects in the Construct Window. The comments for any object can be viewed by selecting the object and then choosing Construct > Get Info... (command-I/ctrl-I). Each kind of object will show a slightly different *Get Info...* box. A typical site marker Get Info... box is shown in Figure 3.7. In this Get Info box, you can view the recognition sequence (dotted line in sequence) and actually alter the cutting site for the enzyme. The cut site can be changed by dragging the arrows adjacent to the sequence or by clicking the mouse at the new location to which you want the triangle to move. Be careful when you do this, however, since the site name will remain the same, but the cut site will be different from the standard site. If you were to copy or cut the fragment delineated by the altered site and paste it into a new construct, the end of the new fragment would not in reality



**Figure 3.7: Site Marker Get Info... Dialog**

be a correct end. If you alter the cut site for a restriction enzyme, it is best to change the name of the site also.

Regions also have Get Info... boxes as shown in Figure 3.8. This dialog con-



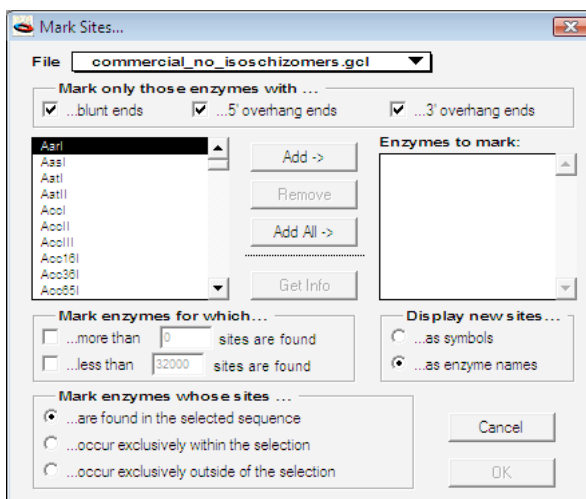
**Figure 3.8: Region Get Info... Dialog**

tains information about the selected region. You can change the direction of

the region by clicking the **Top Strand** or **Bottom Strand** radio buttons. Changing the numbers in the **First** or **Last Nucleotide** box will change the length of the region. The **Protein Sequence** check box determines if the region is to be translated or not when viewed as a sequence

## Marking Sites

Sites can be marked along a construct by using Construct > Features > Mark Sites... (comment-M/ctrl-M). This will present you with Figure 3.9. Using this dialog



**Figure 3.9: Mark Sites Dialog**

box, you can mark sites in the construct using enzymes from many different lists, and define your own parameters of the kinds and numbers of sites to actually be marked. Here are the options in this dialog:

- The **popup menu** at the top of the dialog allows you to specify which enzymes will be displayed in the left side list in this dialog. The enzyme lists that are in the **GCK data** folder are the only ones that will show up in the popup menu<sup>b</sup>.
- The **Mark only those enzymes with...** cluster contains three check boxes that allow you to show any combination of **...blunt ends**, **...5' overhang ends**, and/or **...3' overhang ends**. Check all three of these

b. Note that you can always download the most recent lists from Textco's web site <<http://www.textco.com/>>. The lists are updated quarterly.

boxes to see all of the sites. If you do not check any of the boxes, no sites will be shown.

- Clicking on an enzyme in the left list and then pressing the **Add->** button will move that enzyme to the *Enzymes to mark:* list on the right. All enzymes in the right list will be used to search the targeted DNA for corresponding sites. The **Add All->** button will add all of the enzymes in the left list to the right list of enzymes to mark. If an enzyme is selected in the right list, pressing the **Remove** button will remove that item from the Enzymes to mark list (the enzyme will still remain in the restriction enzyme file for use in other digests – this item just removes it from the list of sites to search for.).
- The **Mark enzymes for which...** cluster allows you to specify a range of cutting frequencies for which you would like to see sites actually marked. If you leave these fields blank all sites are shown. The top field, **More than 'n' sites are found**, lets you define the minimum number of times an enzyme recognition site must be present in the target DNA in order for that enzyme to be marked. To activate a minimum number, make sure that the check box at the left of the text has an 'x' in it. The bottom field, **Less than 'n' sites are found**, lets you define the maximum number of times an enzyme recognition site must be present in the target DNA in order for that enzyme to be marked. By checking both check boxes, you can specify a range of cut frequencies to be found. For example, if you want to see all enzymes that cut between 2 and 5 times, you would put a "1" in the top box and a "6" in the bottom box.
- The **Display new sites...** cluster allows you to specify how you would like to mark the new sites – as symbols or as text. Choose one of the radio buttons to make your choice.
- The **Mark enzymes whose sites...** cluster allows you to define which parts of the construct are to be used as the target for the searching. The only time the radio buttons are active in this cluster is when you actually have a selection in the DNA – otherwise the choices do not make any sense. If no DNA is selected the search will take place on the entire construct DNA. Choosing **...are found in the selected sequence** will mark only those enzymes that have a recognition sequence within the selected DNA. These enzymes might also have sites outside of the selected segment. Choosing **...are found exclusively within the selection** will mark only those enzymes that have recognition sequences within the selected

DNA and have no other sites elsewhere in the construct. Choosing **...are found exclusively outside the selection** will mark only those enzymes that have no sites within the selection but do have sites outside the selection. Using this cluster of choices together with the “Mark enzymes for which...” cluster allow you to find exactly the right enzymes for the task you want to accomplish.

## **Site Markers**

If you choose to mark sites as symbols, **GCK** will choose symbols from the symbol table (see “Symbol”, page 7-22) and will mark all sites for a given enzyme with the same symbol. If there are more enzymes chosen than there are symbols in the table, more than one enzyme will end up having the same symbol. Note also, as a result of how the symbols are chosen, if you mark sites for a specific enzyme at different times, the site marker symbols might end up different<sup>c</sup>. If you want to be sure that all the sites for a given enzyme have the same symbol, you should: (1) first make sure that no sites are marked for that enzyme, and then (2) mark sites for the particular enzyme using symbols (i.e. – there will only be one enzyme in the right list in Figure 3.9, page 3-10 and **mark the sites as symbols** should be chosen).

Once sites are marked on a construct you can select any site by clicking on it once. If you **double-click on a site marker** **GCK** will select all other sites that are marked in the same way. For example if you double-click on a site marked *EcoRI*, all other sites labeled exactly as “*EcoRI*” will be selected – no sites marked as symbols will be selected and no sites marked as text having different text will be selected. On the other hand, if you double-click on a site marked as ♦, all sites having that exact symbol in that exact size will be selected – sites with different symbols, the same symbol in a different size, or sites marked as text will not be selected.

Once a site is marked, changing the label display can be accomplished using the **Format > Site Markers submenu**. **Format > Site Markers > Show Site As Text** will force

- 
- c. The only way for **GCK** to avoid this inconsistency would be to define a different symbol for each enzyme in the list and then maintain that symbol as the list grows and enzyme names change. This would be an impractical task.

any selected site markers to be displayed as text. `Format > Site Markers > Show Site As Symbol` will force any selected site markers to be displayed as symbol. Other items under the `Format > Site Markers` submenu can be used to define what information is to be shown (if the site marker is shown as *text*) and how the site marker is connected to the construct (see “Site Markers”, page 7-23 for a complete discussion).

Site markers may be quite crowded together under some circumstances. To spread out the site markers, select the site markers you wish to spread out and then choose `Format > Site Markers > Automatic Arrangement`. This will spread out the selected markers and will designate them to remain spread out even if you resize the window. To prevent them from automatically rearranging themselves, you can either move the site by hand or choose `Format > Site Markers > Manual Arrangement`.

Moving site markers manually can be accomplished by clicking on a site marker to select it and then dragging with the mouse. You can also move a site marker one pixel at a time with the arrow keys. To change a property for a group of site markers (e.g. to make them all red or Times) you can use the mouse to drag a selection rectangle over the sequences you are interested in selecting. This makes it easy to select a set of markers, such as those at a polycloning site, and set them to auto-arrange while leaving the rest of the (non-crowded) sites to be arranged manually.

Other choices under the `Format` menu will alter the appearance of the selected site marker. `Font`, `Color`, `Style`, and `Size` of the site marker text can be changed using the `Format` Menu. In addition, if the site is being shown as a symbol, the symbol can be changed using `Format > Symbol`.

When a restriction enzyme site is marked on a sequence, convention usually places the first letter of the enzyme name above the first letter of the recognition sequence. In the sequence view of a construct, when an enzyme site is chosen, you will see an indication of the actual cut site as a series of “marching ants” as shown in Figure 3.10. In this case the *Sall* site is GTCGAC so the “S” of *Sall* lines up above the first G in the recognition sequence. The cut is clearly indicated as occurring between the G | T.

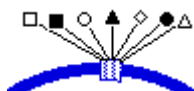
```

HindIII  PstI  Sall  BamHI SmaI  SstI  EcoRI
GATACAGCTTGGGCTGCAGGTCGACTCTAGAGGATCCCGGGCGAGCTCGAAT
CTTATGTTTCGACCCGACGTCCAGCTAGATCTCCTAGGGGCCGCTCGAGCTTA

```

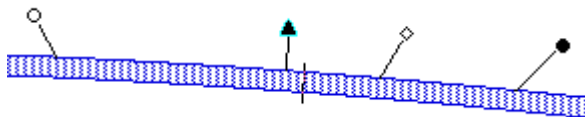
**Figure 3.10: Selected Site Marker**

Because GCK relies on being able to cut DNAs at the actual cut site and not



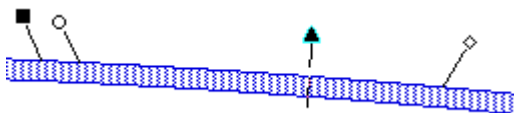
**Figure 3.11: Site Markers at Magnification of 1x**

the start of the recognition site but displays the markers by default at the recognition site, sometimes confusing graphics can result. For example, Figure 3.11 shows the same piece of sequence (a polycloning site from the vector pGEM-1), but viewed graphically with the sites marked as symbols rather than as text. At the 1x magnification shown in Figure 3.11 everything looks



**Figure 3.12: Markers at Recognition Sequence**

normal. However, if the view is magnified (see “Magnification and Construct Scale”, page 3-44) to 3 nts/cm, it appears that the insertion point in the actual construct does not agree with the selected site marker as shown in Figure 3.12, page 3-14. In fact, the drawing is correct in that the symbol is placed above the first nucleotide of the recognition sequence, while the insertion point in the DNA is at the actual cut site.



**Figure 3.13: Markers at Cut Site**

Using the Construct > Features > Place Site Indicators at... provides you an option to specify exactly how markers will be placed along a construct, as shown in Figure 3.13. Using the dialog that appears, you can specify that the site



maker be placed either the beginning of the recognition site for the enzyme, or just to the left of the actual cut site. In some extreme magnifications, these options might actually change the order of some closely spaced restriction site markers.

In addition to marking sites using Construct > Features > Mark Sites..., specific positions on the construct can be labeled by using the Construct > Features > Mark Location.... Marking a location allows you to specify a precise location (e.g. nucleotide 2377) at which a marker will be placed. Since you can define the name for the marker, this is an ideal way to label specific locations on the DNA. The details of Mark Location are discussed in “Mark Location...”, page 7-51.

Construct > Features > Place Sites... allows you to place specific restriction enzyme sites at specific locations. This is very useful for working with generic constructs, whose restriction maps are known, but whose sequence is incomplete or missing entirely. This was discussed in Tutorial 11: “Working With Generic Constructs”, page 2-56 and is also detailed in “Place Sites...”, page 7-52.

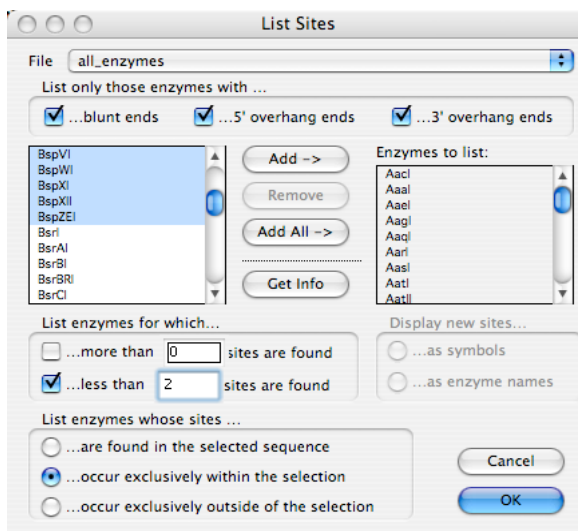
## **Listing Sites**

The setup for this is almost identical to the setup for Marking Sites (“Marking Sites”, page 3-10). Choosing Construct > Features > List Sites... (command-L/ctrl-L) will bring up Figure 3.14. Using this dialog box, you can list sites in the construct using enzymes from many different lists, and define your own parameters of the kinds and numbers of sites to actually be marked. Here are the options in this dialog:

- The **popup menu** at the top of the dialog allows you to specify which enzymes will be displayed in the left side list in this dialog. The enzyme lists that are in the **GCK data** folder are the only ones that will show up in the popup menu<sup>d</sup>.
- The **List only those enzymes with...** cluster contains three check boxes that allow you to show any combination of **...blunt ends**, **...5' overhang ends**, and/or **...3' overhang ends**. Check all three of these

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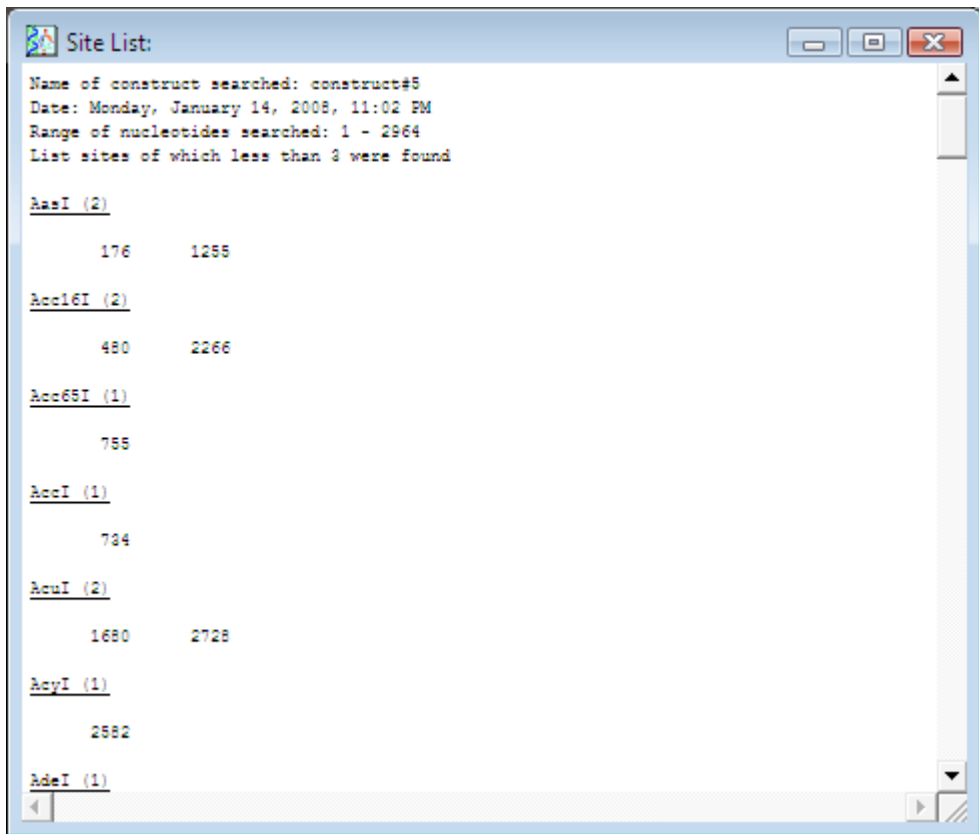
d. Note that you can always download the most recent lists from Textco's web site <<http://www.textco.com/>>. The lists are updated quarterly.



**Figure 3.14: List Sites Dialog**

boxes to see all of the sites. If you do not check any of the boxes, not sites will be shown.

- Clicking on an enzyme in the left list and then pressing the **Add->** button will move that enzyme to *Enzymes to list:* on the right. All enzymes in the right list will be used to search the targeted DNA for corresponding sites. The **Add All->** button will add all of the enzymes in the left list to the right list. If an enzyme is selected in the right list, pressing the **Remove** button will remove that item from *Enzymes to list:* The enzyme will still remain in the restriction enzyme file for use in other digests – this item just removes it from the list of sites to search for.
- The **List enzymes for which...** cluster allows you to specify a range of cutting frequencies for which you would like to see sites actually listed. If you leave these fields blank all sites are shown. The top field, **More than 'n' sites are found**, lets you define the minimum number of times an enzyme recognition site must be present in the target DNA in order for that enzyme to be marked. To activate a minimum number, make sure that the check box at the left of the text has an 'x' in it. The bottom field, **Less than 'n' sites are found**, lets you define the maximum number of times an enzyme recognition site must be present in the target DNA in order for that enzyme to be marked. By checking both checkboxes, you can specify a range of cut frequencies to be found. For example, if you



**Figure 3.15: List Sites Output**

want to see all enzymes that cut only once (“unique cutters”), you would put a “0” in the top box and a “2” in the bottom box.

- The **List enzymes whose sites...** cluster allows you to define which parts of the construct are to be used as the target for the searching. The only time the radio buttons are active in this cluster is when you actually have a selection in the DNA – otherwise the choices do not make any sense. If no DNA is selected the search will take place on the entire construct DNA. Choosing **...are found in the selected sequence** will list only those enzymes that have a recognition sequence within the selected DNA. These enzymes might also have sites outside of the selected segment. Choosing **...are found exclusively within the selection** will list only those enzymes that have recognition sequences within the selected DNA and have no other sites elsewhere in the construct. Choosing **...are**

**found exclusively outside the selection** will list only those enzymes that have no sites within the selection but do have sites outside the selection. Using this cluster of choices together with the “List enzymes for which...” cluster allow you to find exactly the right enzymes for the task you want to accomplish.

A typical output from the List Sites analysis is shown in Figure 3.15, page 3-17. The output has a header which names the sequence and the date on which the analysis was conducted. If there were enzymes chosen that do not have any sites in the DNA, they will be listed at the end of the output.

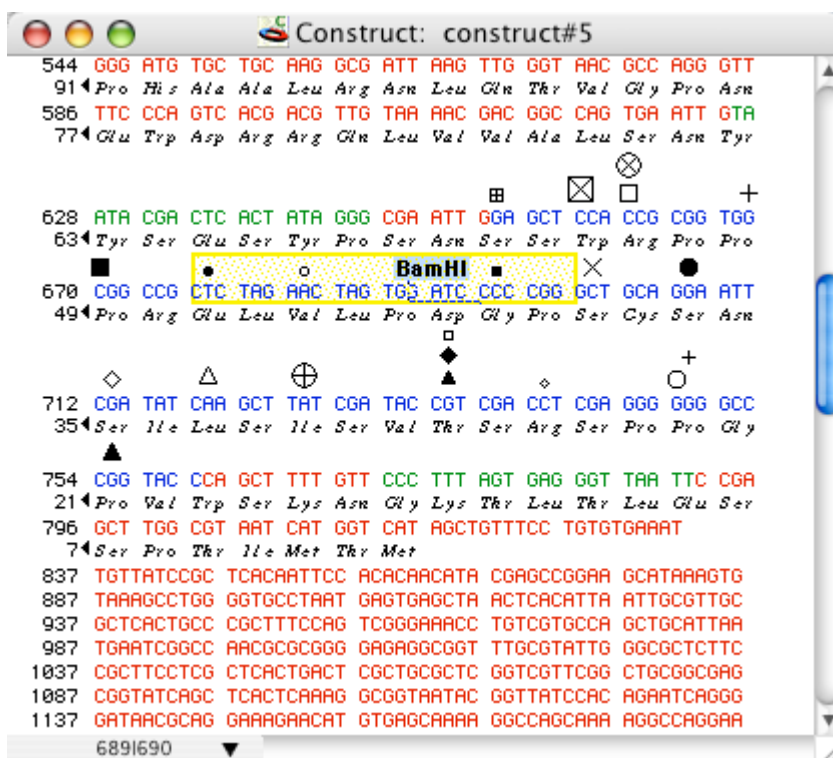
## **Viewing the Construct as a Sequence**

Any construct can be viewed and formatted as a sequence. Most of the menu options under the Construct menu remain active, a few new ones appear, and some become inactive. To view the construct as a sequence select Construct > Display > Display Sequence. The graphical view of the construct will be replaced by a view of the sequence itself. The menu item you just selected will change to Construct > Display > Display Graphics.

The sequence view of the construct will display the actual sequence along with other attributes you have selected. Show/Hide Generations, Show/Hide Regions, Show/Hide Sites, and Show/Hide Comments are still available. These choices function the same in the sequence view as in the graphics view of the construct. Sites will be displayed with the first letter of the site name aligned to the start of the recognition sequence in the DNA. If the region being displayed has been defined as a protein region, it will be displayed as an amino acid sequence below the DNA sequence.

## **Formatting A Sequence Listing**

As described in Tutorial 4: “Viewing the Construct as a Sequence”, page 2-15, any GCK construct can be viewed and edited as a sequence. The sequence display of the sequence can be modified in a number of ways, enabling you to produce a formatted output suitable for publication or for slides or posters. A typical sequence view is shown in Figure 3.16. This view



**Figure 3.16: Sequence View of Construct**

illustrates a number of features.

The nucleotide and amino acid positions are indicated at the start of each line in the sequence view of the construct. These position indicators can be hidden by choosing Construct > Display > Hide Positions. The numbers shown reflect the position for the character at the left end of each line – the A at the top of the window is nucleotide 637, the 70 just below the 637 is the 70<sup>th</sup> amino acid, Ser. The triangles at the end of the amino acid sequence indicate that the coding region is running from right to left and therefore corresponds to the “bottom” DNA strand (not shown). Choosing Construct > Display > Show Double Stranded will show the “second” DNA strand in addition to the one currently being shown.

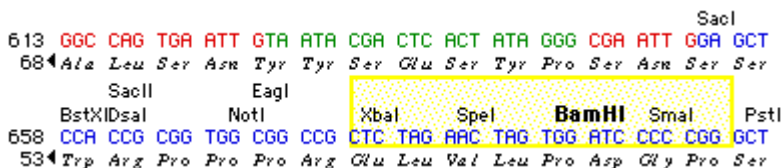
You can modify the output format by choosing Format > Grouping > (see also “Grouping”, page 7-29). To change the grouping of a segment of DNA, start

by selecting a segment of DNA and then choose an item under Format > Grouping > to define the number of nucleotides to be placed between each space. This option is particularly useful for displaying coding regions of DNA in groups of three while other regions of DNA might be in groups of ten (as shown in Figure 3.16) or not grouped. If your particular sequence has an octamer motif, it might be best to display it in groups of eight.

Also available for formatting is Format > Grouping > Insert Line Break. This item provides the ability to insert a line break anyplace in the sequence and therefore gives you complete flexibility to format the sequence in precisely the way you desire. In addition, the Construct > Display > Set Line Spacing... item allows you to define the minimum number of pixels between each line of sequence.

The last sequence formatting option is Construct > Display > Show Line Borders. This option will display a dashed line between each set of lines representing a DNA sequence and its associated marked sites and regions. It is useful in helping you visualize the information more consistently and can clearly indicate which amino acid sequence is related to which line of DNA sequence.

**Frames** are discussed in Tutorial 5: “Modifying the Construct Appearance”, page 2-20. Frames provide a way to enclose part of a sequence display to



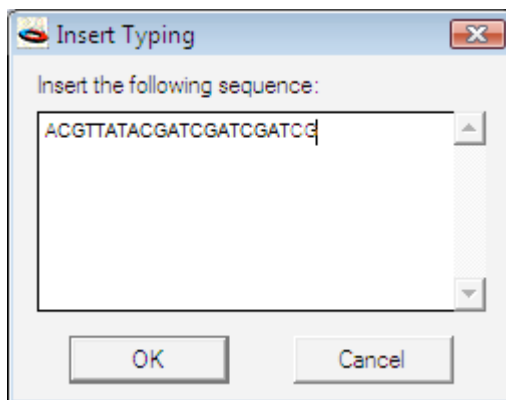
**Figure 3.17: A Framed Segment**

highlight a specific feature. A Framed segment is shown in Figure 3.17, page 3-20. To create a frame around a sequence, first select the sequence you wish to frame and then choose Construct > Features > Make Frame.... A new frame will appear and will be selected as shown in Figure 2.19, page 2-20. When a frame is selected you can alter its appearance by choosing various items under the Format menu. A frame can be made to include the DNA sequence, marked sites and regions or any combination of these items by choosing Construct > Features > Set Frame Contents... (see Figure 2.20, page 2-21).

Once a frame has been deselected, you might need to select it again to modify its properties. Just dragging the mouse in the DNA will select nucleotides along the DNA (as it should) and will not select the frame. How then, can you select a frame? A frame can be selected by option-clicking (holding down the option key and clicking) on the frame or by choosing **Construct > SelectFrame** and then clicking on the frame you want to select. For those of you who are “quick-fingered” you can also select a frame by triple-clicking on the DNA contained within a frame. A selected frame will have its border highlighted (Figure 2.19, page 2-20).

### **Inserting New Nucleotides by Typing**

Because of the complexity of the sequence display window, it would be unacceptably slow to allow typing in new nucleotides while the program updates the screen including all the graphic elements in the sequence view after each nucleotide is typed. Therefore, the Gene Construction Kit allows you to place the cursor in the DNA sequence at the site you wish to add new nucleotides and then start typing, as you normally would. Instead of entering each letter into the sequence display window as it is typed, the program opens up a dialog box (Figure 3.18, page 3-21) into which you type your sequence. Each



**Figure 3.18: Insert Nucleotides**

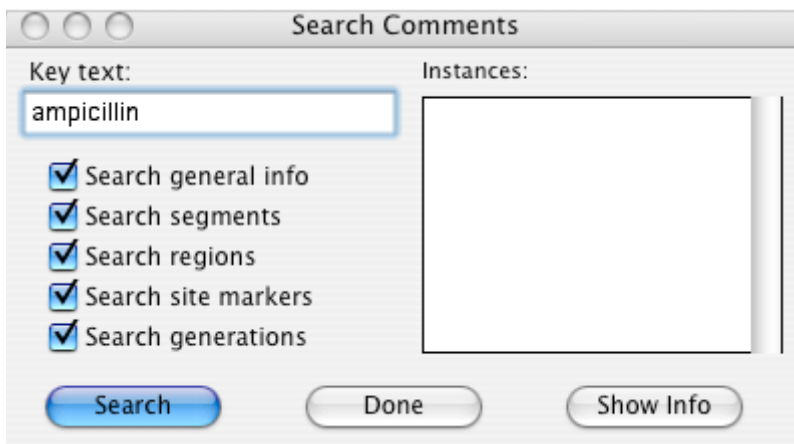
letter is checked as it is entered to ensure that it is a valid nucleotide character (see “Legal Nucleotide Characters”, page 8-1), and then placed into the box. Once you have typed in the desired sequence, clicking the **OK** button will place the new nucleotides into the sequence at the location of the cursor.

Pressing the **Cancel** button will leave the sequence unchanged. You can also bring up this dialog by choosing Construct > Insert Typing....

Nucleotides can be deleted in the sequence view by first selecting the characters to be deleted and then Cutting them. It is not possible to delete nucleotides by backspacing (delete key) because of potential ambiguous situations<sup>e</sup>.

## **Searching Comments**

All the comments associated with constructs can be searched for key words<sup>f</sup>. Comments (and names) associated with regions of interest, marked sites, general information, and with DNA segments can all be searched. This is



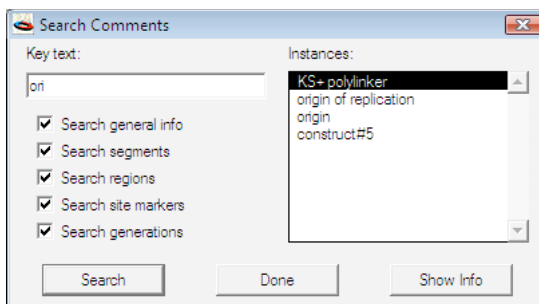
**Figure 3.19: Search Comments Dialog**

accomplished using Construct > Search Comments.... You will be presented with a

- 
- e. For example, it is not clear if **GCK** should stagger a deletion on each strand at an actual cut site. Also, if a site marker is selected, the insertion point will reside at the actual cut site while the site marker remains selected. When the delete key is pressed in this situation, it is not clear if the site marker should be deleted or if the nucleotide to the left of the insertion point should be deleted. Not enabling the delete key eliminates these potential ambiguous areas.
- f. Searching comments was the subject of Tutorial 8: "Finding Comments and File Searching", page 2-38 – reviewing this tutorial will provide a hands-on perspective for getting the most out of **GCK** comments.



dialog box like the one shown in Figure 3.19. If no segment is selected, the search will apply to the entire construct. If a segment is selected, however, the search will be confined to only the selected segment of the DNA and all of its associated sites, segments, and regions in all generations. To initiate the search, type in the text you would like to search for in the **Key Text** box, define which parts of the construct you want to search (using the check boxes along the left), and press the **Search** button. In Figure 3.20, the word



**Figure 3.20: Search Results**

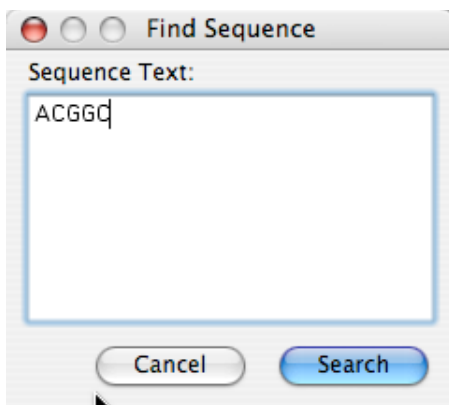
“ori” was used and matches were found in four instances as listed in the right of the dialog box. Double-clicking on the item in the *Instances* list, or selecting the item and then pressing the **Show Info** button will open the Get Info box for that particular match.

Searches are conducted on all generations of the construct and therefore can be used to search through all your notes about a construct. By carefully documenting each step you perform, this electronic notebook can not only contain the same info as your regular lab notebook, but the notes can be associated with specific features of the construct, can be saved in chronological order by taking advantage of the Chronography feature (“Chronography - Tracking Construct History”, page 3-32), and can be rapidly searched. Using these comments in conjunction with an illustration window to diagram the cloning strategy provides a very reliable way to document your project.

## **Finding a Sequence**

When viewing the construct as a sequence, you can place the cursor anywhere in the construct and then search for the next occurrence of a sequence.

