

# ZMap User Manual

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## Revision History

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## Introduction

ZMap is a software package that provides a visualisation and editing tool for genomic features. The software is written in C/C++, utilising the gnome toolkit (GTK2) to draw features on a canvas. ZMap accepts input from multiple sources in multiple formats across multiple genomes and is written in a way so that the addition of further formats is made as trivial as possible. Currently the list of formats includes GFF and DAS, which may reside in any one of; a file, an acedb instance, an http server. Multiple genomes and their associated features can be displayed in a single view as aligned blocks providing support for comparative annotation. ZMap provides facilities for editing and creating features, which can then be saved to a GFF file.

ZMap is part of the Otter annotation suite and contains a protocol for two-way communication with Otter so that it operates seamlessly as an additional display within Otter.

ZMap can call tools from the SeqTools suite of sequence visualisation tools to analyse alignments in more detail.

## Using ZMap

### Running ZMap on a GFF file

To run ZMap standalone on one or more GFF files, simply pass the file name(s) on the command line:

```
zmap file1.gff file2.gff
```

The sequence region will default to the region available in the GFF file. Alternatively, you can specify the sequence/region by passing the `-sequence`, `-start` and `-end` arguments on the command-line. If all of the files contain the same reference sequence, then their contents will be shown in the same view. If the files contain multiple sequences/regions, then they will be shown in multiple views.

### Running ZMap with a config file

If you have several sources, it can be convenient to specify them in a config file rather than typing them on the command-line. You can also then set up many different options and styles to use for these sources. To use a local GFF file as a source, a simple config file would look something like this:

```
[ZMap]
# The list of sources
sources=chr6-18

# Sequence name/start/end can be specified here, on the command-line, or
# omitted if they are to be taken from the file
sequence=chr6-18
start=2696324
end=2864370

# Source stanzas (one per source)
[chr6-18]
url=file:///Users/gb10/work/checkout/zmap/examples/features.gff
```

ZMap can then be started by passing the config file on the command line:

```
zmap --conf_file=/path/to/file
```

Alternatively, the config file can be saved as `/.ZMap/ZMap` and will be picked up by default.

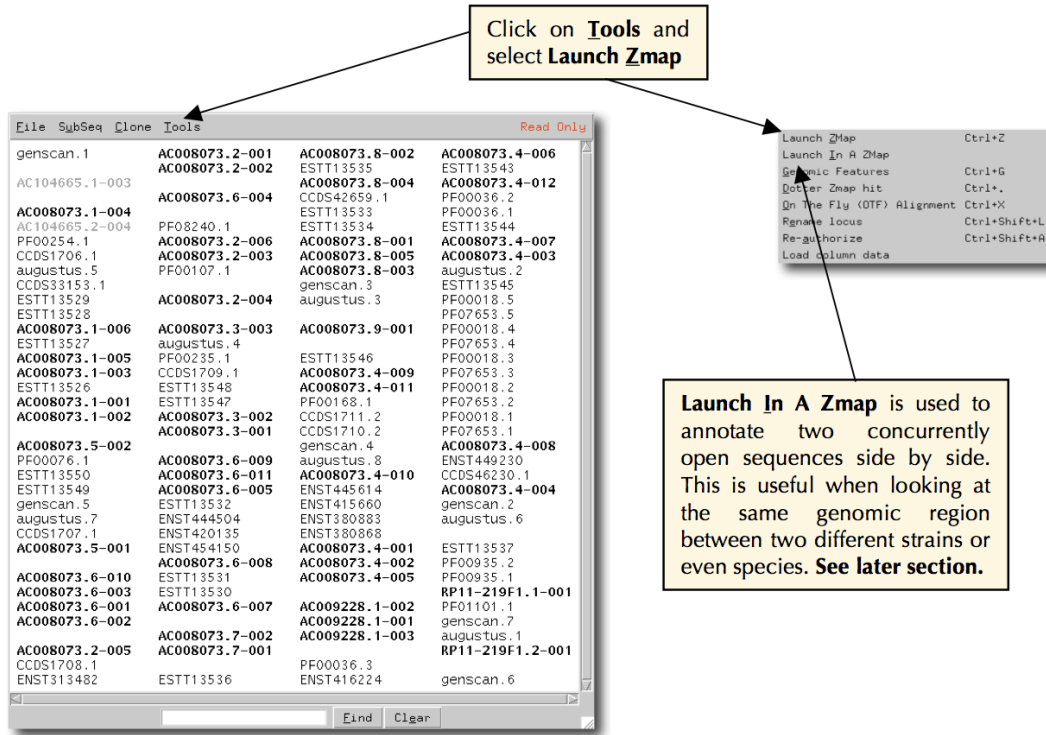


Figure 1: Opening ZMap from Otter

## Operation within Otter

ZMap is opened via the Tools menu bar in Otter (figure 1).

## Main ZMap interface

### Navigating in ZMap and zooming options

1. Navigate by using the scroll bars or the middle mouse button. By clicking the middle mouse anywhere in ZMap you will see a horizontal line. You can move this up and down and the relative position in bp will be displayed along the line. When the button is released, the window will refresh, centering on the position of the line. You can also click in the window to make it active and use the scroll wheel to navigate up and down or achieve the same result using the scroll bar on the right hand side of the window. If you release the mouse outside the ZMap window, you can then check the sequence position displayed, without re-centering. See Figure 3.
2. Zoom in by using the Zoom in/Zoom out buttons at the top, or by drawing a rectangle around the area of interest with the left mouse button. Use the "z" key on the keyboard to zoom to whatever feature is highlighted. Use the "Z" key to zoom to a whole transcript if you have an exon (s) highlighted or all HSPs if you have one HSP highlighted (HSPs are the "blocks" that you see in the homology columns, such as ESTs and protein hits). See figure 4.