This is accomplished by choosing Construct > Find Sequence (command-F/ctrl-F). You will see the dialog box shown in Figure 3.21. Type in the sequence to



**Figure 3.21: Find Sequence Dialog**

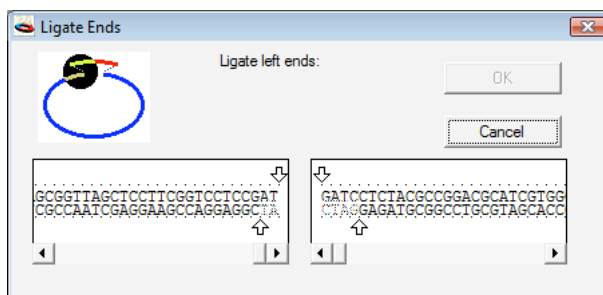
find and press the **Search** button. All valid DNA characters are accepted in the search sequence. **GCK** will search the sequence starting from the cursor and will highlight the next occurrence of that sequence in the sequence listing itself.

## **Cutting, Copying, and Pasting Segments**

You can use the Gene Construction Kit to graphically cut, copy, and paste segments of DNA. Copying (or cutting) a fragment from one construct and pasting it into another is a three step process. First, select the segment by dragging with the mouse or by double-clicking the fragment you wish to move (cut or copy). Once the desired fragment is selected, choose Edit > Cut (command-X/ctrl-X) or Edit > Copy (command-C/ctrl-C) to place that segment in the clipboard. Second, select the location into which you want to insert the fragment. This can be an empty Construct Window or any marked site or end on any other construct, or even a location between two nucleotides in the sequence view. Third, select Edit > Paste (command-V/ctrl-V) to place the segment from the clipboard into the selected site. If you had selected a whole segment in your target construct (rather than just having an insertion point in the target), the selected segment will be replaced by the fragment in the clipboard. This is just like replacing a selected word in a word processing program.

When you copy segments of DNA, all the attached sites, chronography, and regions are copied with the DNA. Note that with regions, however, the entire region must be included within the DNA for it to be copied. Copying a piece of DNA that includes only part of a region will not copy that region.

If you are pasting in a fragment with ends that are not compatible with the insertion site, you will be presented with the ligation dialog box shown in Figure 3.22. The diagram in the upper left of the dialog box tells you which



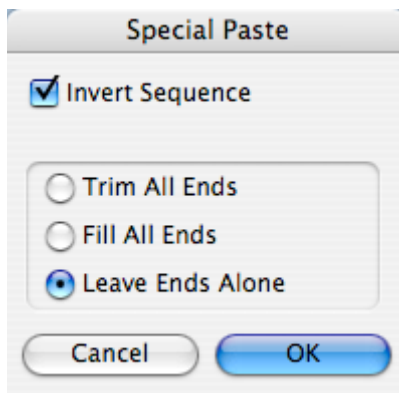
**Figure 3.22: Ligation Dialog**

junction of the ligation you are viewing and will vary depending on whether you are pasting into a linear or circular construct, as shown in the figure. The black circle highlights the actual junction that is being presented in the dialog. The **OK** button cannot be selected until the segment ends are made compatible. This can be done by moving each of the four arrows to the desired locations by dragging with the mouse, or simply by clicking at the desired new location for the cut site. After adjusting the left junction for the insertion, you will be presented with a similar dialog box for the right junction for the insertion if necessary. In the current figure, the left end of the segment to be inserted (the sequence shown on the right of the dialog box) is blunt ended, while the sequence shown on the left of the dialog represents the left side of the opening in the construct which is receiving the segment. This end has a staggered cut. Dragging the top arrow in the left side sequence to the end of the sequence (to the right of the TT) will create a blunt end and allow the ligation to occur.

Note that if you cut a segment out of a construct and that segment has different sites on each end, you will also get a ligation dialog, since the program

needs to know how the two resulting ends should be joined together. This feature forces you to accurately keep track of segment ends and maintain an accurate sequence – just as you would have to do in the lab to recircularize the construct after removing a segment.

You can also paste fragments using the Edit > Special Paste.... The Special Paste dialog is shown in Figure 3.23. **Invert Sequence** will invert the sequence



**Figure 3.23: Special Paste**

before pasting it into the new site. **Trim All Ends** will trim any single stranded tails from all four DNA ends before pasting (just like treating with S1 nuclease). **Fill All Ends** will fill in all the complementary nucleotides on all four ends before pasting (just like filling the ends with DNA polymerase, but be careful not to fill the ends in the 3' to 5' direction unless you have figured out a way to do this biologically as well). **Leave Ends Alone** will present you with a dialog box if the ends are incompatible, as in the normal pasting of a segment. **Leave Ends Alone** in the absence of inversion is equivalent to a normal pasting operation.

Sometimes you will notice that when two segments are joined together, a small marker consisting of a white “j” in a black circle is placed at the junction of the two segments. This marker is called a **junction marker** and is placed at the location of the joining of the two fragments to preserve the boundary between the two joined segments, as shown in Figure 3.24.

Each time that two segments are joined together, the program must decide



**Figure 3.24: Junction Marker**

how to indicate the junction between the two segments. This is important for archival purposes as well as to provide you with the ability to manipulate these segments in the future. If the ends that are being joined are the results of restriction enzyme cuts and the recognition site is regenerated during the ligation, the restriction enzyme marker(s) will be regenerated. If, however, the ends are adjusted with the ligation dialog box and the original recognition sequences altered, the original sites may no longer be appropriate. In this case the program will generate a junction marker at the boundary to retain the information about the boundary. Likewise, if blunt ends are joined together (no ligation dialog will be shown) the program will create a junction marker.

There are two other situations in which a junction marker will be created. First, if a segment is removed from a construct by selecting and then cutting, the two new ends that are left will have to be joined together using the rules discussed above. Second, a junction marker will be created when a linear construct is converted to a circular construct. This is just a special case of joining two ends together.

Obviously, if you do not have a need for a junction marker, it can be deleted (select it and press the delete key). It is put in to preserve the junction information in case you do need it. The philosophy is that it is better to provide the information and let you dispose of it, than not to provide it at all.

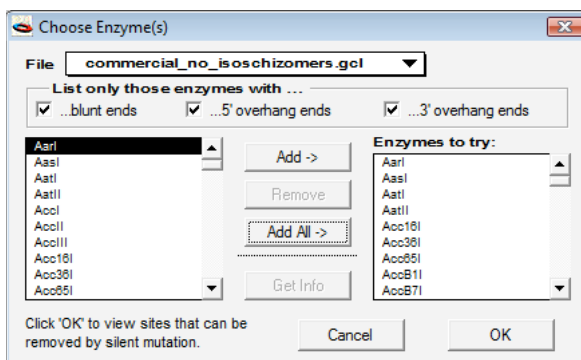
Text sequence can be maneuvered within the sequence view of the construct by cutting and pasting. Sequences can be selected, cut (or copied) and pasted into another window, another construct, or elsewhere in the same construct. It can also be pasted into another program through the clipboard. If a segment of a construct is pasted into a text editor, the DNA sequence will be pasted; if it is pasted into a graphics program, the clipboard will be treated as a graphic and will include all elements of the selection. If you have selected text from another program and transferred it by pasting from the clipboard into

the Gene Construction Kit, the text will be placed at the location of the cursor.

## **Silent Mutations**

In cloning projects, there is often a need to manipulate fragments that do not have restriction sites at convenient locations. It is therefore useful to make a change in the DNA sequence to allow the creation or removal of a restriction site. In addition, it sometimes happens that the site of the sequence change lies within a coding region. In this case, it is desirable to be able to create the sequence change at the DNA level without changing the protein sequence coded for by that stretch of DNA. This type of mutation is called a **silent mutation** because it does not alter the encoded protein. Tutorial 6: “Cloning a DNA Segment and Silent Mutations”, page 2-23 provides a hands-on illustration of removing a site through silent mutation.

GCK allows the creation of new restriction sites by silent mutation and also allows the removal of existing restriction sites through silent mutation. Before a silent mutation can be created, you need to select a protein region so that the program knows which reading frame is to be preserved. If no protein region is selected, the menu options will be disabled.

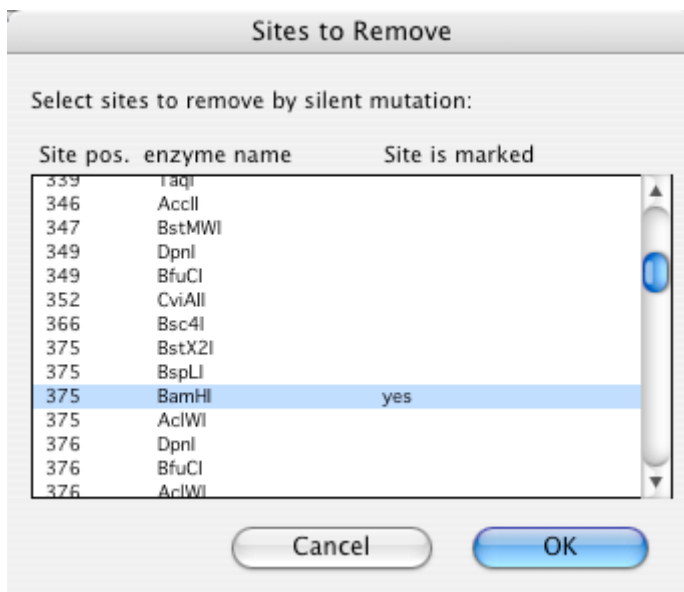


**Figure 3.25: Removing Sites#1**

## **Removing a Site Through Silent Mutation**

Choose Construct > Features > Remove Sites by Silent Mutation to start the process. You will see Figure 3.25. This first dialog lets you choose which enzymes are to

be candidates for removal of a site. It is the basic restriction enzyme chooser. **GCK** needs to be sure that any enzyme site you choose to remove is actually there and also will provide you with other alternative enzymes if you choose to see them. In Figure 3.25, a number of enzymes have been chosen to be viewed as candidates for removal. Pressing the **OK** button will bring up Figure 3.26.



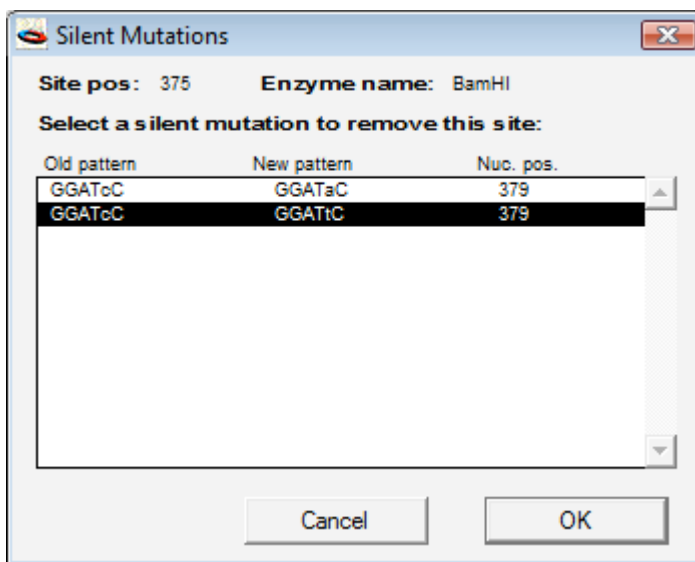
**Figure 3.26: Removing Sites#2**

Figure 3.26, page 3-29 has three columns of data listing possible sites that can be removed by silent mutation. The first column shows the positions of all the possible cuts in the DNA<sup>g</sup> using the enzymes selected in the first step (those that were chosen in Figure 3.25). The second column shows the names of each of the enzymes that have recognition sites in the DNA indicated in the first column. The third column either shows a *yes* or is empty depending on whether the site is actually marked in the current view of the construct. In the list provided in Figure 3.26, you need to specify which site(s) you want to remove by silent mutation. In the figure shown the *BamHI*

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g. Note that this DNA corresponds to the DNA which codes for the region you have selected.

recognition site starting at position 375 has been selected. Pressing **OK** brings up Figure 3.27, page 3-30.



**Figure 3.27: Removing Sites#3**

At the top of this figure are indicated the recognition site position and restriction enzyme being considered. Note that the site position at the top of the window is the first nucleotide in the recognition sequence for the enzyme shown in the window. Like the previous dialog, this dialog also has three columns of data representing the possible changes that can be made in the recognition site (starting at 375) without altering the coding capacity of the DNA corresponding to the selected region. The first column is the current recognition pattern for the chosen enzyme, with the nucleotide to be changed in lower case. The second column is the suggested new pattern that would represent the mutation. The third column shows the actual position of the changed nucleotide. In this case, changing GGATcC to GGATtC has been selected (the C at position 379 will be changed to a T). Once this substitution is made, it is highlighted in the sequence view of the construct as shown in Figure 3.28



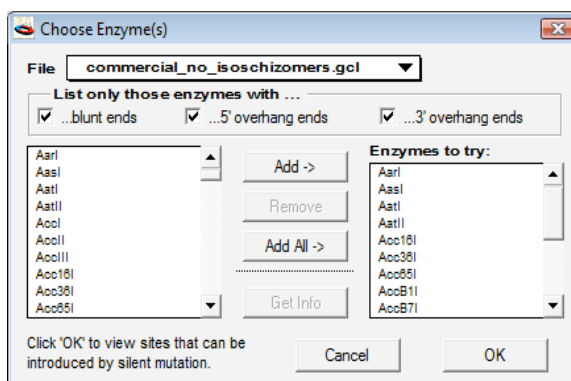
## Introducing a Site Through Silent Mutation

This process is similar to that for “Removing a Site Through Silent Mutation”, page 3-28. Select a protein sequence and then choose Construct > Features >

CTG TGG ATC CTC TAC GCC  
Leu Trp Ile Leu Tyr Ala

**Figure 3.28: Removing Sites#4**

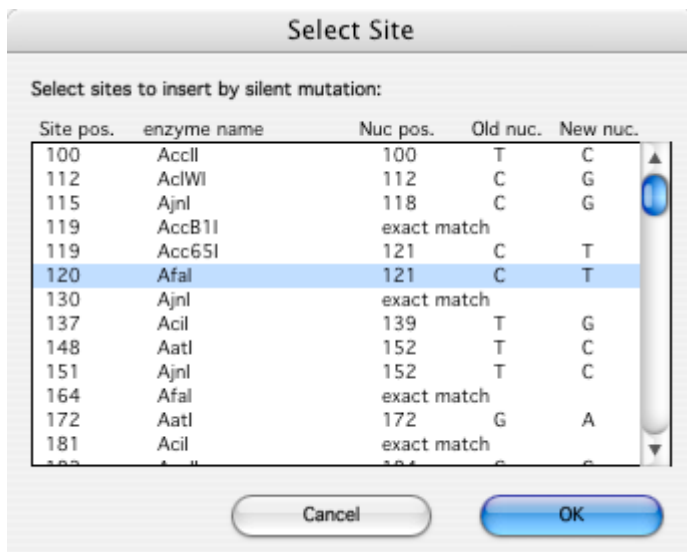
Insert Sites by Silent Mutation to start the process. You will see Figure 3.29, page 3-31, in which you can choose those enzymes that are candidates for possible sites in the DNA corresponding to the region you have selected. In this



**Figure 3.29: Inserting Sites#1**

step you might limit your selection of enzymes to those you have in the lab, or to one or two specific enzymes that might meet other criteria you have in your cloning project. Pressing **OK**, takes you to the next dialog, shown in Figure 3.30, page 3-32

There are 5 columns in this list. The first column (*Site Pos.*) shows the position of the start of the recognition sequence for the new enzyme site to be created. The second column shows the name of the enzyme whose recognition site will be created. Column 3 lists the position of the individual nucleotide that is to be altered; in case there is an exact match with the recognition sequence, “*exact match*” is shown. Columns 4 and 5 show the current nucleotide and the nucleotide it will be changed into. In Figure 3.30, page 3-32, an *AfaI* recognition site will be created starting at nucleotide 120 by changing the C to a T at position 121. Pressing **OK** will make the change-



**Figure 3.30: Inserting Sites#2**

Figure 3.31 shows the result of this operation. Note that the altered nucleotide is in lower case and the restriction site is marked..

AfaI  
 GTC ATC CTC Ggt ACC GTC ACC  
 Val Ile Leu Gly Thr Val Thr

**Figure 3.31: Inserting Sites#3**

## **Chronography - Tracking Construct History**

Chronography was discussed in length in Tutorial 7: “Chronography – Tracking Cloning History”, page 2-30. Please see that tutorial for details on how chronography works. Chronography is one of the most powerful features of the Gene Construction Kit. Because of its uniqueness, however, it is also one of the most challenging to understand and utilize effectively.

Chronography is designed to perform two major functions:

1. to provide a mechanism to allow you to **maintain alternate views** of a construct, each having different sets of features displayed, and

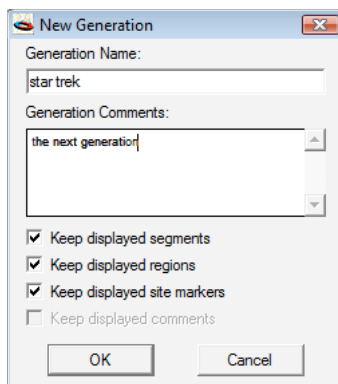
2. to provide a mechanism to **archive the steps utilized in developing a construct**.

Keep in mind that the sequence of the DNA represented in the construct is not changed by any Chronography operations. All Chronography does is change the appearance of what is visible on the screen. Chronography allows you to define different **generations** or views of the construct. Each generation represents a way of displaying different aspects of the construct without changing the DNA sequence in any way.

Using Chronography, one view of a construct might be defined in which the different pieces of DNA (e.g. vector, promoter, cDNA) used to derive a construct are indicated. Another view might show the transcribed regions of the DNA along with initiation and polyadenylation site markers, and yet another view might show some important restriction enzyme sites. It would prove awkward and distracting to show all these features at the same time in the same diagram. It would also be quite tedious to have to indicate each of the features every time you wanted to display them. By defining each of these views as a different generation, you can show any attributes or combinations of attributes you desire, while maintaining the capability to view the alternate displays at any time.

One use for Chronography is to actually use generations to keep track of the different steps used in getting to the current stage of the construct. For example, you decide that you want to place a polycloning sequence into **pBR322** to give you a primitive expression vector for a number of other experiments. You would like to define a new generation (view) of the construct which illustrates those parts of the construct that were derived from **pBR322** in contrast to those that contain the promoter and the polycloning site.

Viewing the current generation of **pBR322** (see Figure 2.29, page 2-30) and choosing **Format > Chronography > Start New Generation...** will bring up Figure 3.32, page 3-34. You can choose which properties of the current generation you wish to appear in the newly created generation by using this dialog. The settings shown in Figure 3.32 will lead to a new generation containing just the red circle of DNA and no sites or regions marked. This new generation



**Figure 3.32: Starting a New Generation**

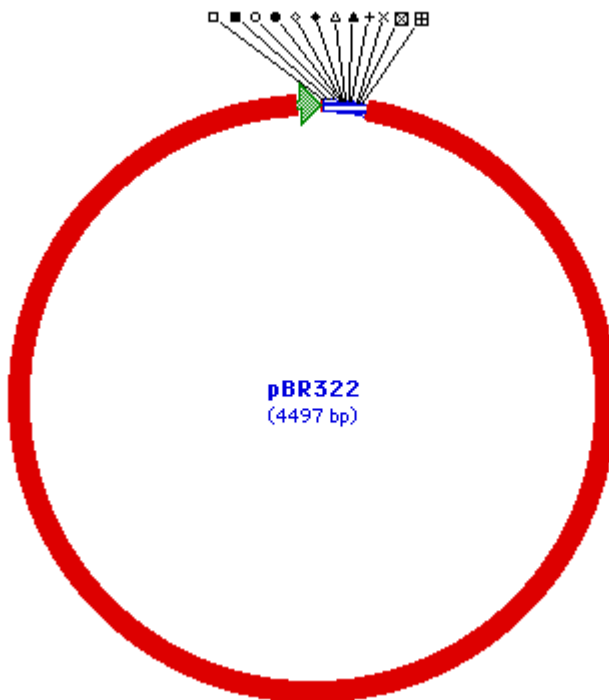
becomes the current generation and, like all “current generations”, is referred to as generation#0. The view that was shown in Figure 2.29, page 2-30 (what used to be the current generation) is now one generation in the past and is called generation#1.

Once the promoter and polycloning site are pasted into the current generation that was just created, a construct with the appearance of that in Figure 3.33, page 3-35 will be created. This shows a construct whose current generation includes the original pBR322 along with the two fragments just pasted in (along with any comments associated with the fragments).

Choosing Format > Chronography > Show Previous Generation will reveal these fragments as a thin gray line (Figure 3.34), indicating that they were not present in the previous generation. Chronography, therefore, can archive the steps taken in generating the construct and can be very useful, especially since all comments can be searched rapidly.

If fragments pasted into a construct contain their own history, that history will be pasted when the DNA is pasted and will appear when the previous generation is shown. This was demonstrated in Tutorial 7: “Chronography – Tracking Cloning History”, page 2-30.

It is possible (likely) that multiple generations will exist in some of the constructs you accumulate. Given the possibility of displaying different segments

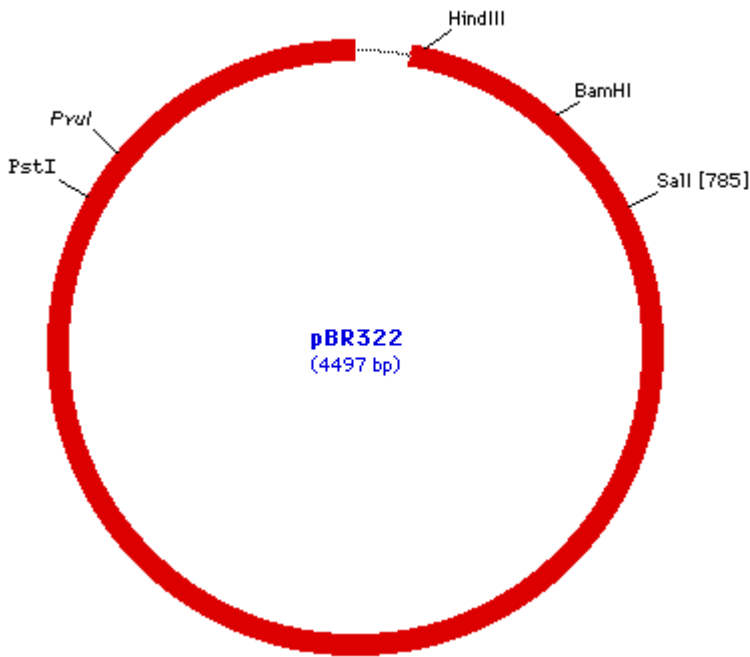


**Figure 3.33: New Fragment in pBR322**

in different generations<sup>h</sup>, it might become difficult to keep track of which generations are being displayed. To address this problem, **GCK** allows you to indicate the display of different generations using the Construct > Display > Show/Hide Generations. When you are showing generations, you will get an extra set of lines on the outside of the construct for all segments not being displayed in the current generation, as shown in Figure 3.35, page 3-37. Those segments being displayed in the current generation will not have an indicator line next to them, which is the case for the new segment just pasted into the construct. The adjacent fragments are being displayed in previous generations, and therefore are indicated as such by the gray indicator around the outside of the circle.

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h. Once any segment is selected, it can be displayed in any generation by using the Chronography menu. In this way it is possible to display different segments in different generations.

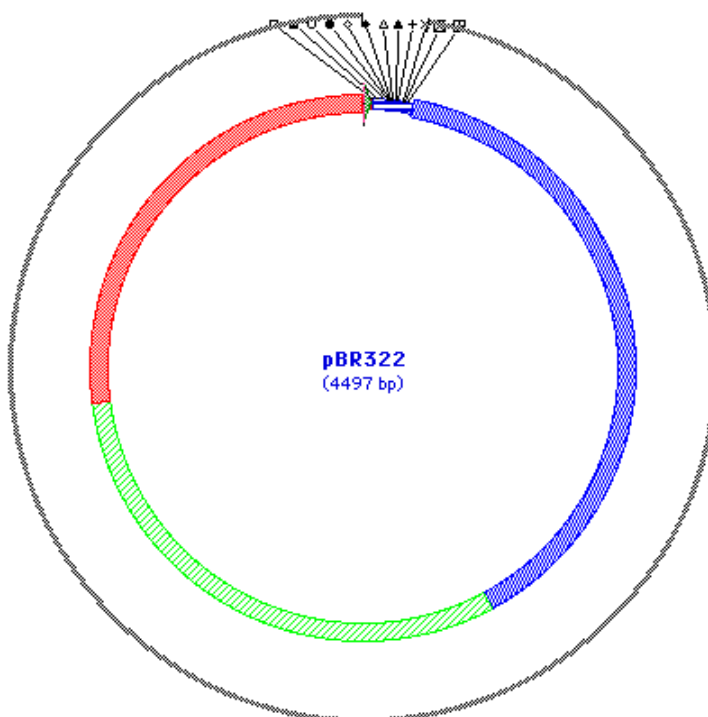


**Figure 3.34: New Fragment, Generation #1**

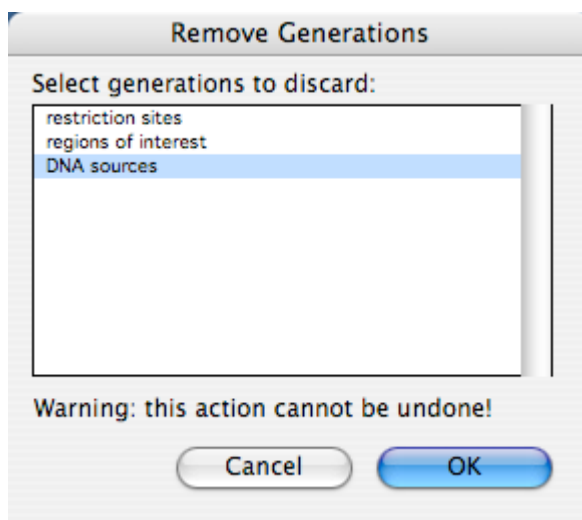
Although you are not likely to ever reach the limitation of 32,000 generations viewable in **GCK**, you may on occasion want to dispose of some of the extra generations you no longer wish to keep track of. This can be done by using Format > Chronography > Discard Generations.... This presents you with the option to throw away any generations of the selected fragments. You will be presented with the dialog box shown in Figure 3.36. It is possible to discard any generation by selecting the name of the generation in this dialog and then pressing the **OK** button. Note that this operation is not undoable and once discarded, this information is lost forever.

## **Finding Open Reading Frames**

It is often desirable to find open reading frames that might exist within a cloned DNA. You can do this using the Construct > Features > Find Open Reading Frames... option. This will bring up Figure 3.37. This dialog explains that **GCK**

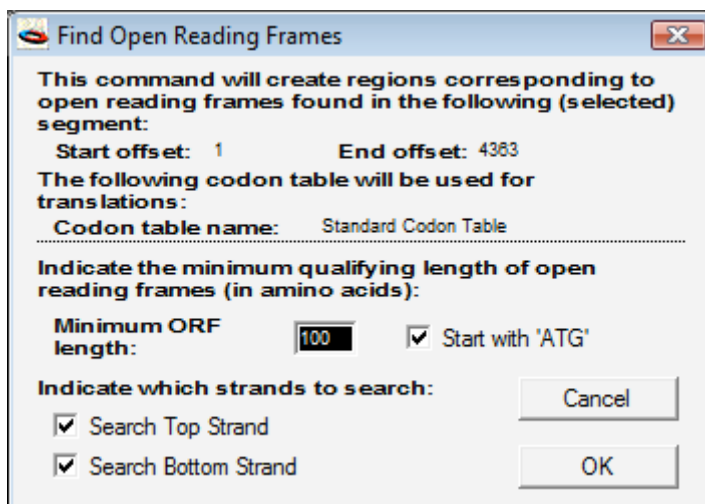


**Figure 3.35: Showing Generation Indicators**



**Figure 3.36: Discard Generations**

will search the construct from positions 1-2964 for open reading frames starting with ATG that will produce proteins of at least 100 amino acids. In this case, the check boxes at the bottom specify that both the top and bottom strands of the DNA will be examined. The default of 100 amino acids for a



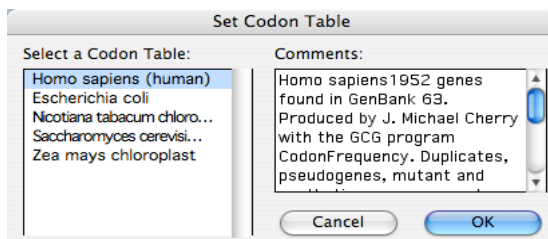
**Figure 3.37: Open Reading Frame Dialog**

minimum can be changed by typing in a new value in the text box. The **Standard Codon Table** is used by default, but this can be changed by choosing Construct > Features > Choose Codon Table..., which will bring up Figure 3.38, page 3-39. In this case a *Petunia* codon table has been chosen. As each table is selected, descriptive text is displayed on the right hand side of the dialog box. The Gene Construction Kit can read Gene Inspector codon tables for use in GCK. This will happen automatically when you place a Gene Inspector codon file into the GCK Data folder. If you open and edit a codon table that is in GI format (see “Edit Codon Tables...”, page 7-44), it will be saved in GCK format from then on. Most of the codon tables are the same, with only the mitochondrial and some trypanosome tables utilizing different codons. Because these files are also used in GI for codon preference analyses, they also contain additional information used by GI.

## **Importing Sequences Into GCK**

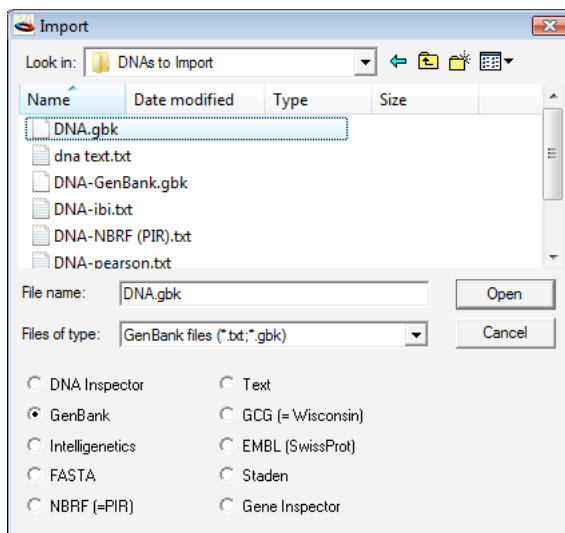
Tutorial 12: “Importing and Exporting Sequences and Other Information”, page





**Figure 3.38: Choosing a Codon Table**

2-60 discusses some issues regarding importing of sequences. Sequence information can be imported into the Gene Construction Kit in several ways. Any sequence information you have generated in other applications can be copied to the clipboard from the other application and then pasted directly into an empty Construct Window. This will give you a sequence that can be manipulated in the standard **GCK** way, either graphically or as text sequence. This procedure will not check the sequence for non-valid characters, however. It will accept whatever you paste in as a valid DNA sequence.



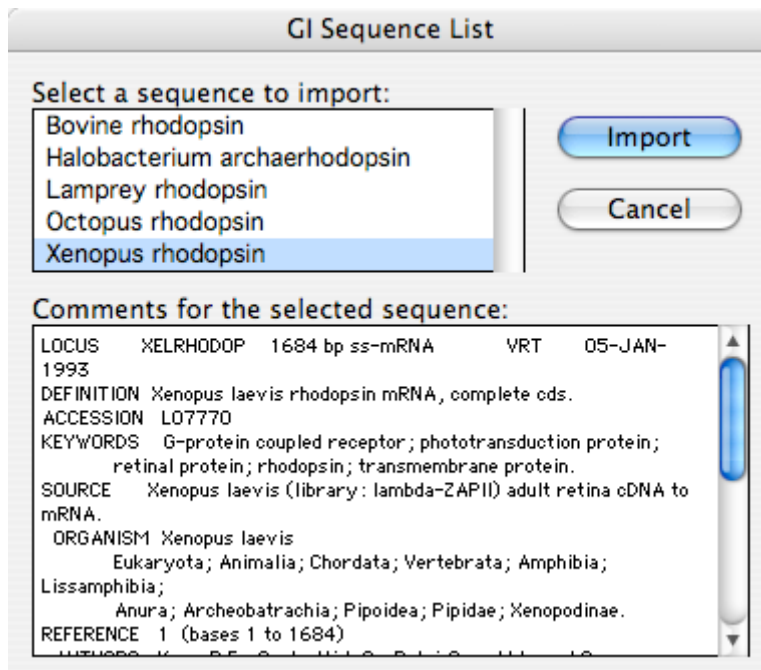
**Figure 3.39: Import Dialog**

Using File > Import... provides you with the ability to import sequence information from files in several other types of formats (see Appendix, "Import File Formats", page 8-2). As shown in Figure 3.39, when you select this

option, you will be presented with the standard file selection dialog box that contains radio buttons for several different types of files. Use the mouse to define the type of file you want to import (in Figure 3.39, Text format is selected), and then select the file from the scrollable list shown in the window. When the **Import** button is pressed, the file will be read in using the assumption that it is in the specified format. If it is in another format, results will be unpredictable as to the actual sequence read in. All files, except Staden format files, that are read in by this method will be checked for invalid characters as they are entered into the Gene Construction Kit, and all Us will be converted to Ts.

All files to import, except DNA Inspector™ and Gene Inspector® files, are stored on disk as TEXT (ASCII) files. The only difference in the files is the precise way in which the sequence and comment information are organized within the file (see Appendix, “Import File Formats”, page 8-2). Files of type TEXT (extension of “.txt” on Windows) can be created by many programs including most word processors (using an Export... or Save As... option), so you should be careful to ensure that the file you are reading is really in the format you have chosen. Any comments associated with the sequence will be read in as General Info to be saved with the entire construct. If you press the TEXT button, GCK simply will take every valid nucleotide character from the file and use it to construct a DNA sequence. It is a simple routine that reads characters one at a time from the disk file and will accept the character if it is a nucleotide and will not accept the character if it is not a nucleotide. You could read in *any* information this way, so once again, be careful.

There are two non-TEXT import options: DNA Inspector™, and Gene Inspector® files. Selecting these radio buttons will only show you DNA Inspector or Gene Inspector files in the scrollable area and will import both the comments and the sequence and determine if the new construct should be linear or circular. Since Gene Inspector files can contain multiple sequences, you will be prompted to choose a specific sequence once you have selected a Gene Inspector file. This is shown in Figure 3.40. The file selected in this case contained a number of rhodopsin sequences, all of which are listed in the scrollable area on the left. Clicking on an item in the list, like *Xenopus rhodop-*



**Figure 3.40: Importing a Gene Inspector Sequence**

*sin*, will show comments (if any) associated with that sequence in the bottom of the window and then allow you to import that particular sequence by pressing the **Import** button.

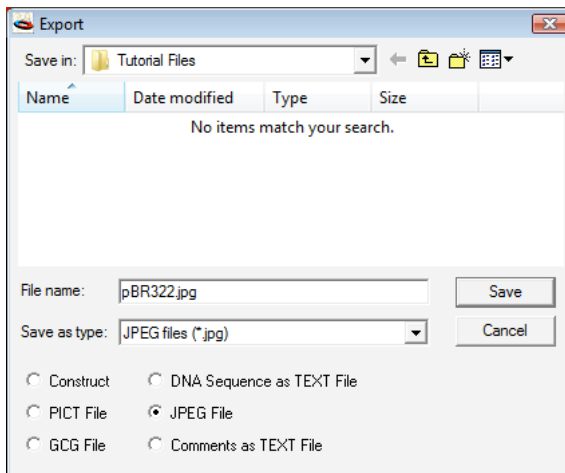
Finally, you can **import graphics** images into the Illustration Window of the Gene Construction Kit through the clipboard. Once the images are placed into the Illustration Window they can be moved around like any other object, but you will not be able to edit them any further.

## **Exporting Data From GCK**

Information from the **GCK** can be exported in a number of ways from each of the different window types of the program. This is accomplished using the File > Export.... Tutorial 12: "Importing and Exporting Sequences and Other Information", page 2-60 discusses some issues regarding exporting of sequences.

## Exporting Constructs

When viewing the construct as graphics in the Construct Window, choosing File > Export... will give you the window shown in Figure 3.41, page 3-42. The



**Figure 3.41: Export Graphics Construct**

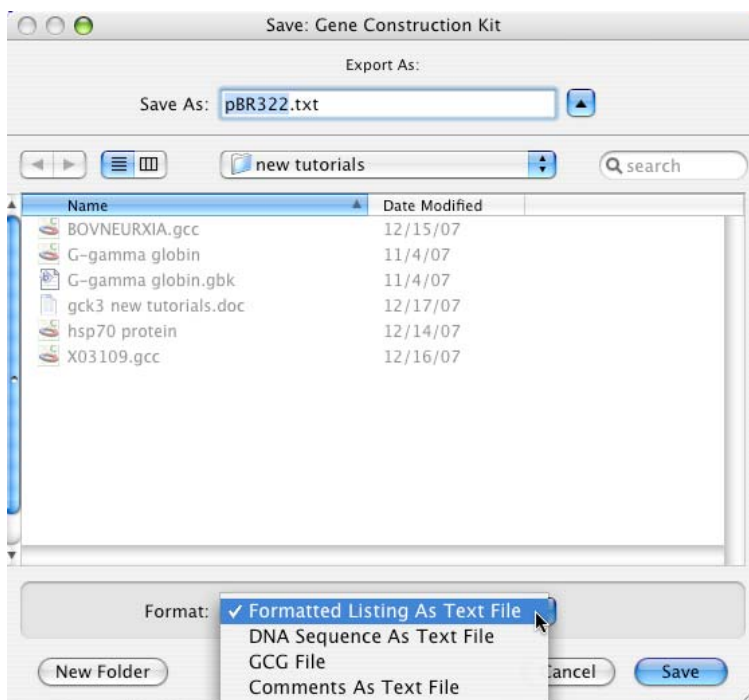
default condition (that which appears when the dialog box first opens up) is to save as a Construct. This is the normal **GCK** format. Choosing **PICT** will save the image of the construct in the currently active window as a Macintosh PICT file. PICT files can be read by most Macintosh and Windows applications that deal with graphics. If you want to work with the image in a Windows program that does not open PICT files, you can export the construct image as a JPEG. This option will allow you to edit the construct graphics using sophisticated graphics programs for special effects. Note that the PICT file will contain the image of the construct but will not contain any sequence information.

Selecting **DNA Sequence as TEXT File** will create a TEXT file (\*.txt on Windows) containing the sequence of the construct. This file can be opened by any text processor. The TEXT file format is also called an ASCII file format and can be transferred to any other computer operating system that can deal with ASCII files, including most mainframe computers. This can serve as a means of transferring the sequence information generated in the **GCK** to other programs for analysis. Selecting the **GCG File** button will save the

sequence and associated comments (for the current view of the construct) as a “Wisconsin Package” GCG file. This file format is a common format used by many programs to store information and is very useful for sharing sequence files with colleagues who might have different analysis programs.

The **Comments as TEXT File** option will save a complete list of all the comments associated with the current construct for all generations. The list will be arranged by generation and by position from the origin (or 5’ end). This provides a convenient way of archiving a hard copy of the information contained in all the objects from all the generations of the construct. This comment file format is listed in the Appendix, “Comment Export Format”, page 8-4.

When viewing the construct as sequence in the Construct Window, choosing File > Export... will give you the window shown in Figure 3.42, page 3-43. In this

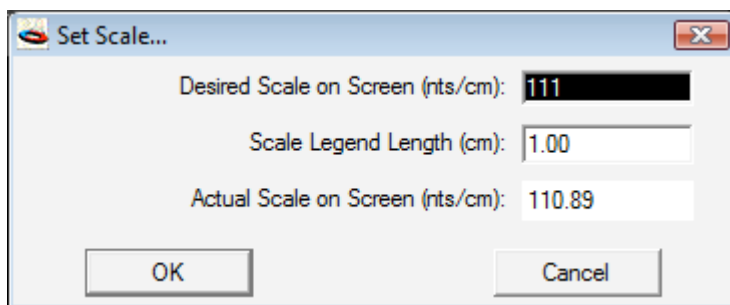


**Figure 3.42: Export Sequence Construct**  
case, there is an extra option called **Formatted Listing as TEXT file**. This

option will produce a text file containing all the line numbers, restriction sites, spacing, and amino acid sequences. Note that in order to view these output files correctly formatted in another application, you will need to use a mono-spaced font like Monaco or Courier.

## **Magnification and Construct Scale**

The scale legend (see Figure 3.1, page 3-1) indicates the scale at which the construct is drawn in nucleotides/centimeter. The scale legend can be shown or hidden using Construct > Display > Show/Hide Scale. When you resize the window and you have selected Construct > Magnification > Fit To Window, the scale legend will be updated to show the new scale. Double-clicking on the scale legend will allow you to set the desired scale as shown in Figure 3.43, page 3-44. The scale can also be set using Construct > Magnification > Set Scale.... Once the scale is set manually, the construct size will not automatically change scale when you change the window size. To have the construct fit the window



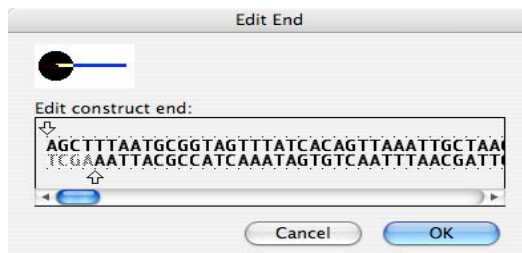
**Figure 3.43: Set Scale**

choose Construct > Magnification > Fit To Window. Construct > Magnification > Fit to Printer Page will resize the construct to fit within the currently selected page size. Construct > Magnification > Zoom In/Zoom Out will enlarge the current construct size or reduce the current construct size. Only the Fit to Window option will allow the construct scale to be resized automatically when the window is resized, since the construct will always be redrawn to fit the window size.

## **Editing Construct Ends and Redefining the Origin**

Linear constructs have ends which can be edited by placing the cursor at the

end of the construct and then choosing Construct > Edit Construct End. A dialog box for the right (3') end of a DNA fragment is shown in Figure 3.44. The ends



**Figure 3.44: Editing a Construct End**

of the DNA strands can be adjusted by dragging the arrowheads to the desired location. Using this option, one can create and then store linkers or adapters as short constructs having precisely defined ends. These could then be used to paste into other constructs when appropriate.

Circular constructs do not have ends, but it is possible to redefine the origin for these constructs by choosing Construct > Redefine Origin. This option will rotate the construct so that the current insertion point becomes the origin of the circular construct and is displayed at 12 o'clock. If you have a selected segment of DNA and choose Construct > Redefine Origin, the 5' end of the selection will become the new origin.



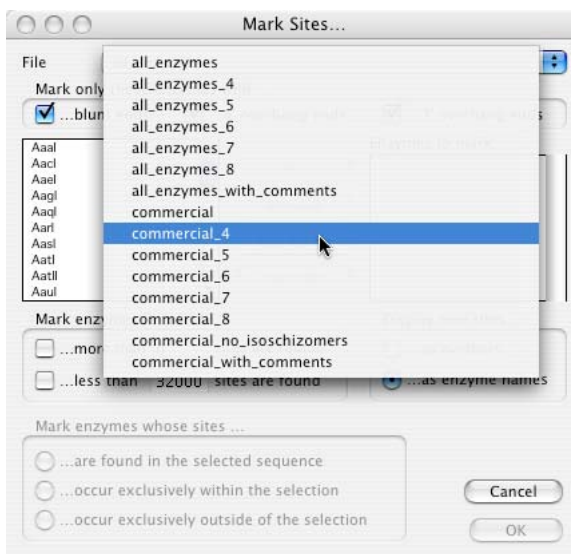


# **Chapter 4**

## **List Window**

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The List Window is the simplest window type in the Gene Construction Kit®. Lists are accessed by **GCK** when Construct > Features > Mark Sites... is chosen. All lists that are in the **GCK Data** folder will be available in a popup menu within the Mark Sites... dialog, as shown in Figure 4.1. In this case the

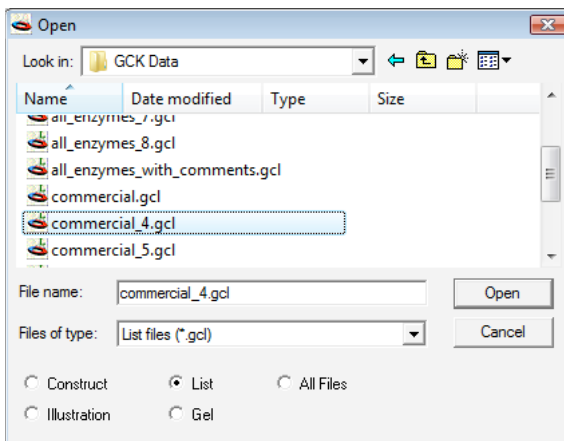


**Figure 4.1: Lists Available in Mark Sites Dialog**

**commercial\_4** list is selected and will be the list that is shown in the scrollable area on the left of this dialog once the mouse button is released. The lists you see might be different from those shown in this figure. You can always obtain the most recent restriction enzyme lists from Textco BioSoftware's web site. The lists are usually updated several times a year and will contain all known enzymes.

## **Opening List Files**

List files can be opened by choosing File > Open..., and pressing the **List** button as shown in Figure 4.2, page 4-2. All of the list files supplied with GCK can

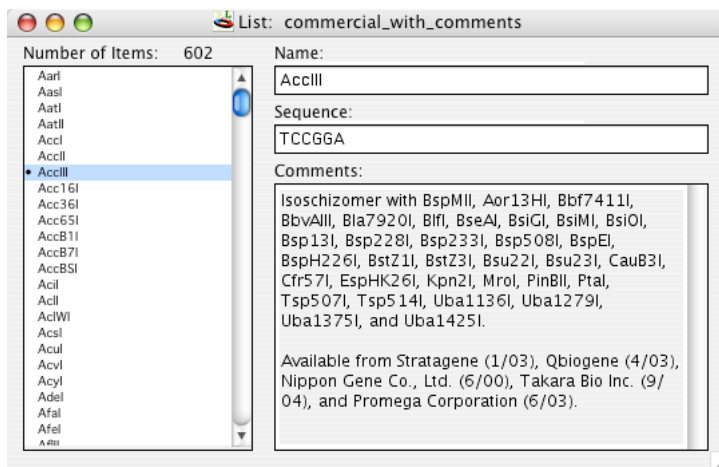


**Figure 4.2: Opening a List File**

be found in the **GCK Data** folder. If you create your own list files, they should be placed in the **GCK Data** folder to become available from within the Mark Sites dialog (Figure 4.1). If you obtain new files from Textco's web site, they should also be placed directly into the **GCK Data** folder. The **GCK Data** folder is a common place to store lists and codon tables. If the files are not in this "blessed" folder, they will not be available for use in the program (although you will still be able to open and edit them).

## **The List Window**

An example of a List Window is shown in Figure 4.3. It is designed to store lists of sequences to be recognized in constructs. Each entry in the list contains a name, a sequence, comments about the entry, and the position of the cut site within the sequence. List windows can store lists of restriction enzyme sequences, linkers, and specific gene features such as poly(A) signals, TATA boxes, etc. Lists might also be used to store primer sequences for PCR. Cut sites are relevant only to restriction enzymes and are of no consequence in other kinds of sites. If no cut site is defined for the entry in the list, the recognition marker will be placed at the beginning of the recognition sequence.



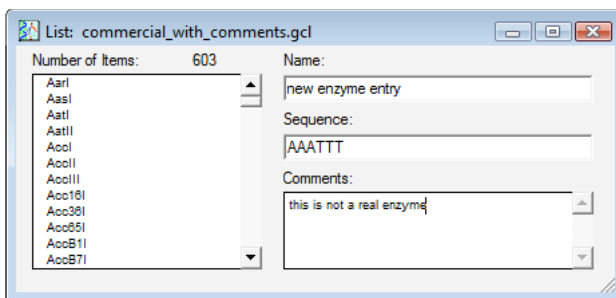
**Figure 4.3: A List Window**

Comments are optional and can be used to list isoschizomers or other information.

When an item is selected in the list, the three fields on the right of the window become filled. In Figure 4.3, *AccIII* in the **commercial\_with\_comments** list provided with **GCK** has been selected. Selected items in lists will have a bullet (•) next to them to indicate that the information on the right side of the window pertains to the bulleted item on the left side of the window. This list (with comments) contains information about isoschizomers of the selected enzyme and sometimes contains possible sources for the enzyme. Note that if you mark sites with this enzyme list, all of the enzyme comments will be saved for each site marked in the construct.

## **New List Entries**

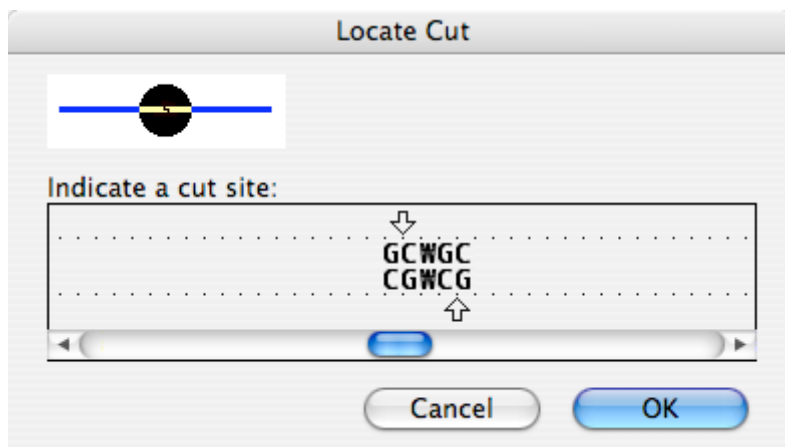
Choosing List > New List Entry will provide a blank entry form to enter the name, sequence, and comments to be associated with your new entry in the list. It will also create a new list entry temporarily indicated by a bullet with an ellipses (...) in the list at the left. Fill in the name of the new entry, the sequence you want this entry to represent, and any comments to be associated with sites on the construct when that entry is used to mark sites. This is illustrated in Figure 4.4.



**Figure 4.4: Creating a New Entry**

## **Defining a Cut Site**

The list entry does not have to represent a restriction enzyme, but can repre-



**Figure 4.5: Defining a Cut Site**

sent a linker, a PCR primer, or other sequence to which you want ready access. If the new entry represents a restriction site, you must also define the cut site for the enzyme. This is done by choosing **List > Define Cut Site** which will bring up a dialog box like that shown in Figure 4.5. The actual cut site is defined by either dragging the arrows to the appropriate locations along the DNA sequence, or by clicking the mouse at the position to which you wish to move the arrow. If an arrow is moved away from the actual sequence into neighboring DNA, an indicator will appear displaying the number of nucleotides between the end of the recognition sequence and the current position

of the arrow.

**GCK** can use as many lists as are present in the **GCK Data** folder. Textco has included some standard lists, including lists of commercially available enzymes that represent 4-, 5-, 6-, 7-, or 8-base recognition sequences.

## **Creating New Lists**

List entries may be selected by choosing them in the list at the left side of the dialog (Figure 4.3, page 4-3). Once selected, list entries can be copied (or cut), and pasted into other lists. As in most standard lists, you can select adjacent multiple entries by clicking on each additional entry with the shift key down. To select non-adjacent entries, hold down the command key (Macintosh) or the ctrl key (Windows) and then click with the mouse on the new item to select.

To create a list of enzymes that you use often, you might start with the commercially available list and then cut out those enzymes that you do not use. Choosing **File > Save As...** will allow you to save the new (edited) file under a new name. If you only have a small set of enzymes you want in a new file, it might be easiest to create a new list file (**File > New...**) and then paste in the enzymes you want in that new file.

The list **commercial (+comments)** contains comments listing any isoschizomers and commercial sources for the enzyme selected.

## **Alphabetical Lists**

Lists can be defined as being alphabetical or not by selecting **List > Alphabetical List**. When this item is checked, the list will always be maintained in an alphabetical order. This means that creating new entries, renaming entries, cutting from and pasting into the list will all result in alphabetical lists. **GCK** will sort the list after you have performed your operation. If the **Alphabetical List** option is not checked, pasting, cutting, etc. will not necessarily result in an alphabetical list.

A non-alphabetical list might be used for storing linkers in order or size or

date of creation or some other criterion.

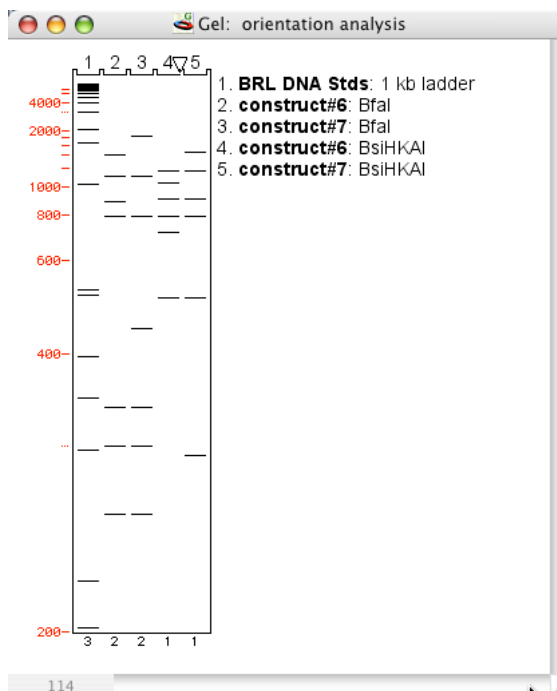
## **Exporting Lists**

Exporting from the List Window provides similar choices to those just described already. Selecting File > Export... will give you a dialog box similar to that shown in Figure 3.41, page 3-42, but only **List** and **TEXT** buttons will be available.

# Chapter 5

## Gel Window

Gel electrophoresis is a standard analytical tool used to track construction project progress. It is often desirable to examine potential fragment sizes from a digest to facilitate the selection of appropriate construction pathways. The Gel Window, shown in Figure 5.1, allows you to view predicted gel electrophoresis patterns for any construct(s) together on a single gel.



**Figure 5.1: A Gel Window**

The cursor changes into a horizontal I-beam when it is over the gel and functions as a **size indicator**. The lower left corner of the window itself pro-

vides a constant readout of the size (in nucleotides) a fragment would be at the current location of the cursor. It provides a convenient way to see an approximate size for any band in the gel.

## **Legend**

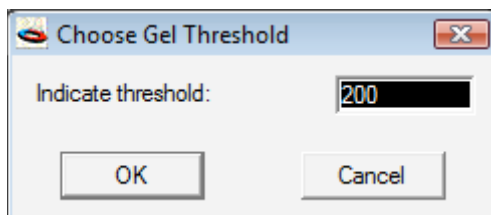
The **legend**, which is visible just to the right of the gel, indicates which construct and enzyme(s) were used to produce the banding patterns in each lane, and in the case of partial digests, specifies the extent of digestion (e.g. - see lane 4, which indicates that 7 out of 8 possible sites were cut). The legend consists of lane numbers and the actual legend adjacent to the gel. The legend can be made visible or not using `Gel > Show/Hide Gel Legend`.

At the bottom of each lane is an indication of the number of fragments that have migrated off the bottom of the gel. These numbers can be made visible or not by using the `Gel > Show/Hide Runoff Count` menu option.

Along the left side of the gel (to the left of lane 1) are a set of standard size indicators. You can show or hide the standard numbers by using `Gel > Show/Hide Standard Sizes`. Each solid band in the standard represents a multiple of the threshold value (see “Threshold”) for the gel. The dotted lines represent 1.5x and 15x the threshold value. We have also included a number of standard digests that you can copy from the “standards gel” in the Tutorial folder to the particular gel you are examining.

## **Threshold**

The threshold value, which is set using `Gel > Set Threshold...`, represents the minimum sized fragment that remains on the gel. This will bring up Figure 5.2.



**Figure 5.2: Set Threshold**



By changing the threshold, you can do the equivalent of running different percent agarose gels. It is very useful for pointing out which size range might be useful for the analytical gels that you actually run in the lab.

The gel pattern is determined using the algorithm of Schaeffer and Sederoff [*Analytical Biochem.* **115**:113-122 (1981)], which calculates the mobility as inversely proportional to the molecular weight of the molecule. This has proven to be more accurate than the more common method of calculating the mobility in proportion to the logarithm of the molecular weight and functions equally well on agarose and acrylamide gels.

## Viewing a Fragment Size Table

As in the Construct Window, the display can be changed from a graphics view to a TEXT view using Gel > Display Table. Shown in Figure 5.3 is a repre-

Gel: orientation analysis		
706	658 (BsiHKA I)	1363 (BsiHKA I)
501	1364 (BsiHKA I)	1864 (BsiHKA I)
85	4846 (BsiHKA I)	4930 (BsiHKA I)
Digest #5, construct#7 (5177) : BsiHKA I		
size	from	to
1448	2237 (BsiHKA I)	3684 (BsiHKA I)
1161	3685 (BsiHKA I)	4845 (BsiHKA I)
904	4931 (BsiHKA I)	657 (BsiHKA I)
786	950 (BsiHKA I)	1735 (BsiHKA I)
501	1736 (BsiHKA I)	2236 (BsiHKA I)
292	658 (BsiHKA I)	949 (BsiHKA I)
85	4846 (BsiHKA I)	4930 (BsiHKA I)
Digest #1, BRL DNA Stds (83885) : 1 kb ladder		
position	site	
75	1 kb ladder	
209	1 kb ladder	
363	1 kb ladder	
564	1 kb ladder	
784	1 kb ladder	
1081	1 kb ladder	
1426	1 kb ladder	
1822	1 kb ladder	

**Figure 5.3: Gel Window as Text**

sentative listing of the two formats for the Table listing of the digest fragments. This view is much more precise than the cursor in the graphics view in giving the exact sizes of the fragments. It provides a complete list of every fragment and which enzymes led to the production of the particular fragment.

The first listing is given in descending size order and the second listing is given from 5' to 3' end for digests of linear constructs, or clockwise from the origin for digests of circular constructs. Especially in the case of partial digests, the identification of each end of a fragment is very useful.

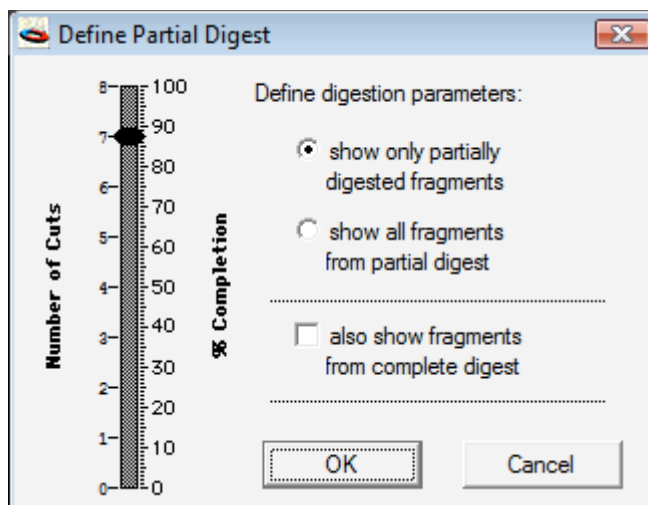
## **Creating and Rearranging Gel Lanes**

Gel lanes are created by pasting in information from a Construct Window or from another gel lane. To create a new gel lane from a construct is simple. In the Construct Window select the marked sites that you wish to use to create the digest and then copy them to the clipboard using **Edit > Copy**. Make sure to select the site markers and not the fragments themselves (an example is given in Tutorial 9: "Running Gels and Orientation Analysis", page 2-45). Multiple sites can be selected by holding down the shift key and clicking on all desired sites. All sites of a given type (e.g. all *HindIII* sites) can be selected by double-clicking on any one of them. Once the sites are copied to the clipboard, either create a new Gel Window using the **File > New...**, or open an older Gel Window, select the location to paste the new lane by clicking at the top of the gel between two lanes, and then paste the digest from the clipboard.

Once the digests have been defined, you might want to rearrange or even delete some of the lanes. This can be done by following the same logical steps you know from the Macintosh interface. Click on a lane to select it (use shift-click to select more than one lane) and then use **Edit > Copy**, **Edit > Cut**, and **Edit > Paste** to move and replace lanes on the gel. You can rearrange lanes on a single gel and even paste lanes from one gel into another. This might be useful if you have a specific set of standards that you run on your gels. You could set up a gel containing those standards and save it. Anytime you want to put a standard lane on a new gel, simply open up your saved standards, Copy the lane you want, and paste it into the new gel.

## **Partial Digests**

Occasionally, restriction enzyme digests do not go to completion, resulting in extra bands on the gel. Using the **GCK**, you can simulate these partial



**Figure 5.4: Partial Digest Dialog**

digests to see if they can explain your laboratory gel results. Select the lane containing the digest that you want to view as a partial digest and then choose Gel > Show Partial Digest.... You will see a dialog box like the one shown in Figure 5.4, page 5-5. The extent of the digest can be set by using the slide control on the left of the dialog box. The left scale is the number of cuts to be made, and the right scale shows the corresponding percentage completion. The **“also show fragments from complete digest”** check box draws the fragments that would be present in a complete digest as solid lines in addition to the partial digest fragments that are shown as dotted blue lines. The **“show only partially digested fragments”** radio button will show only the partially digested fragments (those containing internal sites), while the **“show all fragments from partial digest”** will draw both partial and complete digest fragments from the digest as dotted lines.

You can change the partial digestion parameters for any lane by choosing Gel > Show Partial Digest... again. Choosing Gel > Show Full Digest from the Gel Menu will display the lane as a complete digest.

## Exporting Gels

Exporting from the Gel Window also provides similar choices: **Gel**, **TEXT**,

and **PICT**. TEXT format will save a list of all the cut sites and fragment sizes as if they were viewed in the Gel Window as a Table (see page 5-3).

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# Chapter 6

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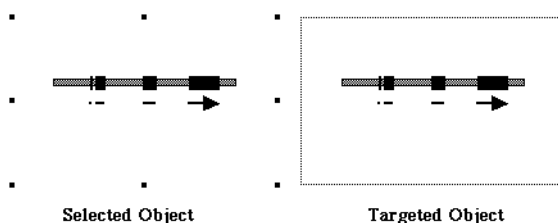
## The Illustration Window

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The Illustration window is designed to fill three needs: 1) to facilitate creation of figures for papers or presentations, 2) to keep track of complex construction projects and allow you the flexibility to change and update the project information easily, and 3) to serve as a library of commonly needed constructs, linkers, adaptors, polycloning regions, gel standards, etc. Constructs, gels, and sequences can be selected in their own windows and pasted into the Illustration Window.

### Selection vs. Targeting

An important concept regarding objects in the Illustration window is the selection or targeting of an object. Clicking once on an object will select the object,



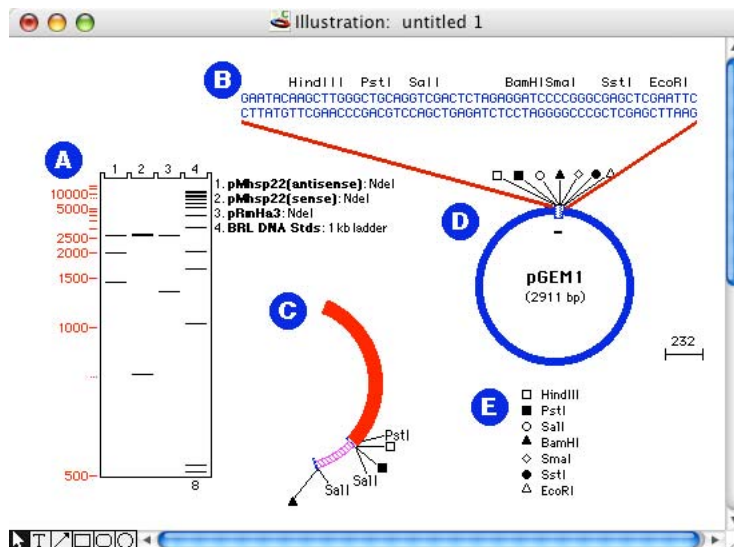
**Figure 6.1: Selection vs. Target**

which will then appear with 8 “handles” around it as shown in the left side of Figure 6.1. A selected object can be resized by dragging one of the handles as in other applications. Clicking in the middle of a selected object allows the object to be dragged to a different location within the illustration. Constructs or Gels that are pasted into the Illustration window can be selected as just described. In addition, these objects can also be targeted by double-clicking on them. A targeted object will have a gray border around it and items within

the object can be edited. If a Gel object is targeted, a Gel menu will appear to the right of the Illustration menu. If a construct is targeted, a Construct menu will appear to the right of the Illustration menu. When either of these objects is targeted, all items in the Construct or Gel menus will be available except for those which change the view of the object. This means that you cannot change from a graphic view to a text/sequence view. This is because a change between graphics and text will most likely require either a change in size of the object or necessitate the appearance of scrollbars. Resizing the object will alter the appearance of the illustration and might cover up other objects, while putting scrollbars on the object will violate the WYSIWYG (What You See Is What You Get) guidelines - what you see on the screen is not what will be printed. Therefore, you are not allowed to make this change.

## Using the Illustration Window

Figure 6.2 shows a typical Illustration Window. There are a number of items



**Figure 6.2: An Illustration Window**


in this window. "A" shows a gel. The lanes present in this illustration were originally selected in a GCK Gel window, copied, and then pasted into the Illustration window to give the figure shown here. "B" shows a sequence that was copied from the sequence view of a construct. This sequence represents


the polycloning site from the construct pGEM1, which is shown in part “D”. The Illustration window has some built in intelligence. If you copy a part of a circular construct from a Construct window and then paste it into the Illustration window, the segment will be displayed as an arc, as shown in part “C”. This makes it easy to display cloning strategies involving segments of circular constructs. The last part of this Illustration is part “E”, which is a legend to the sites shown in “D” for pGEM1. Legends like this can be created automatically in the Illustration window by selecting the site marker symbols in a Construct window, copying them, and then pasting into the Illustration window. The built in intelligence of **GCK** will interpret the site markers being pasted into the Illustration window as a legend<sup>a</sup>.

The Illustration window **tool palette** resides in the bottom left corner of the window border, and is shown in Figure 6.3. Six tools are available. The first



**Figure 6.3: Illustration Tool Palette**


one, the **selection tool** - , allows you to click once on any object with the mouse to select that object within the Illustration window. You can select constructs, gels, sequences, or any objects generated with the other tools in the Illustration Tool Palette. You could also select any graphic objects pasted into the Illustration from outside the Gene Construction Kit®. By dragging with the selection tool, you can select multiple objects in the Illustration window. Once selected, objects can be moved around the window.


The **TEXT tool** -  - allows you to type text anywhere in the window - this ability is sometimes called **sidebar text**. Click the mouse at the location you wish to place the text and then just type. The text generated can be edited using standard cut and paste operations and attributes such as Color, Font, Style, and Size (available under the omnipresent Format menu). To move typed text around the window as an object, make it the *selection* (Figure 6.1, page

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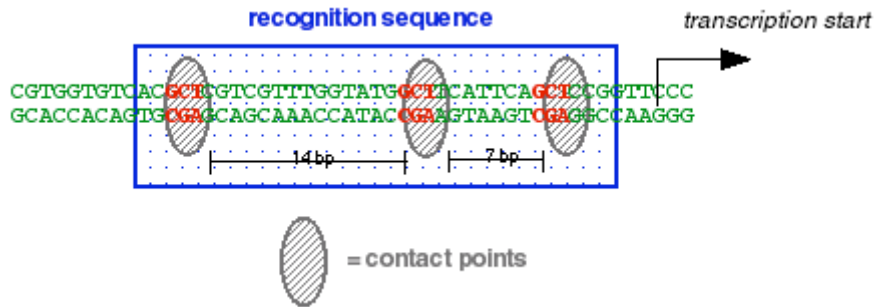
a. Note that if the site markers are pasted into a Gel window, the Gel window would interpret the markers as fragment sizes and would create a new gel lane.

6-1) by clicking it with the selection tool and dragging it to the desired location. To edit the text after it has already been entered, double-click on the text itself to make it the *target* (Figure 6.1). The targeted text object is now editable using all the options available in the Format Menu. New text will always appear in the same font, size, style, and color as the previously entered text unless it is the first text to be entered in the window, in which case it will use the values set in the Preferences submenu (see “Preferences”, page 7-33).

The **line tool** -  - can be used to generate lines. Select this tool and then drag the mouse within the Illustration window from the starting to the finishing location for the line you wish to draw. Once drawn, the line can be altered by using Format > Lines. In addition to line width and color, you can set the direction of the line, and if the line has arrowheads, you can adjust the shape of the arrowheads (see “Lines”, page 7-19). Once lines are drawn, they can be moved at either end (grab the end to move), or moved intact (grab in the middle of the line with the selection tool and drag). If an end of a line is placed within a construct, the line end will stick to the construct and will be moved automatically when the construct is moved in the Illustration window. Holding the shift key down when moving an end or creating a new line, will confine the angle of the line on the screen line to increments of 45°. Newly created lines will have the same attributes as the most recently created lines, or will reflect the Preferences settings.

The **rectangle**, **round rectangle**, and **ellipse tools** -  - can be used to generate shapes for illustrative graphics, as shown in Figure 6.4, which might represent binding sites for specific transcription factors along a gene. Using the Illustration > Send to Back and Illustration > Bring to Front menu options, the different objects can be arranged as desired. The horizontal DNA is actually a linear construct pasted into the Illustration window. Holding the shift key down when creating or editing a rectangle or round rectangle will constrain the shape to a **square**. Holding the shift key down when creating or editing an ellipse will constrain the shape to a **circle**.





**Figure 6.4: An Illustration**

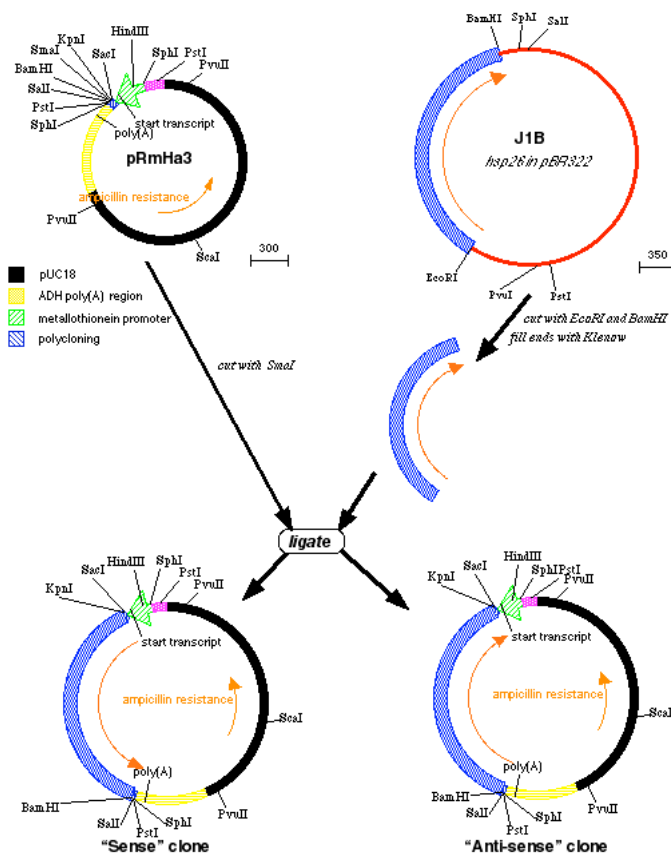
The Illustration window can be used to bring together graphics from other programs with constructs created in the Gene Construction Kit. Any object can be pasted into an Illustration window from the clipboard and can be used to create sophisticated figures in conjunction with constructs created by the Gene Construction Kit. In addition, any Illustration can be saved as a PICT or JPG file to be used by other programs.

## **Tracking Complex Construction Projects**

The Illustration window has some rather remarkable features that make it ideal for tracking complex construction projects. As shown in Figure 6.5, page 6-6, many constructs can be placed in the same Illustration and connected with lines and descriptions. Illustrations are limited to 32,767 pixels on a side, which is about 38 ft. by 38 ft. — large enough for even the most ambitious project. This virtually unlimited size means that projects of extreme complexity can be mapped out and tracked using the Illustration window of the Gene Construction Kit.

Just as TEXT objects in the Illustration window can be edited by double-clicking on them, gel, construct and sequence objects can be edited after being double-clicked. This means that **every construct in the Illustration window can be edited as a construct** after being placed into the Illustration window. Upon double-clicking a construct (making it the target), a new Construct menu will appear to the right of the Illustration menu, as described on page 6-1. This allows you to view different generations, retrieve sequences, change line attri-

### Cloning Hsp26 into pRmHa3



**Figure 6.5: Multiple Constructs in an Illustration**

butes, etc. even after the construct has been pasted into the Illustration window. You can even copy (or cut) a segment from a construct in the Illustration window and paste it into a new Construct window (including all the Chronography that was attached to the segment when it was originally placed in the Illustration window).

The ability to edit a construct within the Illustration window allows you to keep an up-to-date Illustration containing all of the constructs being used in a project and make changes to the constructs as needed. You never have to go back to the original construct files to change them just to maintain a consis-

tent illustration. Because each construct and its associated Chronography can be copied from the Illustration into a new Construct window, you can even use the Illustration as a sort of graphic interactive database for the constructs in a project. Objects like adaptor sequences can be stored in the Illustration window for later retrieval when needed during another project.

## **Changing Magnification**

For large Illustrations that do not fit on the screen, the entire image can be seen by choosing Edit > Show Overview.... This option presents a miniature view of the entire Illustration. The cursor will change to a magnifying glass. Clicking the magnifying glass on a specific location will zoom to normal size with the clicked point centered on the screen.

Magnification can also be changed using the Illustration > Reduction > menu. This menu only allows for the reduction of the view - i.e. “zooming out” so that more of the illustration can be seen. Illustration > Reduction > Reduce is available whenever part of the illustration is out of view. Illustration > Reduction > Enlarge is available whenever the view has been previously reduced. Illustration > Reduction > No Reduction returns the view to its maximum size, and Illustration > Reduction > Full Reduction reduces the entire illustration to fit in the current window size.

Reducing an illustration provides a convenient way to rearrange the components of the illustration on the page. When reduced, it is not possible to target (page 6-1) any of the illustration objects. Targeting an object can only be done when there is no reduction.

The ability to edit constructs from within the Illustration window allows the Illustration window to function as a kind of interactive cloning project management tool. Since each construct’s history is maintained in the Illustration window, it can be used as an automatic archiving facility. The ability to store constructs allows the Illustration window to become a graphical database for constructs. As you become more familiar with the capabilities of the window, you will find more uses of your own for it.

## **Exporting Illustrations**

Exporting from the Illustration Window provides similar choices to those just described for the Construct Window. Selecting File > Export... will give you a dialog box similar to that shown in Figure 3.41, page 3-42, but only **PICT** and **Illustration** buttons will be available.

# Chapter 7

## Menu Items

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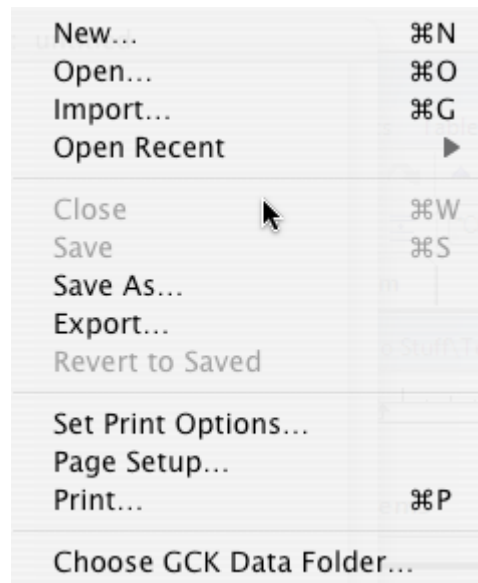
### File Menu

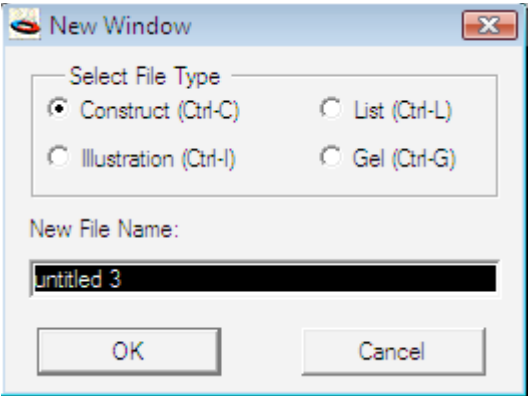
The File Menu deals with reading and writing data to disk or printer, or specifying files to be used by Gene Construction Kit®. The various options allow you to create new windows or files, save files, import or export files, print, and define which data is to be used by GCK.

### New...

File > New... (command-N/ctrl-N) creates an empty window of the type specified by the popup menu item chosen in the new dialog box (Figure 7.1, page 7-2). The

new window can be a **Construct**, **Illustration**, **List**, or **Gel** window. In addition to specifying the type of window, you can enter the name of the window to be created. Note that this does not save the window as a document on disk, it merely gives it a name for identification. If you later choose Save from the File menu, the name you assign in the “New” dialog box will be used to save the file. Instead of using the mouse to select the buttons in this dialog, you can use command key equivalents as shortcuts: command-C/ctrl-C for **Construct**, command-I/ctrl-I for **Illustration**, command-L/ctrl-L for **List**, and command-G/ctrl-G for **Gel**.

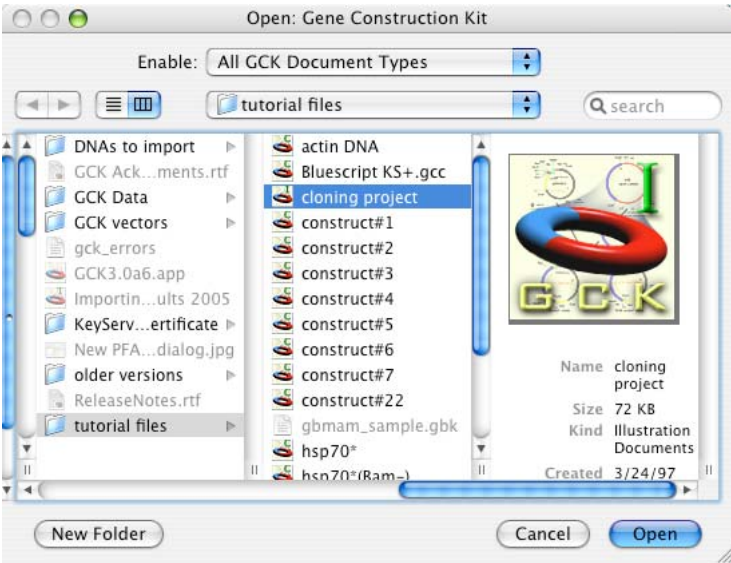




**Figure 7.1: New Dialog**

**Open...**

File > Open... (command-O/ctrl-O) will allow you to select a file to open as shown in Figure 7.2, page 7-2. The buttons on the bottom of this dialog box serve to limit the files that are displayed in the scrollable area to the type that correspond to the selected radio button. In this case, only construct files are shown. As in the New dialog box, the same command key equivalents function

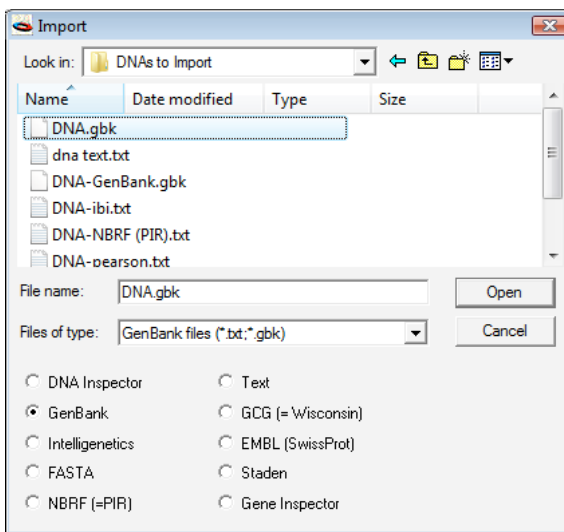


**Figure 7.2: Open Dialog**

as short cuts. In addition, command-A/ctrl-A selects **All Types**. The **Eject** button will eject the disk that is named above the buttons unless it is a hard disk or some other volume that cannot be unmounted. The **Drive** button will show you the files on a different mounted drive. **Open** will proceed to open the file selected in the list on the left and will create a window containing the information from that file, and **Cancel** closes this dialog box. Once you open a file, a new window will appear having the name of the file in the title bar of the window. The type of file will also be indicated in the window title bar. There is a direct correspondence between each window open in the program and a specific file on disk. You can never have open more than one window for any given file on disk, nor can you have more than one file for any given open window.

### **Import...**

File > Import... (command-G/ctrl-G) was discussed in some detail in Importing Sequences Into GCK, page 3-38 and in Tutorial 12: Importing and Exporting Sequences and Other Information, page 2-60. The dialog box is shown in Figure 7.3, page 7-3. The **Import** button imports DNA sequences from dif-



**Figure 7.3: Importing Sequences**

ferent file formats into the Gene Construction Kit. Importing from all formats

except TEXT will preserve the comments associated with the original files. Importing a TEXT format file will simply read the file one character at a time and place the legal DNA characters into a GCK construct sequence. Characters read in from all formats except Staden will be filtered to allow only valid DNA characters (see “Legal Nucleotide Characters”, page 8-1). Any U’s in the sequence will be converted to T’s. DNA Inspector™ and Gene Inspector® files are not TEXT files, but rather are files in a specific format used by those other Textco programs. The other eight file formats contain textual information only. The only difference between these different file formats is the way in which the information is reorganized within the file. These file formats are well established and can be used as a way to transfer information between different programs. If you want to create your own file in a word processing document for importing into GCK, make sure to save it from the word processor as a TEXT only file. If it is not saved as TEXT only, then you will not see it in the list shown in Figure 7.3.

### Importing From File Formats Not Listed

GCK construct files contain sequence information along with other information that GCK uses to display the sequence and allow you to perform certain manipulations on the sequence. The exact organization of the information in that file is dictated by programming needs. Textco will sometimes need to change file formats when new features are added to the program. Since every other software developer faces similar needs and will periodically change their internal file formats, it is difficult if not impossible to support other program specific file formats for importing into **GCK**. Almost every program available, however, can export a sequence as a TEXT file and some can export the sequence into one of the other standard file formats like GCG or GenBank. These exported files can then be read into **GCK**. Another alternative is to copy and paste the sequence from the other program into **GCK**.

### Close

File > Close (command-W/ctrl-W) will close the currently active (front-most) window. If changes have been made since the last Save, you will be shown the dialog box in Figure 7.4. Selecting the **Yes** button will save the changes and





**Figure 7.4: Save Changes?**

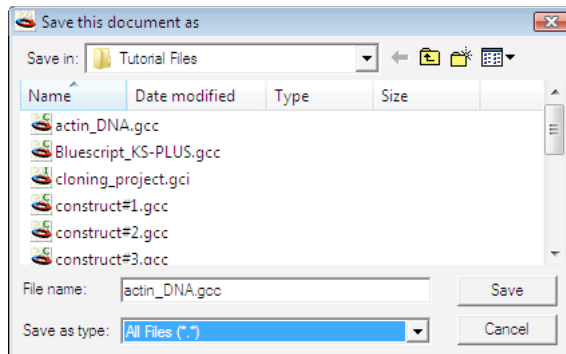
then close the window. Selecting the **No** button will close the window without saving any changes made since the last Save. Selecting **Cancel** will return you to the state you were in before selecting Close.

### **Save**

File > Save (command-S/ctrl-S) will save any changes you have made in the currently active window. It also resets the Revert To Saved menu choice (see "Revert to Saved", page 7-8) to revert to the window conditions that are now being saved.

### **Save As...**

File > Save As... will present Figure 7.5. You do not have any choice in file for-



**Figure 7.5: Save As...**

mats as you do with File > Export...; everything is saved in Gene Construction Kit

format. This option will create a new file of the same type as the original file that was used to create the window (Construct, Gel, List, or Illustration). The original file remains unchanged and the window title changes to the new name. All editing subsequently done in the window is done under the new name. It is always the case that the window title is the same as the name of the file being edited.

### **Export...**

File > Export... will present different options depending on the type of window currently active. This topic was discussed in some detail in Tutorial 12:Importing and Exporting Sequences and Other Information, page 2-60 and in “Exporting Data From GCK”, page 3-41. A number of different output formats are possible in addition to the standard Gene Construction Kit formats. Exported files can be saved as **JPEG** files, along with several other standard file types. These files can be opened by almost any graphics drawing program and should allow you to edit the images you create with Gene Construction Kit. Note that the exported graphics formats cannot be edited in other programs as a collection of objects (e.g. site markers cannot be moved, sequence information is lost, etc.). These manipulations are only possible in **GCK**. Copies created as **TEXT** files contain the textual information available in the currently active window. This could be a sequence of DNA or a table listing gel fragments.

If you are viewing a construct graphically you will see Figure 3.41: Export Graphics Construct, page 3-42. If you are viewing the construct as a sequence instead of graphically, you will see the dialog box shown in Figure 3.42: Export Sequence Construct, page 3-43. One option in this dialog that is not present in the dialog box for the graphics view of a construct is to save a **Formatted Listing as TEXT File**. This option will produce a TEXT file that contains a listing of the sequence as it is currently displayed in the Construct window along with all the other textual information in the window. This would include line numbers (positions) for both DNA and amino acid sequences (if present) and the names of marked sites. Grouping would also be preserved. It would not include any graphical information such as arrows

that indicate regions of interest or dashed lines that indicate line borders. This option can be useful for importing into a word processor later for inclusion in a manuscript.

The last option is selected by choosing **Comments as TEXT File**. This option provides a mechanism to save all the comments associated with a construct. Some constructs can be the result of many steps of manipulation involving DNA from many sources. If you have comments associated with marked sites, Regions of Interest, and DNA segments, and the construct has many generations, it might be difficult to get an overall picture of the history of the construct. This option will create a TEXT file that contains all the comments in the construct from all objects in all generations. The comments will be listed in order, from most recent to most ancient generation and by distance from the origin (or 5' end of linear constructs). The TEXT file can then be opened from any standard word processing package and printed or viewed. The precise format of the output file is shown "Comment Export Format", page 8-4.

### Exporting Illustrations

Exporting from the Illustration Window provides similar choices to those just described for the Construct Window. Selecting File > Export... will give you a dialog box similar to that shown in Figure 3.41, page 3-42, but only **PICT**, **JPEG**, and **Illustration** buttons will be available.

### Exporting Gels

Exporting from the Gel Window also provides similar choices: **Gel**, **TEXT**, and **PICT**. TEXT format will save a list of all the cut sites and fragment sizes as if they were viewed in the Gel Window as a Table (see "Viewing a Fragment Size Table", page 5-3).

### Exporting Lists

Exporting from the List Window provides similar choices to those just described already. Selecting File > Export... will give you a dialog box similar to

that shown in Figure 3.41, page 3-42, but only **List** and **TEXT** buttons will be available.

### **Revert to Saved**

File > Revert to Saved will discard all changes made to the current window since the time of the last save. If changes have been made, you will see the dialog box shown in Figure 7.4, page 7-5. Clicking **Cancel** will return the program to the state it was in when you selected the Revert to Saved option. Clicking **OK** will discard any changes made since the last save and restore the window to the condition it was in at the time it was last saved (or when it was opened if no saves have occurred).

### **Set Print Options...**

Choosing this menu item will bring up a dialog allowing you to set a number of options. The *set margins* values allow you to define the amount of space between the maximum area the printer can utilize and the edge of the usable space in the current window. If this is set to zero, then the maximum area usable by the printer will be utilized in the GCK window. The **Print document name in header** checkbox will force the name of the current document to be printed at the top of every page. **Print page numbers** will cause each page to be numbered when it is printed. These last two options are especially important when printing out multiple page sequence documents.

### **Page Setup...**

File > Page Setup... is the standard dialog box designed to help format printing appropriately. More information on this and other Page Setup... dialogs is available in your Printer and computer manuals. Note that the layout in the dialog box will change when you use different printers.

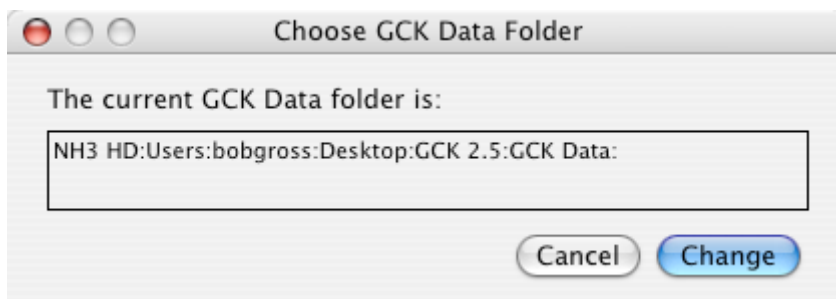
### **Print...**

File > Print (command-P/ctrl-P) will print the currently active window. Note that with the Construct, Gel, and Illustration windows, the image size might be greater

than one page. This can be determined by selecting **Edit > Show Page Breaks** (page 7-15) before initiating printing. If no “page break” lines are visible, then the image will fit on one page. Note also that in the **Construct** window, the image can be scaled to fit on one page by using **Construct > Magnification > Fit to Page** (page 7-43).

### **Choose GCK Data Folder...**

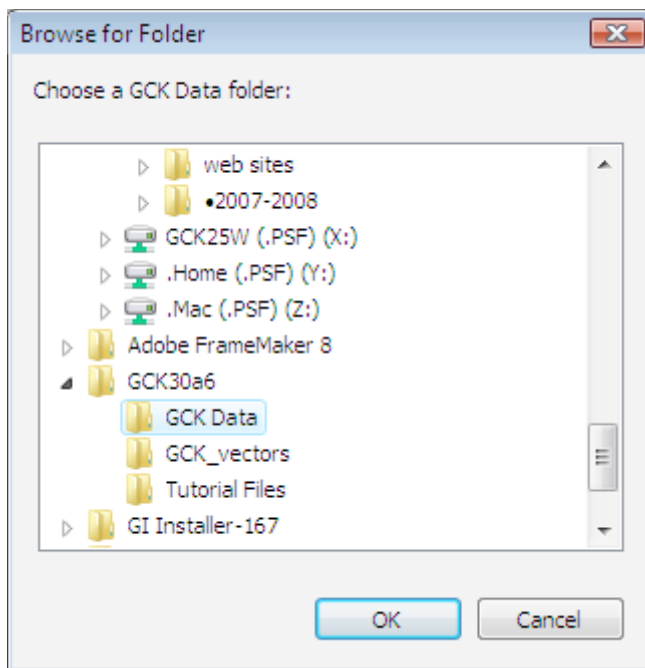
The **GCK Data** folder holds data that is used by **GCK** to perform some of its tasks. This is where restriction enzyme lists and codon tables are stored, for



**Figure 7.6: Current GCK Data Folder**

example. When you first install **GCK**, the **GCK Data** folder will be placed in the same folder as the **GCK** application. **GCK** assumes that the **GCK Data** folder is in this location. If, for some reason, you would like to move the **GCK** application or the **GCK Data** folder to another location, you can tell **GCK** which **GCK Data** folder to use. Choosing **File > Choose GCK Data Folder...** will bring up Figure 7.6, page 7-9. This dialog informs you of the location of the current **GCK Data** folder. In this case it is on a computer whose name is **Ci** and it is located on a volume called **Applications** in the folder called **GCK f**. Two buttons are available. Pressing **Don't Change** will close the dialog and leave everything as it was.

Pressing **Select new GCK Data folder** will bring up Figure 7.7. Use the standard buttons and popup menus to navigate to the folder you are interested in assigning as the **GCK Data** folder. The name in the button at the bottom of this window will change as you navigate to different folders on your disk. Note that the highlighted folder in this figure is **GCK Data**, but the button



**Figure 7.7: Choosing a New GCK Data Folder**

on the bottom shows the name of the current directory being viewed in this list. Pressing the **Select GCK** button in this dialog would tell **GCK** to use the folder named **GCK** as the new “GCK Data” folder. This would mean that the program could not find enzyme tables and other data it might need. To choose the folder named GCK Data in Figure 7.7, you would first have to press the **Open** button to change the list to show the GCK Data directory, and then press the newly named button called **Select GCK Data**, which would appear at the bottom of the new dialog.

## Quit

File > Quit (command-Q/ctrl-Q) will close all windows and exit the Gene Construction Kit. If changes have been made to any of the windows since the last time that window was saved, you will be given the option to save the changes or to discard them (see Figure 7.4, page 7-5).

**Edit Menu**

**Undo**

Edit > Undo (command-Z/ctrl-Z) provides the ability to revert to the state the window was in just prior to your last action, as you are probably familiar with from other applications. The last action can be a cut, a paste or a mouse click followed by typing. The action to be “undone” will be stated in place of the simple Undo (e.g. undo paste, undo style change, undo typing, etc.).

**Cut**

Removes the selected item from the document and places it on the clipboard.

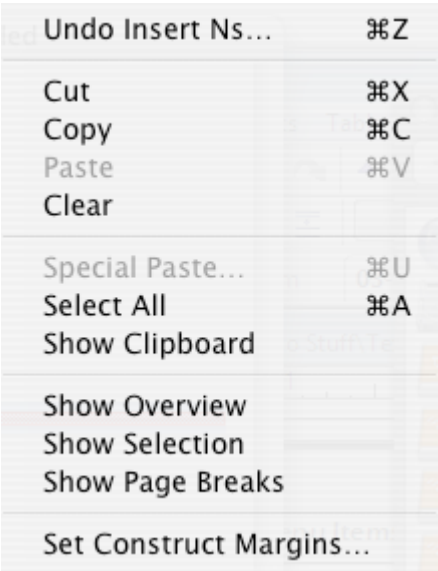
**Copy**

Makes a copy of the selected item and places it on the clipboard.

**Paste**

Edit > Cut (command-X/ctrl-X), Edit > Copy (command-C/ctrl-C), and Edit > Paste (command-V/ctrl-V) function as they do in other applications. Cut will remove the selected item(s) from the window and place it in a temporary holding area, the clipboard. Copy functions similarly to the Cut option, but only places a copy of the selected item(s) in the clipboard — it does not remove anything. Paste will take whatever is in the clipboard and try to place it at the current location of the cursor. Some special situations pertain to cutting, copying and pasting objects in the Gene Construction Kit.

To begin with, it must be kept in mind that when a segment of a construct is selected, it represents a double-strand of DNA. Depending on how the segment was selected and how the ends of the segment are defined, this seg-

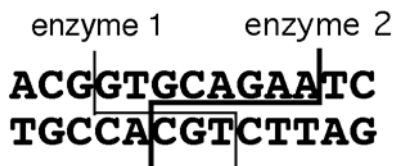


The image shows a screenshot of the 'Edit' menu in a software application. The menu is open, displaying various options. The first option is 'Undo Insert Ns...' with a keyboard shortcut of '⌘Z'. Below this are 'Cut' (⌘X), 'Copy' (⌘C), 'Paste' (⌘V), and 'Clear'. A separator line follows. Then there is 'Special Paste...' (⌘U), 'Select All' (⌘A), and 'Show Clipboard'. Another separator line is present. Below that are 'Show Overview', 'Show Selection', and 'Show Page Breaks'. A final separator line is at the bottom, followed by 'Set Construct Margins...'. The menu has a light gray background and the text is in a standard sans-serif font.

Undo Insert Ns...	⌘Z
Cut	⌘X
Copy	⌘C
Paste	⌘V
Clear	
Special Paste...	⌘U
Select All	⌘A
Show Clipboard	
Show Overview	
Show Selection	
Show Page Breaks	
Set Construct Margins...	

ment might have blunt ends or staggered ends. Double-clicking on a segment in either the graphical view or the sequence view will select the segment between the two nearest “borders”. If no sites are marked, the entire sequence will be selected between style changes. In the sequence view, double-clicking will select a segment of DNA between two marked sites *unless there is a change in font, color, style, size, etc.* In these cases, the selection will stop at the change. This feature allows you to define and display segments of DNA in the sequence view, such as a conserved sequence, that you might not wish to have visible in the graphical view. This facilitates creating informative illustrations of sequence features for presentation while also maintaining the graphical view of the construct. You can have a different formatted sequence listing for each generation of the construct.

There is one situation that has to be dealt with specially. It arises from the



**Figure 7.8: Overlapping Enzyme Sites**

situation shown in Figure 7.8. In this unusual case, two enzymes have recognition sites near each other and their cut sites overlap. If you select this segment by double-clicking or by dragging from cut#1 to cut#2 on the top strand, the top strand will be selected, but *nothing will be selected on the bottom strand*. You can still make font and color changes to the selected (top) sequence, but no changes will be made to the bottom strand. If you now attempt to copy or cut this segment, **GCK** will not let you do so and will present you with an alert box saying that this operation is not possible. The situation is not possible biologically either since as soon as one enzyme makes its cut, the second enzyme recognition site will be gone.

When pasting segments of DNA into sites on other constructs or elsewhere in the same construct, the ends of the segment are examined to see if they are compatible with the site into which the fragment is being pasted. If they are not, you will see the ligation dialog box shown in Figure 3.22, page 3-25 and



be asked to adjust the ends so that the ligation can be done. This parallels what you need to do in the lab. If you have copied a segment of a circular construct to the clipboard and then paste it into a new Construct window, you will get a linear construct containing that stretch of DNA (again, just as if you did the experiment in the lab). If you take that same segment of a circular construct and paste it into the Illustration Window, however, **GCK** realizes that you want it represented as an arc for part of an illustration, and will draw it that way.

The Gene Construction Kit is also smart about the way it puts information in the clipboard — it stores both the picture and textual information. Properly behaving applications (e.g. those that follow Apple's and Microsoft's guidelines) will be able to extract the appropriate information from the clipboard. If you have copied a segment from a graphics view of a construct and then paste it into a text processor, you will paste in the DNA sequence. On the other hand, if you take the same information on the clipboard and paste it into a graphics application, you will see an image of the selected segment and not the sequence. For programs that can display both pictures and text, the program receiving the information from the clipboard makes the decision of how to accept the **GCK** object.

A special case of copying and pasting deals with the copying of marked sites from the Construct window followed by pasting into either a Gel or Illustration window. If you select and copy marked sites in a Construct window and then paste them into a Gel window, **GCK** assumes that you are doing a restriction enzyme digest and will create a lane on the gel showing the fragments that would result from digesting the DNA with those enzymes. If you select and copy marked sites in a Construct window and then paste them into an Illustration window, however, **GCK** assumes that you want to create a **legend**. These two pasting features provide you with an easy way to create gels and to provide legends for constructs in the Illustration Window.

### **Clear**

Edit > Clear will delete whatever is selected. It will not replace what is on the clipboard. Clear is an undoable command.

### **Special Paste...**

Edit > Special Paste (command-U/ctrl-U) was discussed earlier (see discussion around Figure 3.23, page 3-26) and serves to provide special treatment of fragments before they are pasted. Once selected, you get the dialog box shown in Figure 3.23. **Inverting** the fragment before pasting will flip the segment around before pasting it into the target site. In addition to inverting a fragment, options are available to **Trim Ends** or **Fill Ends** of all fragments before pasting (this is equivalent to S1 treatment, and polymerase filling, respectively). The **Leave Ends Alone** option will not treat the ends in any special way — you will see the ligation dialog box if any of the ends are incompatible.

### **Select All**

Edit > Select All (command-A/ctrl-A) will select all of the items of the type you have currently selected (e.g. — if you have selected a marked site, Select All will select all the marked sites). If nothing is selected in the Construct window, Select All will select the entire construct. In the Gel window, Select All will select all the gel lanes. In the Illustration window, Select All will select all the objects in the Illustration.

### **Show/Hide Clipboard**

Edit > Show/Hide Clipboard will either show or hide a window showing the contents of the clipboard. If you are having problems transferring information into or out of **GCK** from/to other applications, examining the clipboard contents can often help sort out where the problem lies.

### **Show Overview**

Edit > Show Overview is active from the Construct and Illustration windows and will shrink the entire window to fit on the screen. The cursor changes to a small magnifying glass. Clicking the magnifying glass at a specific location in the overview window will cause the image to return to normal size and will center the image around the site that was clicked. This action facilitates moving

around large illustrations.

### **Show Selection**

Edit > Show Selection is active in every window except the List window. It will make visible any object that is selected. This might be useful if you have highlighted a sequence in the sequence view of a construct but have scrolled to some other location of the sequence. Choosing Show Selection will make the selected sequence visible.

### **Show/Hide Page Breaks**

Edit > Show/Hide Page Breaks is functional in all windows. It will show a line on the screen indicating the edges of a page. It is particularly useful in the Construct and Illustration windows. Page breaks are automatically shown in the sequence view of a construct.

### **Set Gel/Construct Margins...**

Edit > Set Gel/Construct Margins... will let you define the top, left, bottom, and right margins for Gel and Construct windows and for Gels and Constructs within Illustrations.

## **Window Menu**

### **Stack Windows**

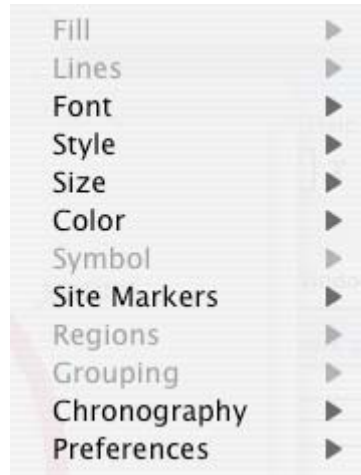
Window > Stack Windows will organize the current set of open windows on the screen. This is useful if you have so many windows open that you cannot locate a particular window. Below the Stack Windows menu item will be a listing of all open windows. The list is organized alphabetically and by window type. Selecting a window name from the list will bring that window to the front.

### **(list of open windows)**

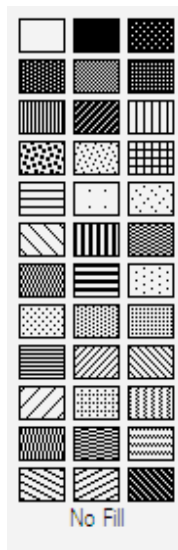
All open windows in **GCK** will appear listed at the bottom of the Window menu. The list will be alphabetical and will contain an icon indicating the type of window.

### Format Menu

The Format menu contains the tools needed to alter the display of information in the various Gene Construction Kit windows. Because of the large number of choices available, all of the items in the Format menu are hierarchical in nature. To view the options in the submenus of the Format menu, simply hold the mouse down on a menu item and a submenu will appear giving you further choices. This is the most logical way to provide access to the large number of choices in an intuitive way; it provides revealed complexity.



Many of the items in the Format menu are available from a number of win-



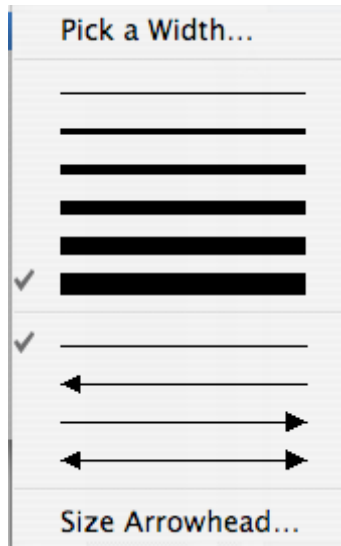
**Figure 7.9: Fill Patterns**

dows. For example, Font is available in all windows except List windows. Fill is available in Construct and Illustration windows. Symbol is available only in the Construct window, and Grouping is available only in the sequence view of the Construct window.

### **Fill**

The Format > Format Fill submenu provides the ability to define the pattern to be used for filling selected objects. The choice of fill patterns is shown in Figure 7.9, page 7-17. Clicking on any pattern with the mouse will fill the selected object with that pattern. This can be used to define segment patterns in the graphic view of constructs or the fill pattern for ellipses, rectangles, and round rectangles in the Illustration window. Selecting none will produce a transparent object in the Illustration or Construct windows (other objects can be seen through objects that have a “none” fill pattern). Fill can be used in conjunction with colors to produce a large variety of effects. This is especially useful if your constructs are used on both black and white and color monitors.

### Lines



**Figure 7.10: Lines Menu**

#### ...Pick a Width

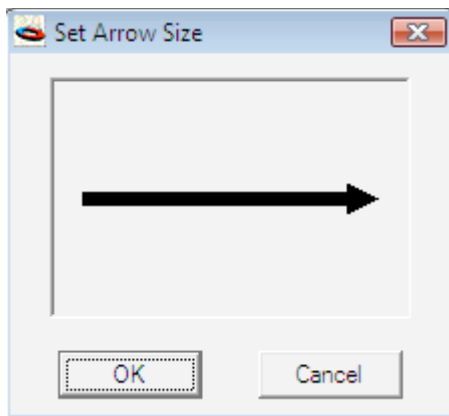
#### ...(predefined line thicknesses)

#### ...(predefined directions)

#### ...Size Arrowhead...

The **Format > Lines** submenu provides control over the attributes of lines and arrows, which are a special kind of line. As shown in Figure 7.10, there are several parts to the **Lines** submenu. This submenu is available when a line is selected in the **Illustration** window or a segment or region is selected in the **Construct** window. Clicking on one of the predefined line thicknesses will change the selected line to the thickness of the line you clicked on. You can specify a line size not present in the submenu by choosing **Format > Lines > Pick a Width...**. Clicking on the bottom part of the submenu will allow you to place arrowheads at one or both ends of the selected line. Choosing a direction to the line that represents a translated region in a construct will specify the cod-

ing direction for protein sequences; left to right (or clockwise) displays the protein coded for on the 5' to 3' (top) strand, while right to left (counterclockwise) shows the protein coded for on the 3' to 5' (bottom) strand. The size and shape of the arrowhead can be adjusted by choosing the `Format > Lines > Size Arrowhead...` option. This will present Figure 7.11. The arrowhead can be



**Figure 7.11: Size Arrowhead**

altered by either dragging a corner of the arrowhead to the desired location, or by clicking the mouse at the new location you desire for the side of the arrowhead. The thickness of the horizontal line in the dialog is the same as the thickness of the line being edited.

## **Font**

The standard `Format > Font` menu lists all of the available fonts from your system. It operates in the same way Font menus work in most other applications. This submenu (along with the `Style`, page 5-21 and `Size`, page 5-21) is available anytime that text is selected in the `Illustration` window or the `Construct` window (either the graphics or sequence views) and will function to alter the display of the selected text. In the `Gel` window, setting these attributes will set the display of the legend text and the text of digests when viewed as a table.

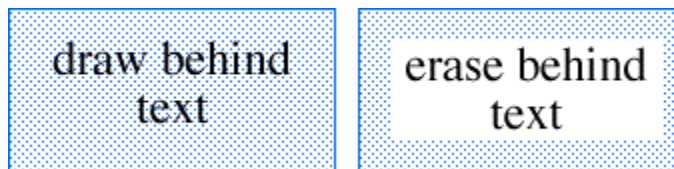
*In this submenu and in all other submenus, the command (Mac)/control (windows) key equivalent to selecting the menus is to press the shift key*



(indicated by the symbol  $\uparrow$ ) in combination with the command/control key (indicated by the symbol  $\wedge$ ). Thus,  $\uparrow$ -command-B/ $\uparrow$ -ctrl-B means to press the shift key, the command/ctrl key, and the B simultaneously. In the Gene Construction Kit, regular menu keyboard equivalents are chosen with the command key and submenu keyboard equivalents are chosen with command and shift keys.

### Style

The Format > Style menu contains the standard style options that are found in other programs including: Plain Text ( $\uparrow$ -command-P/ $\uparrow$ -ctrl-P), Bold ( $\uparrow$ -command-B/ $\uparrow$ -ctrl-B), Italic ( $\uparrow$ -command-I/ $\uparrow$ -ctrl-I), Underline ( $\uparrow$ -command-U/ $\uparrow$ -ctrl-U), Outline ( $\uparrow$ -command-O/ $\uparrow$ -ctrl-O), Shadow ( $\uparrow$ -command-S/ $\uparrow$ -ctrl-S), Condense, and Extend. Left Justify ( $\uparrow$ -command-L/ $\uparrow$ -ctrl-L), Center ( $\uparrow$ -command-C/ $\uparrow$ -ctrl-C), and Right Justify ( $\uparrow$ -command-R/ $\uparrow$ -ctrl-R) are available when text is being edited in the Illustration window. In addition, there are a few other styles that are unique to GCK. When working in a sequence view of a construct, Case as Typed, Upper Case, and Lower Case are available. Note that these are styles just like Bold — they modify the way a character is displayed but do not actually change the character. The last two



**Figure 7.12: Erase Behind Text**

items in this menu also apply to text in the Illustration window. Erase Behind Text will cover up any object that lies behind the text itself. Draw Behind Text will allow the background object to be drawn as shown in Figure 7.12

### Size

The Format > Size submenu allows you to set the size of the currently selected text.

## **Color**

**...Pick a Color...**

**...Add a Color...**

**...Remove a Color...**

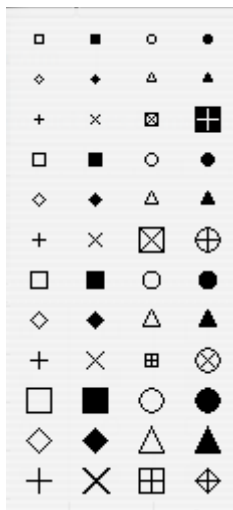
**...(predefined colors)**

**...(user defined colors)**

The Format > Color submenu allows you to define the color of a selected object. You can define an object's color even on a black and white system by choosing the desired color from the Color submenu. You can also use Add a Color... and Remove a Color... to customize the Color menu. The new colors that you add can be defined using the color picker, or by selecting the desired color from a visible window. Color can apply to text, construct segments, marked sites, regions, and symbols in the Construct Window; and to text, lines (arrows), rectangles, round rectangles, and ellipses in the Illustration window. By assigning meaningful names to colors when you create them and add them to the menu (e.g. transcript or coding), it is easy to maintain consistency among all constructs and illustrations.

## **Symbol**

Format > Symbol submenu will display the graphical submenu shown in Figure 7.13, page 7-23. Selecting a symbol will change the currently selected marked site(s) in the Construct window to the selected symbol from this submenu. Note that symbols may be chosen even if the site is being displayed by name. Be careful when changing symbols, however, because when you double-click on a symbol in the Construct window all the sites showing that same symbol will be selected – even if they represent different enzymes. Similarly, when double-clicking on a site that is displayed by name, all sites with that name will be selected.



**Figure 7.13: Symbol Menu**

### **Site Markers**

The **Format > Site Markers** submenu provides access to specifying characteristics of how the site markers will be displayed. In general, a site marker can be displayed as either a symbol or as text. When shown as text, the name of the marker and/or the position of the cut site can be displayed. The items in this menu all deal with determining how the site is marked on the construct, in both sequence and graphics display modes.

#### **...Manual Arrangement**

This option complements the Automatic Arrangement, page 5-23 in the previous section. When **Format > Site Markers > Manual Arrangement** is chosen, any site marker that is selected will not be automatically moved when the window is resized or “zoomed”. It is a property of each site marker.

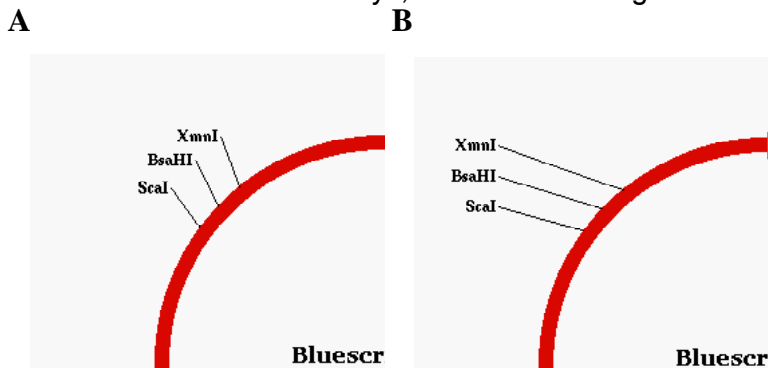
#### **...Automatic Arrangement**

When a site marker is selected and then **Format > Site Markers > Automatic Arrangement** is chosen, **GCK** will automatically move that marker to a position in which it does not overlap any other site markers. If the Construct window is

resized or “zoomed” in or out, these site markers will be automatically placed in non-overlapping positions. This is very useful for locations along the DNA in which there may be many overlapping sites, such as polycloning regions. Note that Automatic Arrangement is a **property** of the site marker – almost like a specific style. This property will stay with the site marker until you either choose Manual Arrangement, page 5-23, or drag the site marker with the mouse.

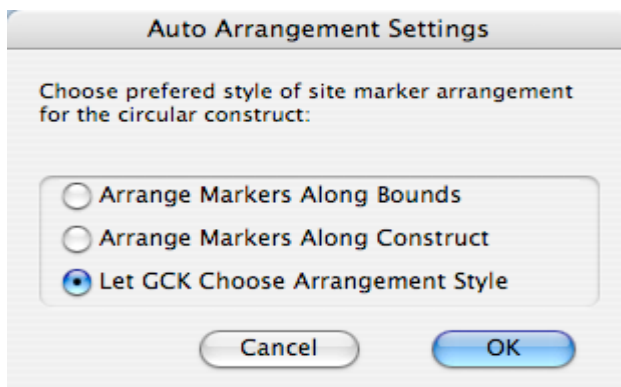
### **...Auto Arrangement Settings...**

Format > Site Markers > Auto Arrange Settings... allows you to define the way in which site markers are arranged for circular constructs. When site markers are displayed using automatic arrangement with circular constructs, the automatic arrangement can be done in two ways, as shown in Figure 7.14. In part A,



**Figure 7.14: Autoarrangement Styles**

the site names are arranged along the construct. In part B, the site names are arranged along the window borders. You can choose which way to display the site markers using the dialog shown in Figure 7.15, page 7-25. Letting **GCK Choose** (the last radio button) will allow GCK to change between different methods depending on the circumstances of the view. This can be particularly useful when zooming in to a high magnification. In this case, arranging the site markers along the construct makes sense in the magnified view, while it might be preferable to view them vertically arranged when the entire construct is showing. Note that this menu item is only available when a construct is circular.



**Figure 7.15: Auto-Arrangement Settings**

### **...Show Site as Text**

### **...Show Site as Symbol**

These two items in the `Format > Site Markers` submenu allow you to set the display of any *selected* marked site(s) to either symbols or text. You can change between these two formats at any time and can restrict the change to an individual site marker.

### **...Show Name Only**

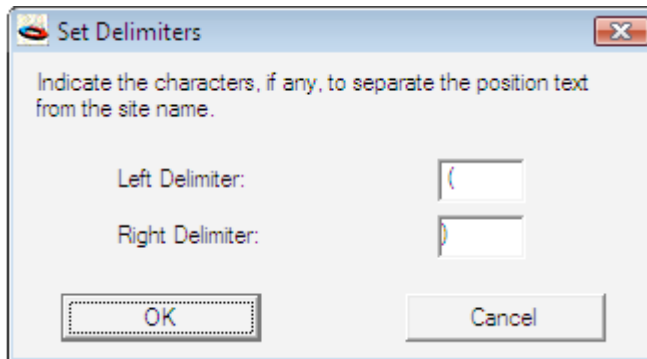
### **...Show Position Only**

### **...Show Name and Position**

`Format > Site Markers > Show Name Only`, `Format > Site Markers > Show Position Only`, and `Format > Site Markers > Show Name and Position` allow you to specify the text to be shown when the site marker is viewed as text. The position will be indicated in the site marker label as it is defined in `Set Position Delimiters...`

### **...Set Position Delimiters...**

`Format > Site Markers > Set Position Delimiters...` submenu item allows you to set the characters that will be used at the left and at the right of the position indicator



**Figure 7.16: Set Position Delimiters**

text. Using the dialog box shown in Figure 7.16, you enter the text you want to be placed to the left and to the right of the actual position number. In this case the site name will be followed by a space and a left parenthesis; after the position number there will be a right parenthesis. The site marker might look like: “HindIII (29)”.

### **...Attach at Left**

### **...Attach at Center**

### **...Attach at Right**

### **...At Nearest Horizontal**

These four items define how the line extending from the construct to the site marker will attach to the site marker text in the left <-> right (horizontal) direction. Choosing **Format > Site Markers > Attach at Left** forces the line to connect to the left edge of the site marker text. Choosing **Format > Site Markers > Attach at Center** forces the line to connect to the horizontal center of the site marker text. Choosing **Format > Site Markers > Attach at Right** forces the line to connect to the right edge of the site marker text. Choosing **Format > Site Markers > At Nearest Horizontal** allows **GCK** to choose the best location for the attachment for you. This automatic attachment allows you to move the site marker around and still have the line connected to the marker text at an aesthetically pleasing location.

### **...Attach at Top**

### **...Attach at Middle**

### **...Attach at Bottom**

### **...At Nearest Vertical**

These four items define how the line extending from the construct to the site marker will attach to the site marker text in the top <-> bottom (vertical) direction. Choosing Format > Site Markers > Attach at Top forces the line to connect to the top edge of the site marker text. Choosing Format > Site Markers > Attach at Middle forces the line to connect to the vertical center of the site marker text. Choosing Format > Site Markers > Attach at Bottom forces the line to connect to the bottom edge of the site marker text. Choosing Format > Site Markers > At Nearest Vertical allows **GCK** to choose the best location for the attachment. This automatic attachment allows you to move the site marker around and still have the line connected to the marker text at an aesthetically pleasing location.

## **Regions**

### **...Move Region Outwards/Upwards**

### **...Move Region Inwards/Downwards**

Once Regions have been created, it is sometimes desirable to rearrange them. This can be accomplished using the Format > Regions > Move Region Outwards/Upwards (⌘-command-+/⌘-ctrl-+) and Format > Regions > Move Region Inwards/Downwards (⌘-command--/⌘-ctrl--) menu items. Selecting a Region and then choosing one of these options will move the selected region to the outermost/top or innermost/bottom location.

### **...Show As Protein**

### **...Don't Show As Protein**

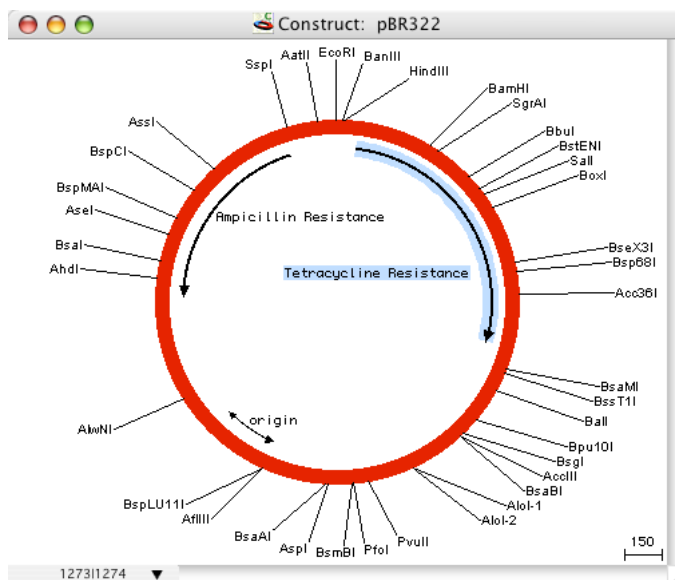
These two menu items are available when one or more regions are selected.

In the graphical view of a construct, only an arrow will be drawn, but in the sequence view, **Format > Regions > Show As Protein** will cause the region to be drawn as a series of amino acids translated from the corresponding DNA in the orientation indicated by the arrow direction. **Format > Regions > Don't Show As Protein** will convert any translated regions back to plain graphic arrows.

### **...Show Region Names**

### **...Hide Region Names**

Selecting a region and then choosing **Format > Regions > Show Region Names** will place the name of the region near the graphic of the region as seen in Figure



**Figure 7.17: Region Names**

7.17, page 7-28. This option is not available in the sequence view of a construct. Note that when a region graphic is selected, the region name is also selected (and vice versa). This makes it easy to see which name corresponds to which graphic when the names are moved around.

### **...Reset Region Name Positions**

The **Format > Regions > Reset Region Name Positions** menu item will return a region



name to its default position along the region to which it corresponds.

### **Grouping**

The Format > Grouping submenu is only enabled when the sequence view of a Construct window is active. It allows you to define precisely the way in which the DNA sequence is displayed (and printed). Grouping refers to the number of nucleotides that are placed adjacent to each other without any intervening spaces.

#### **...No Grouping**

Format > Grouping > No Grouping (⌘-command-J/⌘-ctrl-J) will allow the sequence to wrap around based on the width of the window.

#### **...Group by Threes**

Format > Grouping > Group by Threes (⌘-command-F/⌘-ctrl-F) is useful for displaying the DNA sequence above a translated amino acid sequence.

#### **...Group By Tens**

Format > Grouping > Group By Tens (⌘-command-T/ctrl) provides a useful format for long sequence listings. Having the sequence displayed in groups of 10 makes it easy for the eye to follow sequence positions.

#### **...Insert Line Break**

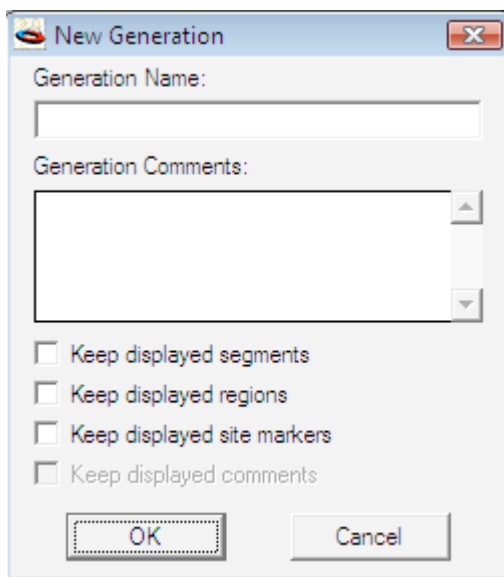
#### **...Remove Line Breaks**

Format > Grouping > Insert Line Break will force the character immediately after the insertion point to begin a new line. If you have selected a segment of the sequence and then choose Insert Line Break, GCK will place the selected sequence in its own formatting segment. This means that the selection will start on a new line, and the end of the selection will be the end of a line (the next formatting segment will start a new line). Format > Grouping > Remove Line Breaks will remove line breaks from the selected segment of DNA. If no seg-

ment is selected, then all line breaks will be removed. The Grouping submenu should be used in conjunction with the Display submenu of the Construct menu to precisely define the layout of the sequence.

### **Chronography**

The Format > Chronography submenu provides you with access to GCK's



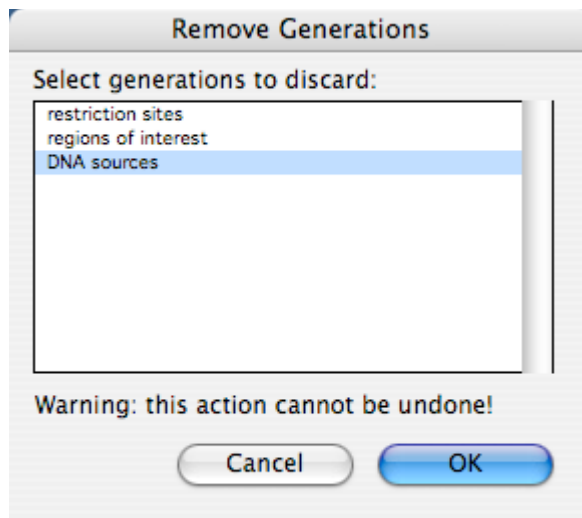
**Figure 7.18: New Generation Dialog**

archiving feature that can track all the steps in the synthesis of a construct and present alternative views of the information in the construct. The uses of Chronography were covered earlier in Tutorial 7:Chronography – Tracking Cloning History, page 2-30.

### **...Start New Generation...**

The appearance of a construct can be changed without losing the current display by defining a new “generation” at any time. The information defined in a generation consists of the line thickness, fill pattern, color and direction of each specific segment of the construct, the marked sites, and the regions (i.e. all aspects of the appearance of the construct). Using the Format > Chronography > Start New Generation (⌘-command-G/⌘-ctrl-G) option will define a new generation.

After selecting this option, you will see a dialog box allowing you to keep (or not) features of the current display in the new generation that is being defined (see Figure 7.18). Checking a box will keep that information displayed in the new generation you are creating. You can provide a name for each generation to help you in tracking all the steps and alternative views in your construct. You can also enter any comments you wish for the generation. This might be a convenient spot to store information about the construction process.



**Figure 7.19: Discarding Generations**

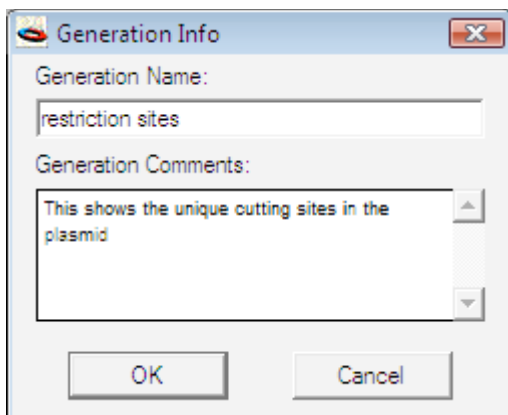
### **...Discard Generations...**

Format > Chronography > Discard Generations allows you to dispose of one or more generations that you no longer need. Disposing of generations will make the file size smaller. The dialog is shown in Figure 7.19. To delete more than one generation, you can hold down the shift key while you make multiple selections from the list with the mouse (shift-clicking). Note that this operation is not undoable. Once a generation is deleted, it can never be recovered.

### **...Get Info For Current Generation**

Each generation can have information stored with it. The information can be

entered when the generation is first created (Figure 7.18, page 7-30), but once the generation is created you can access and edit the information by choosing this menu option. It will bring up Figure 7.20, page 7-32.



**Figure 7.20: Generation Information**

### **...Show Previous Generation**

### **...Show Next Generation**

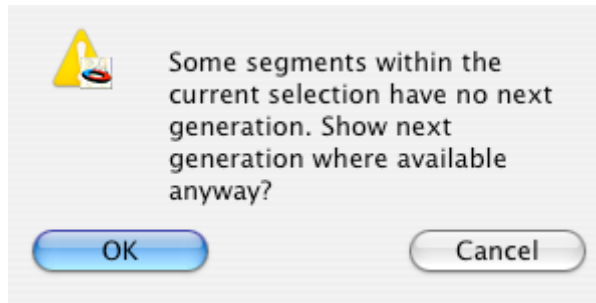
Format > Chronography > Show Next Generation ( $\uparrow$ -command->/ $\uparrow$ -ctrl->) and Format > Chronography > Show Previous Generation ( $\uparrow$ -command-</ $\uparrow$ -ctrl-<) will be selectable only when appropriate (when there is a more recent or more ancient generation to be shown). If your selection in the construct contains segments in more than one generation, you will be shown the alert box in Figure 7.21, page 7-33. In general, if the Gene Construction Kit cannot figure out what you want, you will be shown alert boxes like Figure 7.21.

### **...Show Most Recent Generation**

Format > Chronography > Show Most Recent Generation ( $\uparrow$ -command-A/ $\uparrow$ -ctrl-A) will display the most recent generation for each selected segment.

### **...Show Most Ancient Generation**

Format > Chronography > Show Most Ancient Generation ( $\uparrow$ -command-Z/ $\uparrow$ -ctrl-Z) will display



**Figure 7.21: Ambiguous Generations**

a generation corresponding to the oldest generation in existence for the selected segment of the construct.

### **...(user defined generation names)**

The last section of this menu will contain the names of the various generations that you have defined. You can show the generation by selecting its name from this menu.

## **Preferences**

The Preferences Submenu allows you to set the default values to be used when new windows are created by GCK.

A **Construct**, **Illustration**, or **Gel** window can be opened from the Preferences submenu. To set a default characteristic for any given window, just define the attribute as you would in the normal window. For example, in the Construct preferences window, to define the default segment color, double-click on the segment to select it, then select the desired default color from the **Format > Color** submenu. To change the font for marked sites, select the site marker and then choose the desired default font from the **Font** submenu of the **Format** menu.

### **...Default Construct**

For the Construct window, you can set the following default attributes for graphical display: margins, linear or circular construct, segment thickness,

segment fill pattern, segment color, segment direction (arrowheads), display site as text or symbol, auto-arrange or manual arrange placement of site markers, site marker font, site marker font size, site marker font style, site marker font color, site marker text justification, site marker symbol, and site marker color. For sequence display, you can also set the grouping, line spacing, show/hide positions, font size, style, and color.

### **...Default Gel**

For the Gel window, you can set the following attributes: margins, legend/table font, legend/table font style, legend/table font size, threshold value, and which items should be displayed (legend, runoff count, size standards).

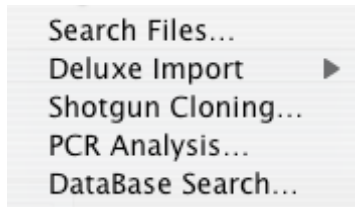
### **...Default Illustration**

For the Illustration window, you can set the following attributes: object color, object fill pattern, object line thickness, line color, line thickness, line direction (arrowheads), text font, text size, text style, text justification, and text color.

### Tools Menu

#### Search Files...

File > SearchFiles... was discussed in Tutorial 8:Finding Comments and File Searching, page 2-38. File searching provides you with the ability to search Gene Construction Kit files on disk for matches with key words in comments or for matches with DNA sequences. Please see the tutorial for details on how to use this option.



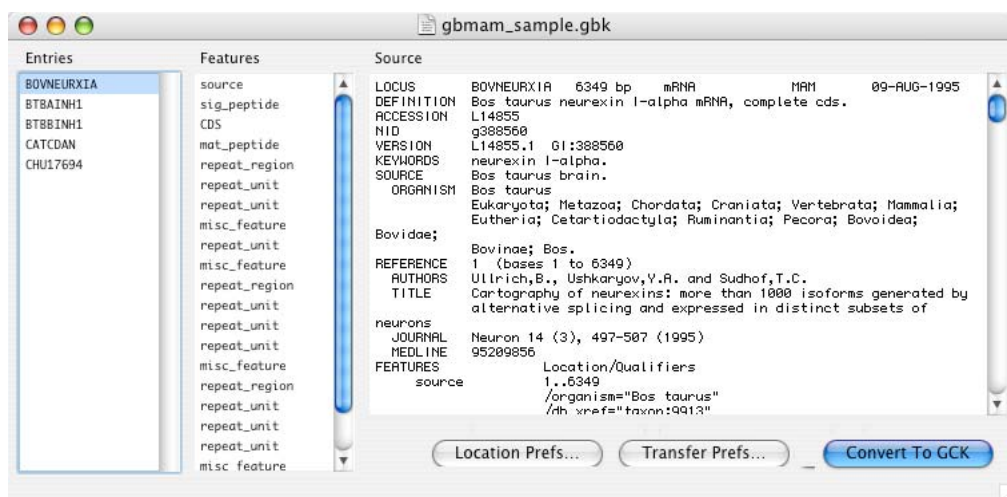
If you are conscientious about entering information with your constructs, SearchFiles can be used to keep track of storage locations for different constructs. For example, if in the general comments associated with every construct, you include the storage location for that particular construct, you could then use File Searching to locate all the actin constructs in freezer#2. Therefore, **GCK** can serve as a simple database management system for constructs providing both storage and sequence information.

#### Deluxe Import

There are six submenus under this menu. **GCK** has the ability to take each item in a GenBank features table and convert it to a specific **GCK** feature. The menu items in this submenu facilitate that process. Tutorial 13:Importing GenBank Sequence Files Using Deluxe Importing, page 2-64 discusses the process in some detail.

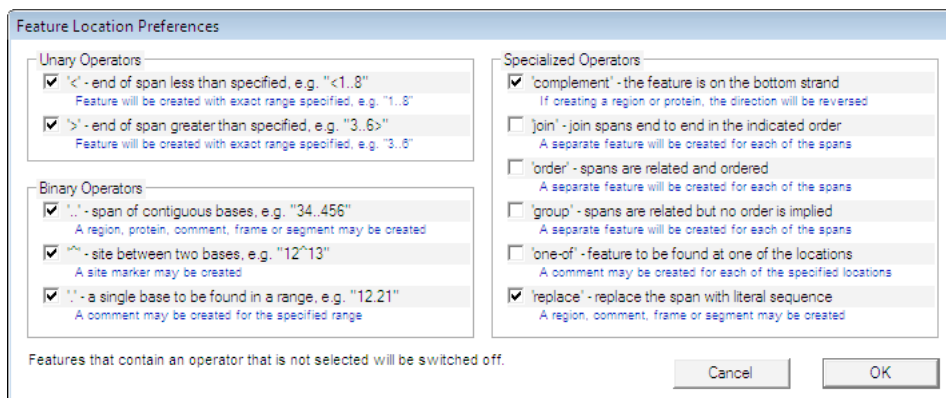
#### ...Open Sequence(s) File

If you already have a GenBank or EMBL file downloaded to your hard disk, choosing this option will allow you to convert it to a **GCK** file. When opening, you will see the dialog shown in Figure 7.22. This particular file has five different sequences in it, the first of which is selected. The **Location Prefs** button allows you to define how different locations specified in the GenBank file are to be converted into **GCK** locations. The choices are shown in Figure 7.23, page 7-36. Using this dialog, you can specify exactly which options



**Figure 7.22: Convert File Dialog**

you want to use and how they are to be deciphered. The **Transfer Prefs** button results in Figure 7.24. Once all of these parameters have been set,



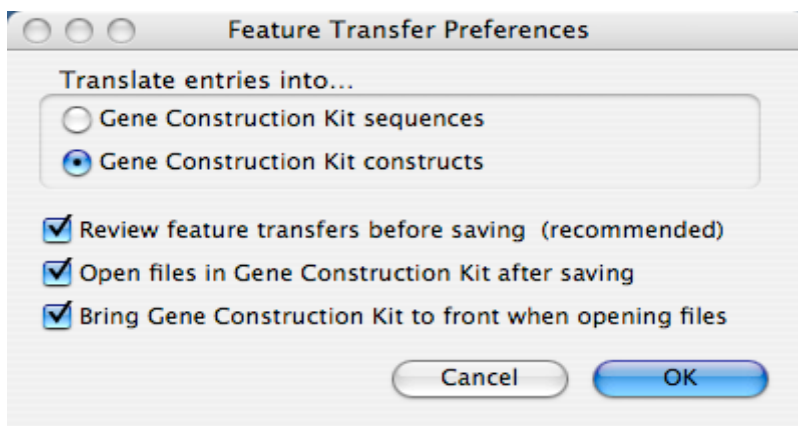
**Figure 7.23: Feature Location Preferences**

GCK will remember them so you will not have to set them more than once. Importing the file uses the Conversion Defaults as illustrated in Figure 2.68, page 2-65.

### ...Retrieve Sequence

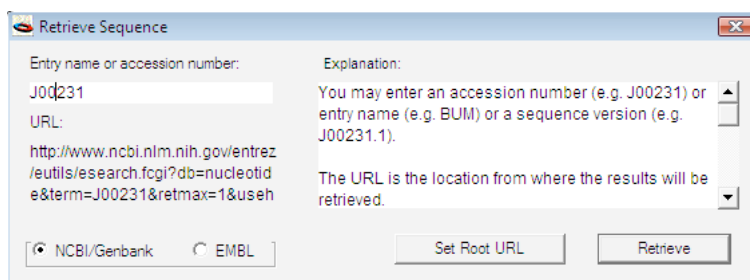
The Retrieve Sequence menu option allows for the retrieval of specific





**Figure 7.24: Feature Transfer Preferences**

sequences from either GenBank or EMBL databases. As shown in Figure 7.25, page 7-37. You can enter either an accession number or a keyword



**Figure 7.25: Retrieve Sequence Dialog**

description for the sequence.

If you have a local database that is set up using the GenBank format, you can choose to search that database by pressing the **Set Root URL** button. This feature is useful if you are searching private databases. Note that the target database must be in the same format as either GenBank or EMBL databases in order for the retrieval to work.

### **...Search GenBank**

This option was discussed in Tutorial 13:Importing GenBank Sequence Files Using Deluxe Importing, page 2-64. You can search GenBank by entering a keyword in any number of categories. The categories are accessible using a

popup menu as shown in Figure 2.71, page 2-68.

### **...Conversion Defaults**

The conversion defaults specify how each feature in the GenBank table will be converted into a **GCK** feature. As discussed in Tutorial 13:Importing GenBank Sequence Files Using Deluxe Importing, page 2-64, you can choose any number of ways to have GenBank features represented in the final **GCK** construct you create. The Conversion Defaults window is shown in Figure 2.68, page 2-65. The correspondences you specify in the Conversion Defaults are used to define how any particular conversion will be done. However, each time you perform a conversion, you have the option of overriding the default conversions individually or as a whole.

### **...Save Conversion Defaults**

Once you have defined a set of Conversion Defaults that you like, you can save them in a file to be recalled later if needed. If you have several different sets of **Conversion Defaults**, you can create several different files using this option.

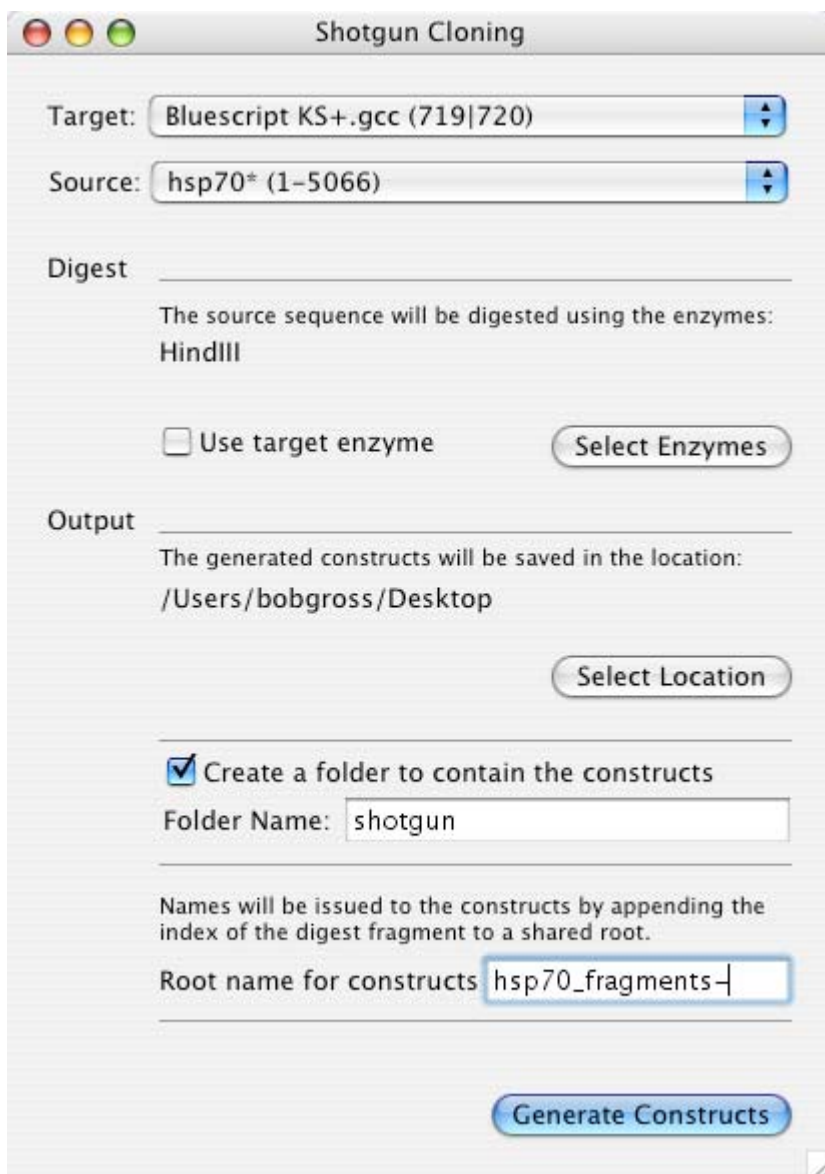
### **...Restore Conversion Defaults**

This menu item allows you to open any of the files you may have saved under **Save Conversion Defaults** to be used as the new set of defaults.

### **Shotgun Cloning**

Shotgun cloning refers to the process of fragmenting a DNA segment and then placing each fragment into a restriction site of a target vector. Gene Construction Kit allows you to choose any enzyme to digest your DNA and to use as a target site within the vector DNA. Thus, there are two DNAs involved in this process: the source DNA and the target DNA. These DNAs and the enzymes used for the digest as well as the details of the output files are all available within the Shotgun Cloning window.

The Target and Source popup menus will display any open constructs that



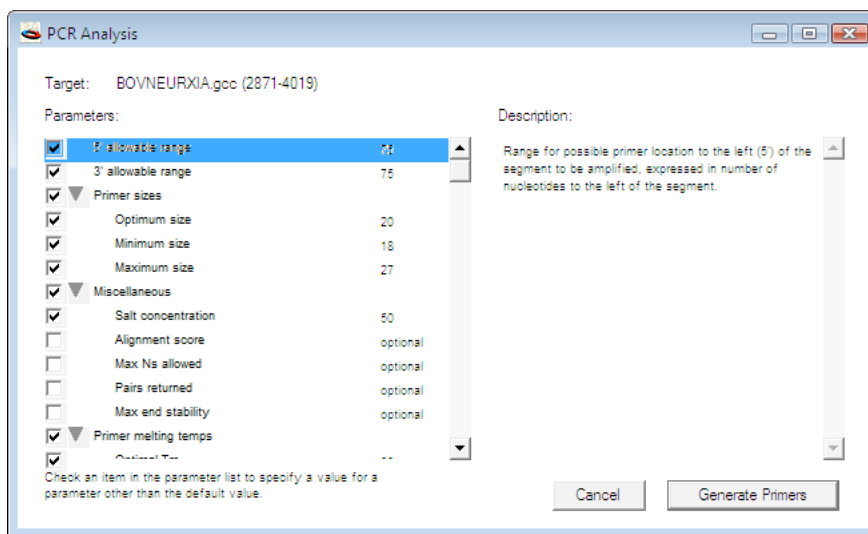
The image shows a software window titled "Shotgun Cloning". It contains several input fields and buttons. At the top, there are three colored window control buttons (red, yellow, green). The "Target:" field contains "Bluescript KS+.gcc (719|720)". The "Source:" field contains "hsp70\* (1-5066)". Below these, the "Digest:" section shows "The source sequence will be digested using the enzymes: HindIII". There is a checkbox labeled "Use target enzyme" which is unchecked, and a button labeled "Select Enzymes". The "Output:" section shows "The generated constructs will be saved in the location: /Users/bobgross/Desktop" and a button labeled "Select Location". Below that, a checkbox labeled "Create a folder to contain the constructs" is checked. The "Folder Name:" field contains "shotgun". A text block explains: "Names will be issued to the constructs by appending the index of the digest fragment to a shared root." The "Root name for constructs" field contains "hsp70\_fragments-". At the bottom right is a large button labeled "Generate Constructs".

**Figure 7.26: Shotgun Cloning window**

you have, and if there are site markers selected, they will be used as the default enzyme(s). If the construct file is not open, you can choose **Browse** from the popup menu to select the construct you want. GCK will automatically

open any chosen file. If **Use Target Enzyme** is checked, the selected site marker in the target DNA will be used as the default enzyme. If it is not checked, you can choose any enzyme you wish by pressing the **Select Enzymes** button. Note that the enzymes you choose need to have an end that is compatible with the target site. Incompatible ends will not be ligated into the target site. **Select Location** will allow you to choose a folder (or create a folder) that will be the home of the newly created files. You can type the name for a new folder in the *Folder Name* field. Finally, the *Root name for constructs* field will allow you to specify the “prefix” that will be used for all of the files generated. Each of the generated files will start with the prefix text and will be sequentially numbered. Press **Generate Constructs** to complete the process.

## PCR Analysis



**Figure 7.27: PCR Analysis Window**

Polymerase Chain Reaction (PCR) is a method used to amplify small DNA segments. GCK uses code developed at the Whitehead Institute with their permission. This code is called Primer 3 and is one of the more highly thought of PCR analysis codes. You can see a practical application of how to use the PCR analysis in Tutorial 16:PCR Analysis, page 2-74.

When you initiate the PCR analysis you will see Figure 7.27, page 7-40. There are a number of parameters that you can define as listed in the *Parameters* list on the left side of the window. Clicking on any item in the list will show a description of the item on the right side of the window in the *Description* field. Once you press the **Generate Primers** button, GCK will return a list of possible primer pairs (Figure 2.80, page 2-76). Choose a pair and press the **Show Info** button to see details about how the various attributes of the primer pair scored. Once a pair is selected, you can add it to the construct by pressing **Add Primer** button. Once you are finished adding the primers, press the **Done** button to close the window. Any primers that you added to the construct are displayed as pairs of opposite facing indicators, one on the top and one on the bottom of the construct DNA. You can click on a primer pair, Get Info..., and see the same information that was available in the Show Info window.

### **Database Searching**

There are many databases available over the Internet which contain information relevant to biologists. For details on how to use the DB searching capabilities, see Tutorial 18:Database Searching, page 2-81. The basic idea is that you can provide keywords, protein or DNA sequences to GCK and the program will send that information to one or more database sites on the web.

The searching is managed through a DB searching engine on Textco BioSoftware's web server. The DB server accepts either DNA, protein or keyword information and sends out queries to various web sites. The DB server contains information on how to format queries for different sites and formats the query information appropriately for each site. If the site changes how it wants new queries formatted, the DB server will be updated negating the need to update individual applications. This arrangement also means that we can update the DB server with new sites and remove obsolescent sites without necessitating updating all of the GCK clients.

If you have suggestions for new sites that you feel would be of general interest, please let us know about them by sending us an e-mail at [info@textco.com](mailto:info@textco.com).

## Construct Menu

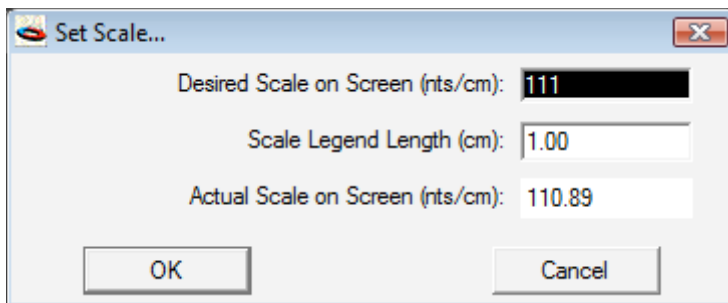
The Construct menu provides complete control over the appearance of a construct. You can choose to show or hide certain features of the display and you can add regions and marked sites as well. Both the graphics and sequence views of the Construct window are controlled from the Construct menu. Essentially all editing of constructs is done in the Construct window using the tools provided in the Construct and Format menus.

## Magnification

This menu will be available whenever a construct is being viewed graphically.

### ...Set Scale

Construct > Magnification > Set Scale allows you to precisely set the scale at which the construct will be drawn. Selecting this menu item will bring up Figure



**Figure 7.28: Set Scale Dialog**

7.28. Since the screen is limited in its resolution to an integral number of pixels/inch on the screen (typically 72 pixels per inch), GCK will show you the actual scale on screen in addition to the value you entered. The *scale legend*

*length* refers to the length of the bar that will be drawn in the window to indicate the scale.

Once you have defined a specific scale with this dialog, you are telling **GCK** that you want to see the construct represented at that particular scale. Therefore, when you resize the construct window, the scale will not automatically change. If you shrink the window, the scroll bars will become active to allow you to navigate around the construct. To have the construct automatically change size when the window is resized, choose “Fit To Window”, page 7-43.

### **...Zoom In**

### **...Zoom Out**

Selecting Construct > Magnification > Zoom In ( $\uparrow$ -command-)]/ $\uparrow$ -ctrl-]) will magnify the view and center it on the location of the insertion point (or the selection). It will provide a 2-fold magnification with each iteration of the command. Construct > Magnification > Zoom Out ( $\uparrow$ -command-[/ $\uparrow$ -ctrl-]) will decrease the magnification by a factor of two with each iteration to allow more of the construct to be seen.

### **...Fit To Window**

Construct > Magnification > Fit To Window ( $\uparrow$ -command-W/ $\uparrow$ -ctrl-W) proportions the construct to fit nicely within the window. If Fit To Window is chosen, the construct will be redrawn and rescaled each time the window is resized. With any of the other options, the construct will not be rescaled if the window size is changed. To keep the construct drawing at a specific scale, choose “Set Scale”, page 7-42.

### **...Fit To Printer Page**

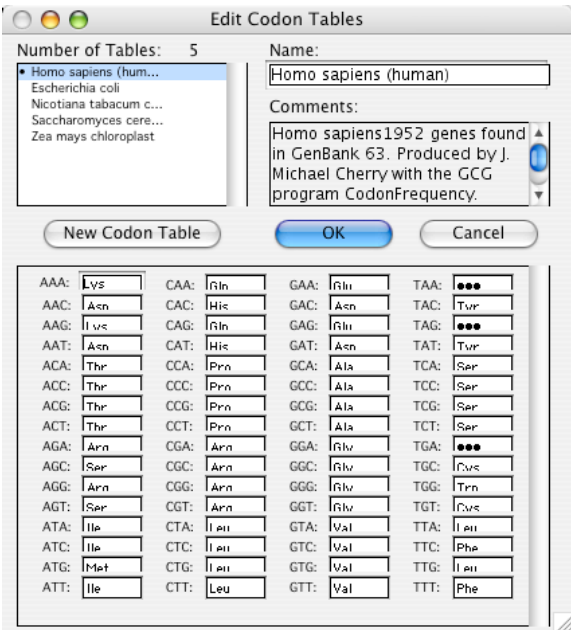
Construct > Magnification > Fit To Printer Page will resize the drawing of the construct to fill up a single printed page, as defined in File > Page Setup.... You can view the off-screen parts by using the scroll bars on the right and bottom of the

Construct window. To make everything visible again, choose Construct > Magnification > Fit to Window.

## Features

### ...Edit Codon Tables...

Choosing Construct > Features > Edit Codon Tables... will bring up Figure 7.29. You



**Figure 7.29: Edit Codon Tables**

will see a list of all the available codon tables in your **GCK Data** folder. In Figure 7.29, there are only 4 tables listed<sup>a</sup>. In the bottom part of this window is a table containing all the codons and the amino acids they code for using the currently selected table. The selected table will be bulleted in the list on the left [**Standard (human)** in this case], its name will be shown in the *Name* box and any comments related to the table will be shown in the *Comments* box. The **OK** button will accept any changes you have made in the table and

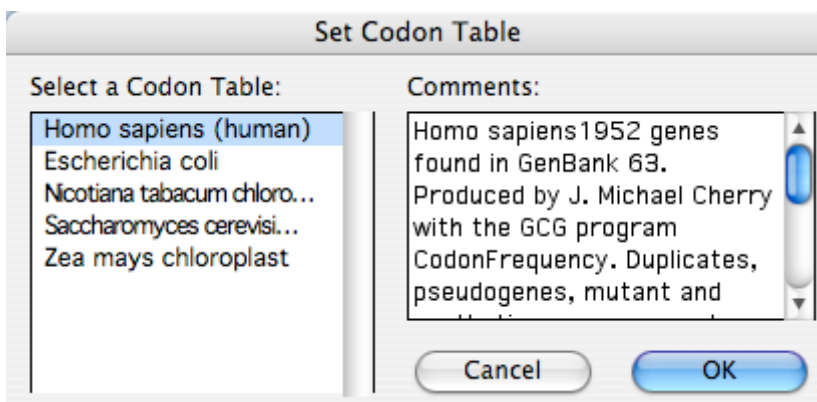
a. If you have codon tables you have generated in the Gene Inspector®, Textco's sequence analysis and molecular biologist's notebook application, they can be placed directly into the GCK Data folder to be used from within **GCK**.



will close the window. The **Cancel** button will close the window without saving any changes. Choosing **New Codon Table** will copy the codon definitions currently visible and create a new table to name and modify. Any new codon tables you create will automatically become available when needed elsewhere in **GCK** (e.g. see “Find Open Reading Frames...”, page 7-46). The new tables will be created in the GCK Data folder.

### **...Choose Codon Table...**

When **GCK** translates a DNA sequence or is asked to find ORFs (“Find Open Reading Frames...”, page 7-46), it uses a codon table to do the translations. The codon table to be used as the standard table can be set here. A

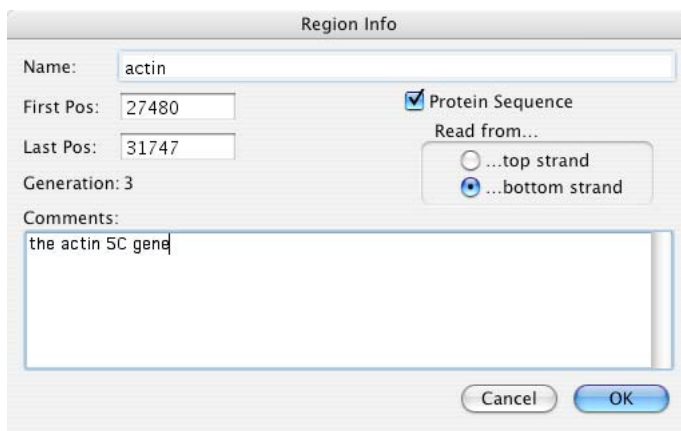


**Figure 7.30: Selecting a Codon Table**

dialog like the one shown in Figure 7.30 will be shown. To choose a default codon table, select the table in the left list and press **OK**. Pressing **Cancel** will close the dialog without changing your previous choice of default table. Some of the supplied tables have comments associated with them. If you decide to add your own table, they can have comments as well.

### **...Make Region...**

Construct > Features > Make Region (command-R/ctrl-R) will create a region corresponding to the selected segment of the construct or sequence. If no DNA segment is selected, you will have to type in the end points of the region in the dialog box shown in Figure 7.31, page 7-46. Once created, the region

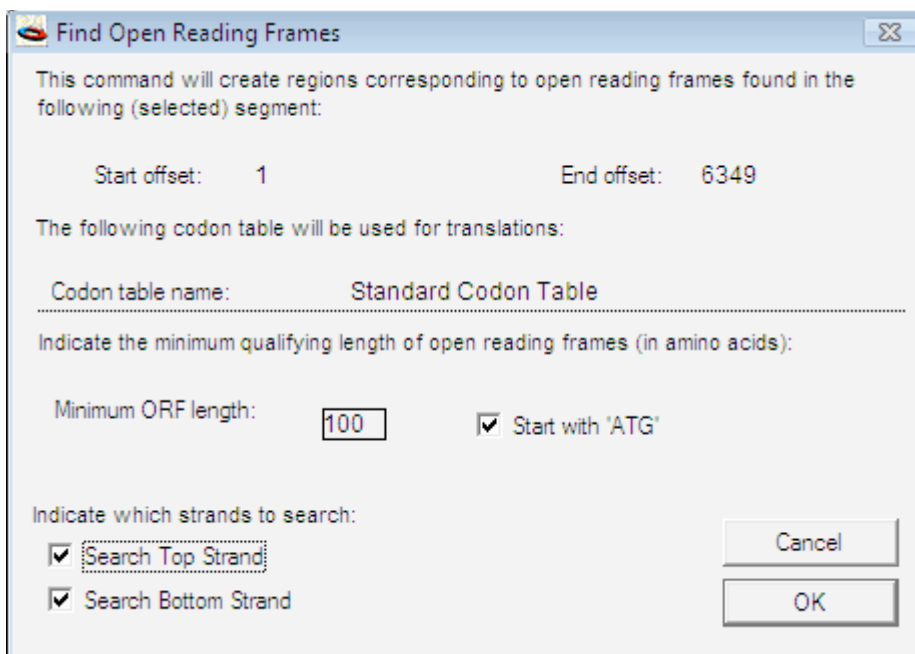


**Figure 7.31: Creating a New Region**

can be selected and then modified using any of the appropriate commands on the **Format** menu. You can adjust the starting and ending nucleotides to be used for creation of this region by typing in the value you wish. Checking **Protein Sequence** in this dialog will cause **GCK** to translate the sequence using the default codon table (“Choose Codon Table...”, page 7-45) whenever the construct is displayed as a sequence. The **Protein Direction** buttons allow you to define which strand will be used to create the protein sequence. Note that the direction of a selected region can also be changed by using the **Format > Lines** menu and choosing a different direction for the arrowhead.

### **...Find Open Reading Frames...**

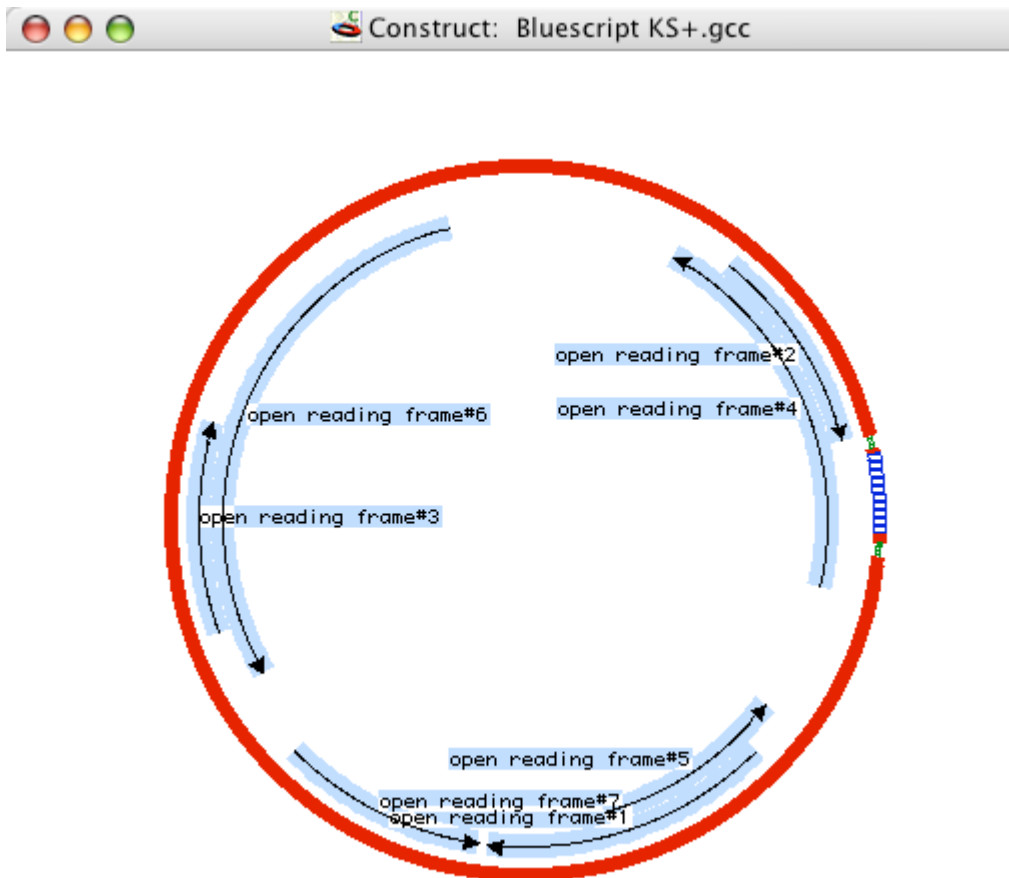
The **Construct > Features > Find Open Reading Frames...** menu item will search the construct and find all open reading frames according to the parameters set in the dialog shown in Figure 7.32. If part of the construct is selected, the ORF search will look only at the selected segment. If no segment of DNA is selected, the entire construct will be searched. The starting and ending offsets (nucleotide positions) are indicated in this dialog along with the codon table to be used in doing the translations (set the codon table using “Choose Codon Table...”, page 7-45). In most instances, the standard table will be appropriate. This corresponds to the standard genetic code found in most text



**Figure 7.32: Find ORFs Dialog**

books. If you are working with some mitochondrial genomes or other unusual organisms, you might need to choose a different table. If there is no table corresponding to your organisms, you can create one using “Edit Codon Tables...”, page 7-44.

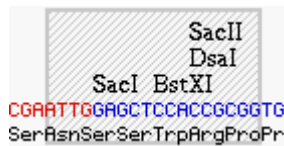
Also in this dialog box, you can type in a number for the minimum length open reading frame to be reported – anything less than this length will not be shown. Use the **Start with ‘ATG’** check box to specify whether all reading frames should start with ATG or not. For prokaryotic sequences, you probably want to check this box. For eukaryotic sequences, be careful not to check this box if you are looking at downstream exons. Finally, you can specify whether the search will examine the top and/or the bottom strand of the DNA sequence. Once the ORFs are found, they are automatically marked and named in the Construct window. If Show Region Names is on (“Show Region Names”, page 7-28), the names will be shown next to the regions themselves as shown in Figure 7.33, page 7-48.



**Figure 7.33: Open Reading Frames**

### **...Make Frame...**

Frames were discussed in some detail in Tutorial 5:Modifying the Construct Appearance, page 2-20. Frames are boxes that can be placed around parts of a construct sequence listing. Construct > Features > Make Frame will create a frame around any selected sequence. An example is shown in Figure 7.34. Once a frame is created, its appearance can be changed using the items in the Format menu as explained in Tutorial 5. In this figure, the frame is a light gray and the fill pattern is striped. This does not obscure the sequence listing.



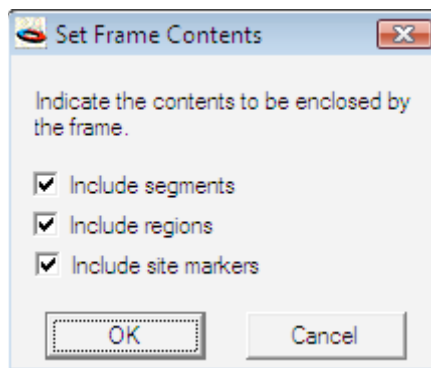
**Figure 7.34: A Frame**

A frame can be selected in any of three ways:

- hold down the option key and click anywhere inside the frame
- choose Construct > Select Frame (“Select Frame”, page 7-67) and then click on the frame you want to select
- triple click on the sequence contained in the frame

### **...Set Frame Contents...**

This topic was discussed in Tutorial 5:Modifying the Construct Appearance, page 2-20. The frame shown in Figure 7.34, page 7-49 encloses not only the sequence but the marked sites and the translated protein sequence. Construct > Features > Set Frame Contents will allow you to adjust what is actually enclosed in the frame. Selecting this option will bring up Figure 7.35. By



**Figure 7.35: Adjusting Frame Contents**

using the checkboxes in this dialog, it can be specified whether the frame should include segments of DNA, regions (graphic arrows or translated sequences), and/or marked sites.

### ...Mark Sites...

This was discussed in great detail in Tutorial 2:Marking Sites, page 2-8. Construct > Features > Mark Sites (comment-M/ctrl-M) is used to search the DNA sequence and place site markers at the locations corresponding to recognition sequences of items in the chosen list. Lists are **GCK** files that contain restriction enzymes or other items that can recognize specific sequences (see Chapter 4: “List Window”, page 4-1). Choosing Mark Sites will bring up Figure 7.36, page 7-50. As explained in more detail in “Marking Sites”, page 3-



**Figure 7.36: Mark Sites Dialog**

10, this dialog allows you to specify how the sites are to be marked on the construct. You can:

- select the enzymes to be used
- choose the kinds of cut sites you want to be marked (blunt, 5', 3')
- define the number of sites an enzyme must have to be shown
- specify whether the site marker should be displayed as a symbol or as a text label
- and specify whether the enzymes should cut only in specific segments of the DNA

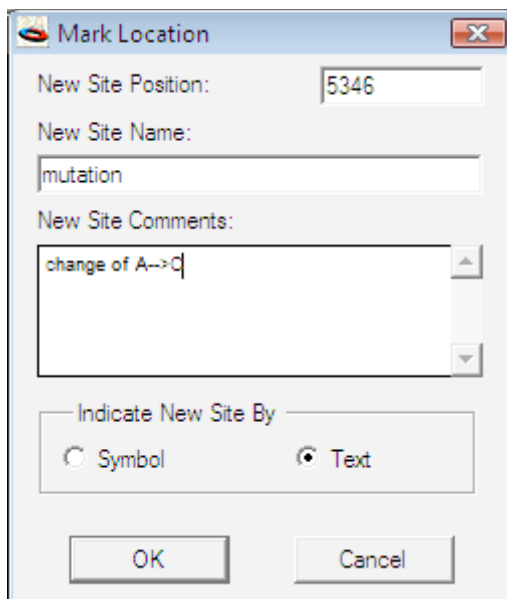
See “Marking Sites”, page 3-10 for details on each of these choices.

### **...List Sites...**

Construct > Features > List Sites (command-L/ctrl-L) performs an operation similar to that of Construct > Features > Mark Sites (“Mark Sites...”, page 7-50). Each choice will search a construct for locations that match a sequence specified in a list file. While Mark Sites will actually place site markers on the construct, List Sites will create a text listing of sites on the construct. List Sites is discussed in detail in “Listing Sites”, page 3-15.

### **...Mark Location...**

Construct > Features > Mark Location... is will place a marker on the construct at a defined location. Unlike Mark Sites..., Mark Location requires the exact position for



**Figure 7.37: Mark Location Dialog**

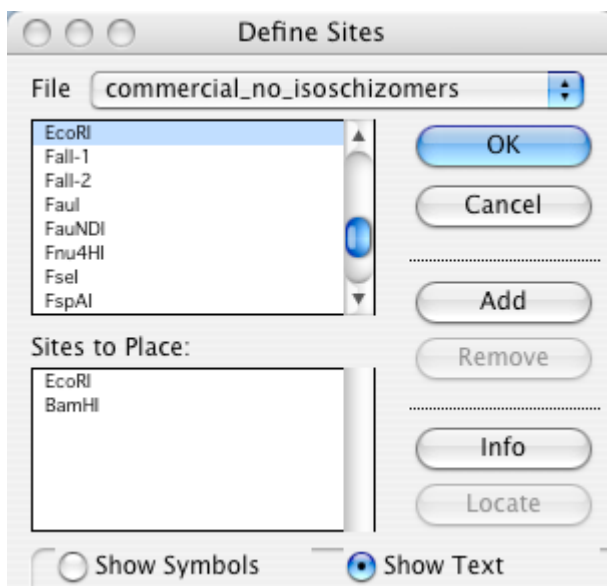
the site marker and the name for the site marker to be typed into the dialog box shown in Figure 7.37. Comments to be associated with the marked location may also be entered. These comments will remain with the site marker during subsequent cutting and pasting operations. Marking a location might be useful for indicating features that are not easy to store in a List window. Examples are transcription start sites, polyadenylation signals, and splice

sites.

Marked locations function the same as marked sites in enabling the selection of segments of DNA. In the most ancient generation of the **pBR322** construct packaged with **GCK**, the three pieces of DNA that were source plasmids for **pBR322** were originally defined by placing marked locations at the specific nucleotides needed to define each segment. Once the segments had been defined, the marked locations were discarded.

### **...Place Sites...**

Construct > Features > Place Sites... (command-J/ctrl-J) was discussed in Tutorial 11: Working With Generic Constructs, page 2-56. Place Sites... provides the opportunity to insert recognition sequences into a construct at specific locations. Using this feature, you can assemble generic constructs with accurately placed restriction sites, even though the sequence is not known. You will see



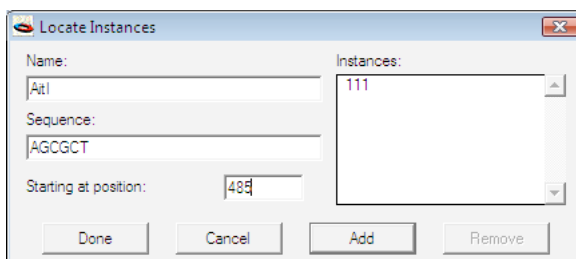
**Figure 7.38: Place Sites Dialog**

the dialog box shown in Figure 7.38. This dialog is the first step in placing the sites into the construct. Select an enzyme from the top list whose recognition sequence you want placed into the construct and then press the **Add**



button (you can also double-click on the enzyme name in the top list to make it appear in the Sites to mark list). In Figure 7.38, *Acil* has been chosen and added to the list on the bottom. An item can be removed from the *Sites to Place* list by first selecting it and then pressing the **Remove** button.

When an item is selected in the bottom list, the **Info** button and **Locate** button become enabled. **Info** will present a simple 'Get Info' type of dialog box indicating the cut site and other information about the selected enzyme in a non-editable window. Pressing **Locate** will present you with the subdialog box shown in Figure 7.39. The enzyme *Acil* was selected in this case and,



**Figure 7.39: Defining Place Sites Locations**

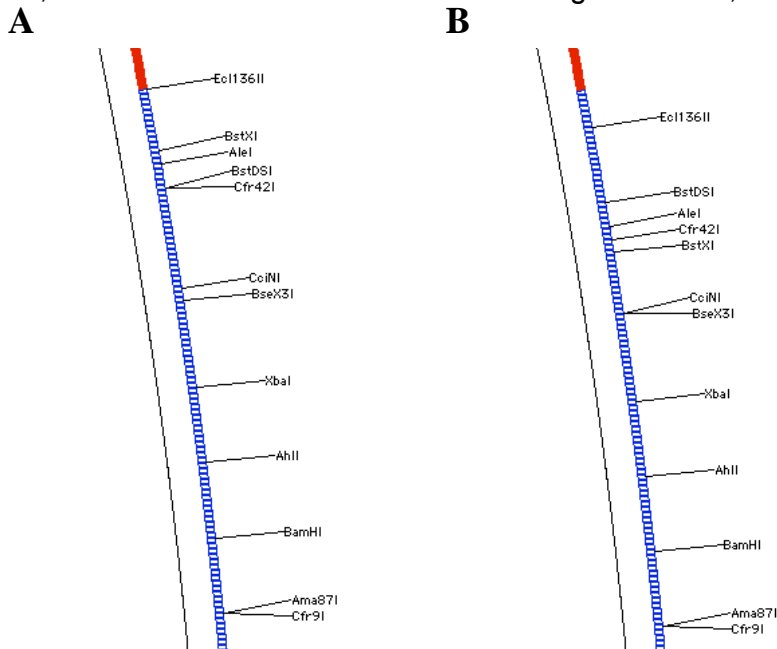
as the figure shows, the recognition sequence for *Acil* (CCGC) will be placed starting at nucleotides 100 and 485. The second location at 485 has been typed into the *Starting at position* box. Pressing the **Add** button will add 485 to the *Instances* list at the right of the window. To remove an item from the *Instances* list, select the item and then press the **Remove** button. After all locations have been defined for this enzyme, press the **Done** button to return to the main dialog (Figure 7.38, page 7-52), where the locations for placing sites for other enzymes can be defined.

**Note that** Place Sites *actually changes the sequence of nucleotides in the construct*. The construct sequence at the specified location(s) is replaced by the sequences representing the enzyme recognition sequences chosen. Any nucleotides in the construct, not just Ns, will be replaced! Be sure that this is what you want to do because it is not undoable.

### **...Place Site Indicators At...**

Construct > Features > Place site indicators at... can be used to specify the placement

of the site markers on the construct. You will be presented with a dialog that gives you a choice of placing the marker at the start of the recognition sequence or at the left of the actual cut site. This will only make a noticeable difference when you are zoomed in to a high magnification in areas that are densely packed with restriction site markers. The difference is shown in Figure 7.40, which is drawn at 3.6 nts/cm. In Figure 7.40A, the markers are

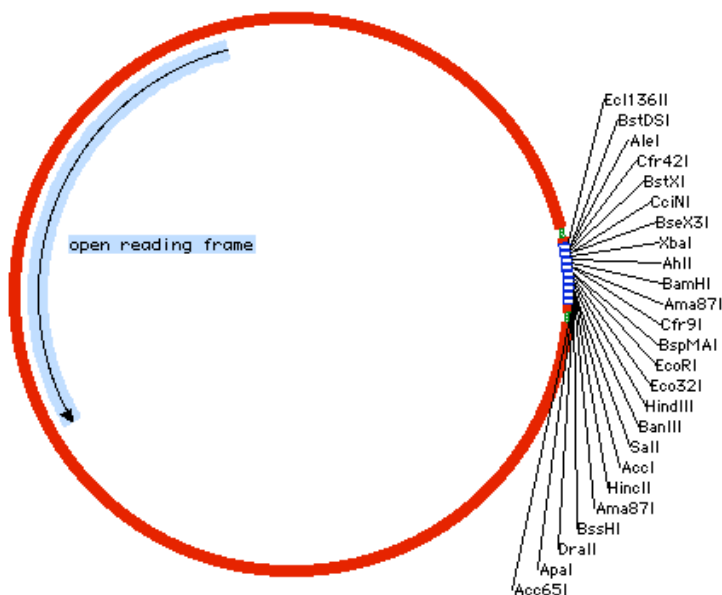


**Figure 7.40: Site Markings**

placed at the recognition sequence. Notice that the insertion point (indicated by the arrow) is not located where the *SmaI* site intersects the construct DNA sequence. The *SmaI* marker text is at the start of the *SmaI* recognition sequence, while the insertion point (where any new sequence would be pasted in) is further downstream of the site. In Figure 7.40B, the *SmaI* marker text is placed at the cut site. Now it corresponds to the location of the insertion point. Traditionally, restriction enzyme names have been placed above DNA sequences such that the first letter of the restriction enzyme name is exactly over the first letter of the recognition sequence for that enzyme. You can choose either method in **GCK**.

### ...Insert Sites By Silent Mutation...

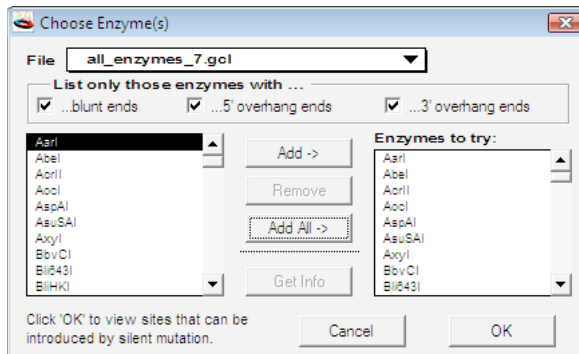
Silent mutations are changes in a DNA sequence that do not alter the coding information contained in the DNA. Silent mutations are often used to create



**Figure 7.41: Selected Region for Inserting Silent Mutation**

convenient restriction enzyme sites or to remove inconvenient sites (see “Remove Site By Silent Mutation...”, page 7-58 and Tutorial 6:Cloning a DNA Segment and Silent Mutations, page 2-23). Construct > Features > Insert Sites By Silent Mutation allows you to define a segment of DNA into which you want to place a new enzyme site. To begin, you must first select a translated region – this is essential because GCK must know which reading frame is to be conserved. This is shown in Figure 7.41. In this case, the goal is to insert a new restriction site for a 7 base recognition site enzyme. When the region is selected (Figure 7.41), Insert Sites By Silent Mutations... becomes enabled. Unless a region is selected, the menu item will be disabled.

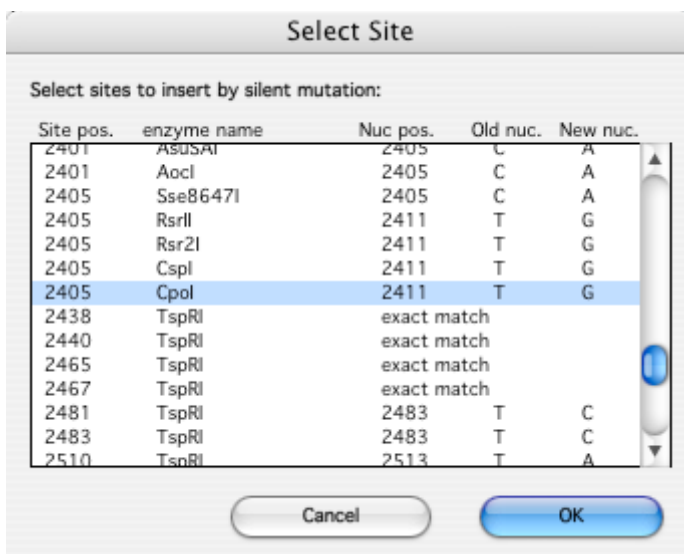
Choosing Insert Sites By Silent Mutations... will bring up a dialog box asking you to



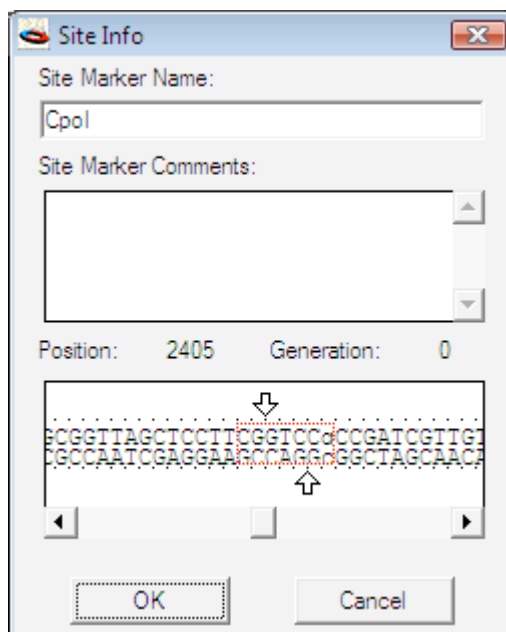
**Figure 7.42: Choosing Enzymes for Inserting Silent Mutations**

to this dialog box. The popup list at the top allows an enzyme list for use as candidate enzymes to be chosen; in Figure 7.42, page 7-56, *commercial* has been chosen. The next cluster of check boxes allows you to choose the kinds of cut sites to be considered in the search. The enzyme list on the left shows all the enzymes in the currently selected list file. Select one or more enzymes in the left list and then press the **Add** button to add them to the right list. All enzymes in the right hand list will be candidates for the insertion of a silent mutation. Pressing **OK** starts the process of examining the DNA sequence and locating sites in the DNA that could be altered by a point mutation to create a new enzyme site while not changing the coding of the DNA. Pressing **OK** will bring up a dialog like that shown in Figure 7.43. In the figure, the *CpoI* site is chosen at position 2405 in the DNA; this position represents the start of the recognition site for *CpoI*. The suggested change is to take place at nucleotide 2411 and will replace a 'T' with a 'G'. Note that *CspI* and *RsrII* sites will also be created in this particular instance because changing the 'T' to a 'G' will also create recognition sites for those enzymes. If the *SapI* site at 2164 were chosen, then only that one site would be created (the change of 'G' to 'A' at 2164 would not produce any other sites corresponding to any of the candidate enzyme recognition sequences).

The changed nucleotide will be converted to lower case to make it easy to identify, as shown in Figure 7.44. This dialog box is the standard “Get Info...” dialog available by choosing Construct > Get Info... (“Get Info...”, page 7-68).



**Figure 7.43: Creating a New Site**



**Figure 7.44: A Silent Mutation**

Note that the last 'g' in the sequence is in lower case because it was changed from the original sequence. The *CpoI* recognition sequence is boxed.

### **...Remove Site By Silent Mutation...**

Construct > Features > Remove Site By Silent Mutation... was discussed in detail in Tutorial 6:Cloning a DNA Segment and Silent Mutations, page 2-23 and complements “Remove Site By Silent Mutation...”, page 7-58. This option will create a single point mutation to remove a restriction enzyme recognition site without altering the coding of that segment of DNA. As is true for the “Insert Sites By Silent Mutation...”, page 7-55, you must first select a region for this menu item to be enabled.

### **Display**

The Display submenu has options pertaining to the display of the construct in both the sequence view and the graphics view.

### **...Show as Double/Single Stranded**

Construct > Display > Show as Double/Single Stranded (⇧-command-D/⇧-ctrl-D) pertains only to the sequence view of the construct. As the menu name implies, this option allows you to toggle between displaying the sequence as double stranded and single stranded. Selection and cut/copy/paste operations function the same in the single stranded mode as in the double stranded mode. The difference lies in the fact that both strands are highlighted in the double stranded mode.

### **...Show/Hide Positions**

Another menu that is available only in the sequence view is Construct > Display > Show/Hide Positions (⇧-command-K/⇧-ctrl-K). Showing positions will place position numbers at the beginning of each line of sequence. It will display the nucleotide position of the first nucleotide in the line of DNA sequence and will display the leftmost amino acid position of the peptide line (if a peptide is visible).

### **...Show/Hide Line Borders**

If the construct is being viewed as a sequence Construct > Display > Show/Hide Line

Borders ( $\uparrow$ -command-E/ $\uparrow$ -ctrl-E) will be available. Line borders serve a function analogous to page borders in a larger document that spans more than one printer page (see “Show/Hide Page Breaks”, page 7-15). Line borders delineate one sequence line from the next. This may not seem necessary under most circumstances, but in situations in which the DNA is being displayed as double stranded and both regions and marked sites are visible, it can sometimes be confusing as to which DNA sequence line is related to which set of marked sites and regions.

### **...Set Line Spacing...**

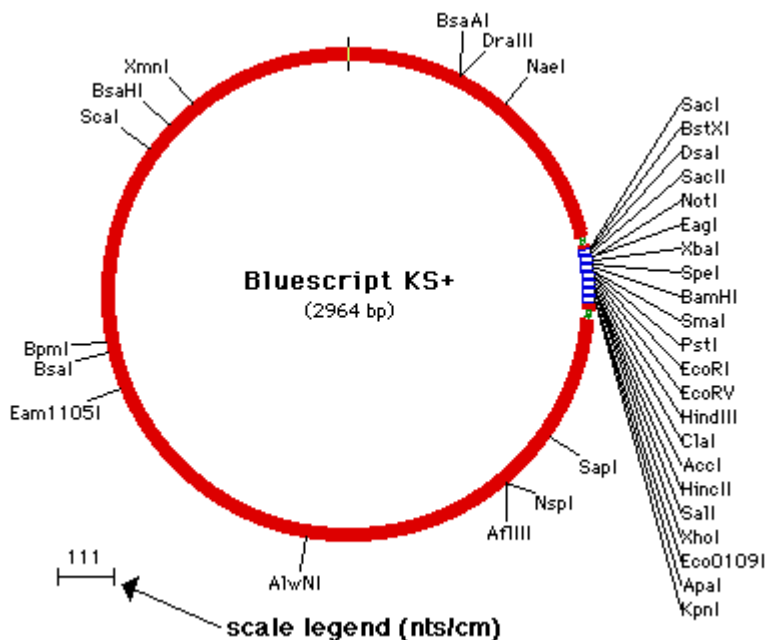
The Construct > Display > Set Line Spacing... menu item (also available only in the sequence view) will let you specify the minimum number of pixels between adjacent sequence lines. The default value is zero leaving the line spacing between sequence lines equivalent to single spacing. As you change the value from zero to a positive value, each line of sequence will be moved apart by that many extra pixels. Line spacing can be set in the default preferences.

### **...One/Three Letter Amino Acid Code**

This pair of menu items will only be available when the construct is being viewed as sequence. Choosing Construct > Display > One Letter Amino Acid Code will display all translated sequences as single letter amino acid code. Choosing Construct > Display > Three Letter Amino Acid Code will display all translated sequences as three letter amino acid codes. This is a global setting so all protein sequences are affected by this change. You cannot show different translations in the same construct using different length amino acid codes.

### **...Show/Hide Scale Legend**

When viewing a construct graphically, Construct > Display > Show/Hide Scale Legend is available. Show Scale Legend will show a horizontal bar in the construct window indicating the scale at which the construct is drawn as shown in Figure 7.45, page 7-60. If the window is resized and the magnification is set to “Fit to Window” (see “Fit To Window”, page 7-43), the scale legend will be updated



**Figure 7.45: Scale Legend**

to reflect the current scale for that window.

Double-clicking on the scale legend will allow you to actually set the scale at which the construct is to be Drawn. This will bring up the dialog box shown in Figure 7.28, page 7-42. The box allows you to define both the scale in nucleotides/cm and the length of the scale indicator. The program will create the scale as close to the desired value as possible. Since there are only a fixed number of pixels per inch on the screen, it is usually not possible to have the on-screen scale exactly as specified. However, the *Actual Scale on Screen* value will define that scale precisely. Once the scale is set using this dialog box, **GCK** assumes that you want the construct to stay at that scale. Therefore, resizing the window will not change the scale. To have the construct resize as the window is resized, you should select Construct > Magnification > Fit To Window (see “Fit To Window”, page 7-43).



### **...Show/Hide Construct Title**

The construct title is text that can be displayed in the center of a circular construct, or centered below a linear construct. This text has the title of the file as its first line, and the length of the construct as its second line of text. You can select each line of text and change the text attributes using the Format menu. Construct > Display > Show Construct Title and Hide Construct Title toggle between making the title visible or not.

### **...Center Construct Title**

You can select the construct title and move it to any location in the window. Sometimes, it is desirable to return the title to its original location. Construct > Display > Center Construct Title will return the title to the center of a circular construct or centered below a linear construct.

### **...Show/Hide Regions**

As the name suggests, this menu item will toggle the display of regions on or off. It is a global setting so that all regions will be either shown or hidden. Regions are features that are not part of the actual DNA, but represent additional information about the DNA (see “Regions of Interest and Translating DNA”, page 3-5). You cannot set this property for just one region. If you are interested in having different regions displayed under different situations and do not want to dispose of any regions, you can do this using Chronography (see “Chronography - Tracking Construct History”, page 3-32). You can use a different chronography generation to display each set of regions you are interested in displaying.

### **...Show/Hide Frames**

As the name suggests, this menu item will toggle the display of frames on or off. It is a global setting so that all frames will be either shown or hidden. Frames are features that adorn the display of the DNA sequence (see “Make Frame...”, page 7-48 and Tutorial 5:Modifying the Construct Appearance, page 2-20). You cannot set this property for just one frame. If you are inter-

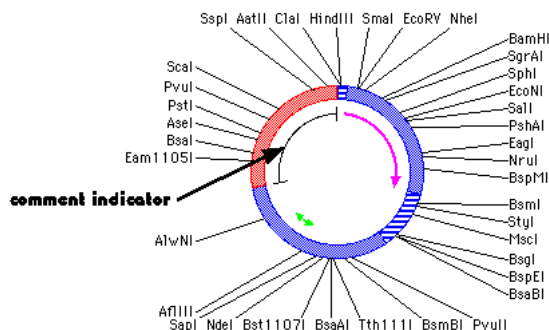
ested in having different frames displayed under different situations and do not want to dispose of any frames, you can do this using Chronography (see “Chronography - Tracking Construct History”, page 3-32). You can use a different chronography generation to display the sequence with a different set of frames.

### **...Show/Hide Sites**

As the name suggests, this menu item will toggle the display of site markers on or off. It is a global setting so that all site markers will be either shown or hidden. Site markers are features that are not part of the actual DNA, but represent additional information about the DNA (see “Marking Sites”, page 3-10 and Tutorial 2:Marking Sites, page 2-8). You cannot set this property for just one site marker. If you are interested in having different site markers displayed under different situations and do not want to dispose of any site markers, you can do this using Chronography (see “Chronography - Tracking Construct History”, page 3-32). You can use a different chronography generation to display each set of site markers you are interested in displaying.

### **...Show/Hide Comments**

As the name suggests, this menu item will toggle the display of comment indicators on or off. Comments can be attached to any segment of DNA to store annotations that are easily available. Sometimes, however, you might



**Figure 7.46: Comment Indicator**

lose track of the comments and not recall which segment of DNA contains the

comments in which you are interested. A comment indicator is shown in Figure 7.46 and is discussed in “Attaching Comments to Objects in Construct Window”, page 3-6. The presence of a comment indicator denotes that there are comments associated with that particular segment of DNA. You can view the comments by clicking once on the segment indicator to select it and then choosing `Construct > Get Info...` You can double-click on the comment indicator to select the corresponding segment of DNA and then view the comments by choosing `Construct > Get Info...` In operational terms, these comment indicators behave very similarly to regions. However, comment indicators and regions can be shown and hidden separately, regions do not have to have comments, and regions can represent protein sequences.

### **...Show/Hide Generations**

As the name suggests, this menu item will toggle the display of generation indicators on or off. Generations are part of the chronography feature of **GCK**. This is discussed in detail in “Chronography - Tracking Construct History”, page 3-32 and in Tutorial 7:Chronography – Tracking Cloning History, page 2-30. If you are viewing a construct that is displaying multiple generations, you can view indicators of which segments are being shown in each generation. Given the possibility of displaying different segments in different generations<sup>b</sup>, it might become difficult to keep track of which generations are being displayed. To address this problem, **GCK** allows you to indicate the display of different generations using the `Construct > Display > Show/Hide Generations`. When you are showing generations, you will get an extra set of lines on the outside of the construct for all segments not being displayed in the current generation, as shown in Figure 3.35, page 3-37. Those segments being displayed in the current generation will not have an indicator line next to them. The adjacent fragments are being displayed in previous generations, and therefore are indicated as such by the gray indicator around the outside of the circle.

---

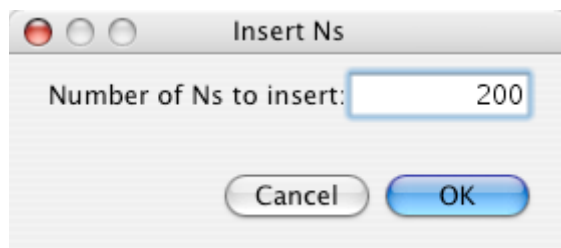
b. Once any segment is selected, it can be displayed in any generation by using the Chronography menu. In this way it is possible to display different segments in different generations.

### **...Display Sequence/Graphic**

Any construct can be viewed as either a graphic representation of the sequence or as an actual sequence listing. The two views are actually different ways of displaying the same information. Construct > Display > Display Sequence and Construct > Display > Display Graphic (shift-command-/shift-ctrl-/) will toggle between the two different views of the construct.

### **Insert Ns...**

A string of generic nucleotides (Ns) can be inserted into a construct at any time. Choosing Construct > Insert Ns... (command-H/ctrl-H) from the Construct menu will present the dialog box shown in Figure 7.47. This is useful even in empty



**Figure 7.47: Insert Ns**

Construct windows to generate restriction maps and figures for constructs that have not yet been sequenced. Used in conjunction with the Place Sites option (“Place Sites...”, page 7-52), complete figures can be made even if your information consists of only the construct size and the locations of some restriction sites.

### **Insert Typing...**

Construct > Insert Typing... is available only in the sequence view of the construct. It can be activated either by selecting it from the menu, or by placing the cursor in the sequence at the location you wish and then starting to type. This is discussed “Inserting New Nucleotides by Typing”, page 3-21.

### **Edit Construct End.../Edit Cut Site...**

When the cursor is placed at the end of a linear construct, the Construct > Edit

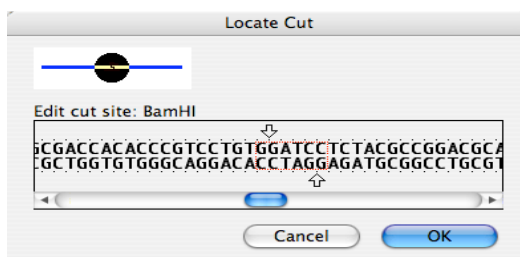
Construct End... (command-E/ctrl-E) menu option is enabled. Selecting this menu item will bring up the dialog box shown in Figure 7.48, page 7-65. As the



**Figure 7.48: Edit Construct End**

picture in the upper left of the dialog indicates, the visible sequence represents the right end of the construct. This end can be edited in the standard Gene Construction Kit fashion – either drag the arrows to trim nucleotides from a strand, or click the mouse at the desired new location for the arrow.

If the cursor is not located at an end in the construct, but is instead at a marked site, the menu changes to Edit Cut Site... and will present you with a dialog box similar to Figure 7.49. By dragging the arrows, you can adjust the cut



**Figure 7.49: Edit Cut Site**

site for the particular enzyme. Be sure to change the name of the enzyme if you alter the cut site.

### **Find Sequence...**

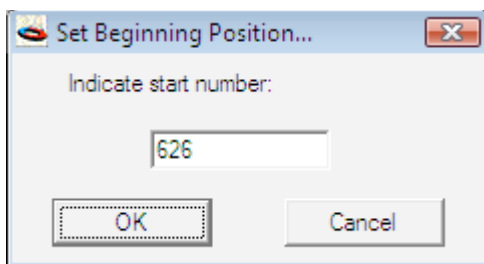
Construct > Find Sequence... (command-F/ctrl-F) was discussed in detail in “Finding a Sequence”, page 3-23. Using the standard letters for DNA (see “Legal Nucleotide Characters”, page 8-1), GCK will search through the construct and

highlight any match with the search sequence. The search starts from the current location of the cursor and continues clockwise (for circular DNAs) or in the 5' --> 3' direction for linear DNAs. Once the end of a linear DNA is reached, the searching will stop. Searching can continue around a circular construct.

### **Set Beginning Position**

#### **Redefine Origin**

In the graphical view of a construct, either `Construct > Redefine Origin` or `Construct > Set Beginning Position` (command-1/ctrl-1) will be available, depending on whether you are working with a circular or linear construct. For circular constructs, `Redefine Origin` will rearrange the construct so that the current position of the insertion point will become the new origin for the construct. This will rotate the construct on the screen to bring the origin to the top of the construct. For linear constructs, `Set Beginning Position` will allow you to define a different number for the starting nucleotide in the sequence using the dialog shown in Figure 7.50. This would be useful, for example, if the DNA you are working with



**Figure 7.50: Set Start Position**

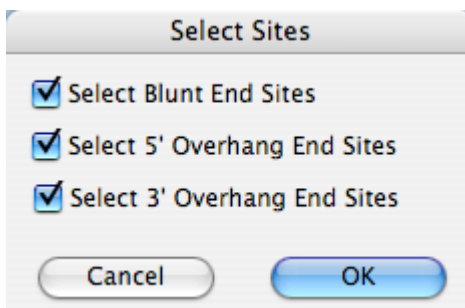
represents a segment of a much larger sequence, but the segment it represents is not the first segment of the DNA. In this case, it might be easier to maintain consistency if the segment numbering started at a position representing the actual position of the nucleotide in the whole sequence (e.g. 86). Another case might be to indicate the position relative to the start of transcription (e.g. -266). For creating figures for papers, this feature is very handy.

### **Select Frame**

Frames represent a way to highlight specific parts of a sequence listing. Frames are discussed in Tutorial 5:Modifying the Construct Appearance, page 2-20 and in “Make Frame...”, page 7-48. The sequence view of a construct allows you to select site markers, segments of DNA and regions as well as frames. Sometimes it might be confusing as to how to select a frame. Choosing Construct > Select Frame will change the cursor to a “cross-hair”; clicking inside any frame will select that frame. Frames can also be selected by triple-clicking on the DNA sequence inside the frame or by holding down the option key and clicking inside the frame you wish to select. Once a frame is selected, you can use the items in the Format menu to modify the appearance of the frame.

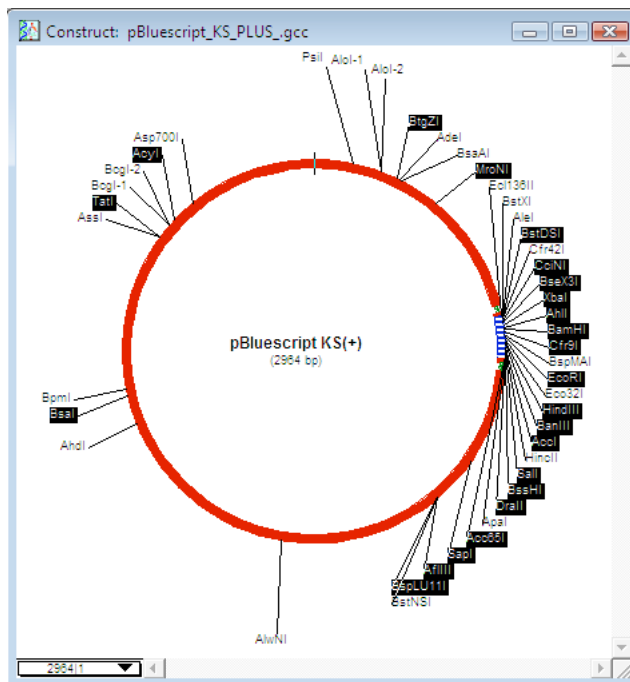
### **Select Sites By Kind**

Construct > Select Sites By Kind will select a subset of the marked sites on any construct depending on the properties of the enzyme recognition site. Using



**Figure 7.51: Sites By Kind Dialog**

the dialog box shown in Figure 7.51, you can specify which kind of site should be selected. The results of this selection on pBR322 with all the unique sites indicated is shown in Figure 7.52, page 7-68. It might be useful to change the color or font of all these sites so that they can be distinguished from 5'-overhang and 3'-overhang sites. The 5' and 3' sites can also be selected and color coded to make recognition easier. Having this kind of information available at a glance will facilitate cloning strategy design.



**Figure 7.52: Blunt End Sites Selected**

## Get Info...

Get Info... (command-I/ctrl-I) will provide information on any selected object in the Construct Window. You can Get Info... on construct segments, marked sites, or regions. Depending on the object selected, you will get a different but similar dialog box containing information relevant to the selected object. Some of these have been discussed before. A region get info box is shown in Figure 3.8, page 3-9; a site get info box is shown in Figure 3.7, page 3-9; and a segment get info box is shown in Figure 3.4, page 3-6. Get Info... also works when a construct is targeted in an Illustration window.

## General Info...

Construct > General Info... differs from Get Info... in that it provides information about the entire construct and is not attached to any particular object within the construct. This dialog is shown in Figure 3.6, page 3-8. General informa-



tion will always stick with the construct – it remains attached to the window (file) and will not be lost when segments are cut and pasted within the window. This is in contrast to the segment comments which are lost whenever the segment is disrupted. Copying an entire construct and pasting it into a new window will copy the general information. Pasting the entire construct as a segment of another construct will result in the conversion of the general information into info attached to the new segment resulting from the paste operation.

### **Search Comments...**

The Construct > Search Comments... option was discussed in “Searching Comments”, page 3-22. It provides the ability to search through all the comments associated with a construct in all generations. As shown in Figure 3.19, page 3-22, general info, segments, regions, marked sites, and chronography generations can be searched in an intuitive way. Matches that are found are listed and comments can then be viewed.

### **Make Linear/Make Circular**

Construct > Make Linear/Make Circular (command-V/ctrl-V) toggles the display of the construct between linear and circular. Making a circular construct into a linear one presents the option of opening the construct either at the insertion point or at the origin as shown in Figure 7.53. Converting a linear construct to cir-



**Figure 7.53: Make Linear Dialog**

cular joins the ends of the linear construct to become the origin. If a ligation dialog box is needed, it will be shown.

### **Gel Menu**

The Gel Window is discussed in detail in Chapter 5: “Gel Window”. A typical gel window is shown in Figure 5.1, page 5-1.

### **Show/Hide Runoff Count**

At the bottom of each lane in a gel window is an indication of the number of fragments that have migrated off the bottom of the gel. If no fragments are smaller than the threshold set for the gel, then no number is placed below the lane. If too many fragments run off the gel, increase the threshold for the gel (see “Set Threshold...”, page 7-72). These numbers are called the runoff count and can be made visible or not by using the Gel > Show/Hide Runoff Count menu option.

Hide Runoff Count	
Hide Gel Legend	Ctrl+L
Hide Standard Sizes	
Show Full Digest	
Show Partial Digest...	
Set Threshold...	Ctrl+T
Display Table	Ctrl+ /

### **Show/Hide Gel Legend**

The Gel legend is discussed in “Legend”, page 5-2. The **legend** is visible just to the right of the gel and indicates which DNA and enzyme(s) were used to produce the banding patterns in each lane, and in the case of partial digests, specifies the extent of digestion. The legend consists of lane numbers and the actual legend adjacent to the gel. The legend can be made visible or not using Gel > Show/Hide Gel Legend (command-L/ctrl-L).

### **Show/Hide Standard Sizes**

Along the left side of the gel (to the left of lane 1) are a set of standard size indicators. Show or hide the standard numbers by using Gel > Show/Hide Standard Sizes. Each solid band in the standard represents a multiple of the threshold value (see “Threshold”, page 5-2) for the gel. The dotted lines represent 1.5x and 15x the threshold value.

### **Show Full Digest**

Selecting a lane on the gel and choosing Gel > Show Full Digest will show the

bands that would result if complete digestion took place. This option is used to convert a partial digest lane (see next section).

### **Show Partial Digest...**

Partial digests were discussed in “Partial Digests”, page 5-4. Choosing Gel > Show Partial Digest... brings up a dialog (Figure 5.4, page 5-5) that allows you to define the extent of digestion by the enzymes chosen. Bands on the gel that are the result of a partial digest will be represented as a dotted blue line instead of a solid black line.

### **Set Threshold...**

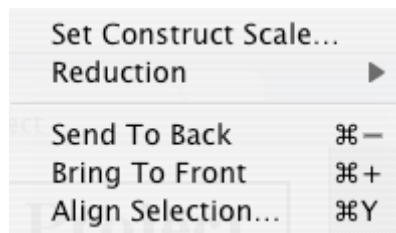
Gel > Set Threshold... (command-T/ctrl-T) defines the minimum sized fragment that will remain on the gel. The dialog box is shown in Figure 5.2, page 5-2 and thresholds are discussed in “Threshold”, page 5-2.

### **Display Table/Display Graphics**

Gel > Display Table/Display Graphics (command-/ | ctrl-/) will toggle the display between listing the fragments for each lane in a tabular form (Figure 5.3, page 5-3) and illustrating the predicted gel pattern graphically (Figure 5.1, page 5-1). Each view presents the same data, but one is graphical and the other is textual.

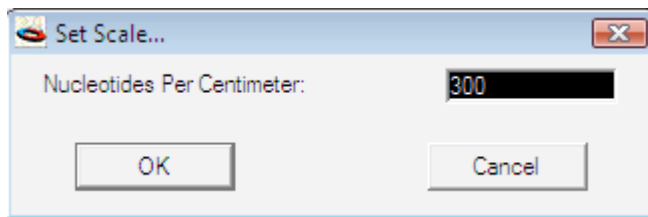
### **Illustration Menu**

The Illustration menu is simple and consists of just a few items. It allows you to set the scale in nts/cm for constructs, to set the drawing scale for the whole window, and to arrange objects in the Illustration window. Note that other controls for the Illustration window are located in the bottom left corner of the window frame itself in the Tool Palette (Figure 6.3, page 6-3).



### **Set Construct Scale...**

The Illustration > Set Construct Scale... menu option is available when one or more construct(s) is selected in the Illustration window. Set Construct Scale... sets the drawing scale for any construct using the dialog box shown in Figure 7.54. It



**Figure 7.54: Set Construct Scale**

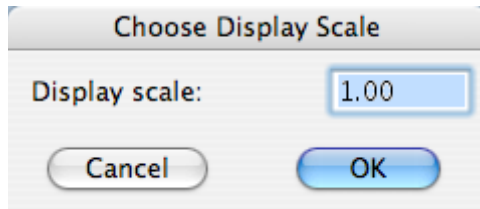
does not place an indicator on the screen. To set all the constructs in an Illustration to the same scale, select one of the constructs by clicking on it, choose Edit > Select All, and then choose Set Construct Scale....

### **Reduction**

The items in the Reduction submenu allow you to set the degree of reduction for drawing the Illustration window. It is not possible to view the Illustration window at a magnification of more than 1x. You can either view the window at full size (no reduction) or at various degrees of reduction.

### **...Set Reduction...**

Illustration > Reduction > Set Reduction... allows you to enter a value in the dialog box shown in Figure 7.55 to specify the actual reduction. In this case a value



**Figure 7.55: Set Reduction Dialog**

of 0.5 has been entered so the drawing will be at 50% normal size. In reduced mode, you will be able to move objects around but you cannot edit them (they are non “targetable”).

### **...Reduce**

Choosing Illustration > Reduction > Reduce will reduce the area of the Illustration window by approximately 50%. Note that this is not the same as a 50% reduction in linear dimensions, which would reduce the area to 25%.

### **...Enlarge**

Choosing Illustration > Reduction > Enlarge will increase the area of the Illustration window by approximately 50%. Note that this is not the same as a 50% increase in linear dimensions, which would increase the area by 400%.

### **...No Reduction**

Illustration > Reduction > No Reduction is the same as setting the reduction scale to 1.0. The Illustration window will be drawn at full size.

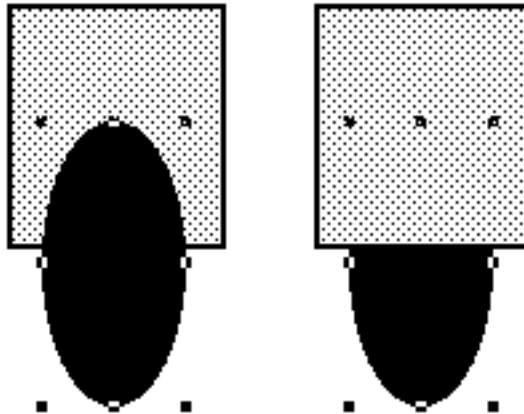
### **...Full Reduction**

Illustration > Reduction > Full Reduction is sometimes called “Reduce to Fit” in other applications. It will reduce the size of the Illustration window to fit entirely on your monitor. It provides a complete overview of the layout of your window.

### Send To Back

### Bring To Front

Illustration > Send To Back (command--/ctrl--) and Illustration > Bring To Front (command-+/ctrl-+) allow the manipulation of objects on the screen as shown in Figure 7.56. On the left, the ellipse is selected and is in front of the rectangle.

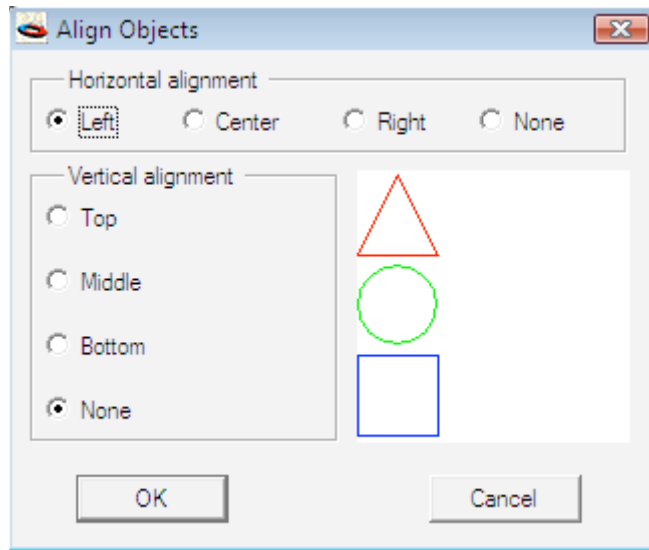


**Figure 7.56: Front-Back Stacking of Objects**

Choosing **Send To Back** will move the ellipse to the back of the picture, in this case, behind the rectangle, as shown on the right. Any object that can be selected in the Illustration window can be manipulated with these two commands.

### Align Selection

When one or more objects is selected in the Illustration window they can be aligned by choosing **Illustration > Align Selection**, which will bring up Figure 7.57, page 7-76. Separate control is provided for horizontal and vertical alignments. By clicking on one of the radio buttons, the triangle, circle, and square sample objects will align. In the figure, the **Left** horizontal alignment button was chosen so the left edges of all the objects are forced to align to the left-most position of any of the selected objects. Pressing the **None** button will leave the selected objects in their original position (either vertically or hori-



**Figure 7.57: Alignment Dialog**

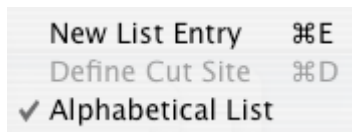
zonally). By choosing different combinations of buttons and examining the sample object alignment, you can easily determine how the objects you have selected in the Illustration window will be aligned.

The alignment of objects using *Align Selection* provides a convenient way to organize the various constructs, gels, and text which are displayed in the Illustration window.



### **List Menu**

The List menu allows you to manipulate the items in a list. All list files are stored in the **GCK Data** folder. If a list file resides outside a GCK Data folder, it will not be available from within the program. List files were discussed in detail in Chapter 4: “List Window”. In conjunction with the Cut, Copy, and Paste options under the Edit menu, one or more list entries can be moved between lists or removed from a given list.



### **New List Entry**

New List Entry (command-E/ctrl-E) was discussed on “New List Entries”, page 4-3. It provides a blank entry form to enter the name, sequence, and comments to be associated with your new entry in the list (see Figure 4.4, page 4-4). It will create a new list entry temporarily indicated by a bullet with an ellipses (...) in the list at the left. Fill in the name of the new entry, the sequence you want this entry to represent, and any comments to be associated with sites on the construct when that entry is used to mark sites.

### **Define Cut Site**

To define the actual site at which the enzyme cuts the DNA, use the Define Cut Site (command-D) option. This will bring up the dialog box shown in Figure 4.5, page 4-4. Use the triangles to define the cut site.

### **Alphabetical List**

If the Alphabetical List menu item is checked, then the list will be maintained in alphabetical order and all new entries into the list will be placed alphabetically. If the list is not alphabetical, new entries will be placed at the end of the list.



# Chapter 8

## Appendix

### Legal Nucleotide Characters

Letter	Nucleotide(s)	Complement
A	adenosine	T
C	cytosine	G
G	guanosine	C
T	thymidine	A
R	A, G (puRine)	Y
Y	C, T (pYrimidine)	R
M	A, C (aMino, in large groove)	K
K	G, T (Keto, in large groove)	M
S	G, C (Strong)	S
W	A, T (Weak)	W
N	A, C, G, T	N
B	C, G, T (not A)	V
D	A, G, T (not C)	H
H	A, C, T (not G)	D
V	A, C, G (not T)	B

## **Import File Formats**

NOTE: All files are imported using the rules defined below. All files with the exception of Staden files (which can use non-standard nucleotide characters) are filtered to accept only legal nucleotide characters as defined by international standards (“Legal Nucleotide Characters”, page 8-1). Lower case letters are converted to upper case and Us are converted into Ts. Any non-nucleotide characters are discarded. This includes all numbers, RETURNS, TABS, and line feeds. Importing sequences is discussed in “Importing Sequences Into GCK”, page 3-38.

### **DNA Inspector**

Standard file format generated by DNA Inspector™ Ile. These file formats are listed in the corresponding DNA Inspector manuals. Note that this file format is provided as an import option for our customers who still have DNA Inspector Ile, which is no longer a current product of Textco BioSoftware’s. Textco BioSoftware has replaced DNA Inspector Ile with Gene Inspector™. See our web site for more information.

### **GenBank**

Using File > Import, everything is put into comments through the line starting with ‘ORIGIN’. Any text after this line is considered sequence. To import all GenBank features as GCK features use Tools > Deluxe Import, page 7-35. See Tutorial 6: “Cloning a DNA Segment and Silent Mutations”, page 2-23 for more details.

### **Intelligenetics**

All lines beginning with a ‘;’ are considered comments. The first line without a semicolon is the sequence ID (name). Everything following this line is considered sequence.

### **Pearson (FASTA)**

Any line starting with ‘>’ is a comment line. Lines starting with > indicate the

start of a new sequence. Pearson files can have multiple sequences in the same file. Sequence lines are those that do not start with '>'.

### **NBRF**

The first line is the HEADER line and the second line is the TITLE; both these lines are put into comments. The sequence follows starting on the third line and continues until an asterisk is reached. Everything following the asterisk is comments.

### **Text**

Everything in a TEXT file is considered sequence. All non-nucleotide characters are removed, lower case letters are converted to upper case letters, and U's are converted to T's.

### **GCG (Wisconsin)**

Everything is comment until two adjacent periods. Everything after the periods is considered sequence.

### **EMBL (SwissProt)**

Starting at the first line of the file, every line is considered a comment including the line that starts with 'SQ'. The line immediately after the 'SQ' line starts the sequence. Sequence information continues until '//'.

### **Staden**

Sequence and comments are mixed. All comments start with '<' and end with '>'. Everything else is considered sequence. The Gene Construction Kit expects Staden sequence files to start with a comment.

### **Gene Inspector**

Gene Inspector is Textco BioSoftware's sequence analysis and molecular biologist's notebook program. **GCK** can directly import any one GI sequence into a construct window. If the GI file contains more than one sequence, you

can select the sequence you are interested in. Only one sequence can be imported at a time.

## **Comment Export Format**

```
<General>
>Name: constructName
>Date: date
>General Info:
All General Info goes here until next marked line (put NONE if no
comments)
    ...
    ...
    ...
<Generations>
>Generation: Int
>Name: generationName
Comments for above generation go here until next marked line
    ...
    ...
    ...
<Sites>
>Name: siteName
>Position: longInt
Comments for above site go here until next marked line
>Name: siteName
>Position: longInt
Comments for above site go here until next marked line
    ...
    ...
    ...
<Segments>
>Name: segmentName
>Direction: --- or --> or <-- or <-->
>Range: longInt to longInt
Comments for above segment go here until next marked line
    ...
    ...
    ...
>Name: segmentName
>Direction: --- or --> or <-- or <-->
>Range: longInt to longInt
Comments for above segment go here until next marked line
```

```
...
...
...
<Regions of Interest>
>Name: regionName
>Direction: --- or --> or <-- or <-->
>Range: longInt to longInt
Comments for above region go here until next marked line
...
...
...
>Name: regionName
>Direction: --- or --> or <-- or <-->
>Range: longInt to longInt
Comments for above region go here until next marked line
...
...
...

... this gets repeated for each generation
```

## **GCK Acknowledgements**

### **National Center for Biotechnology Information (NCBI)**

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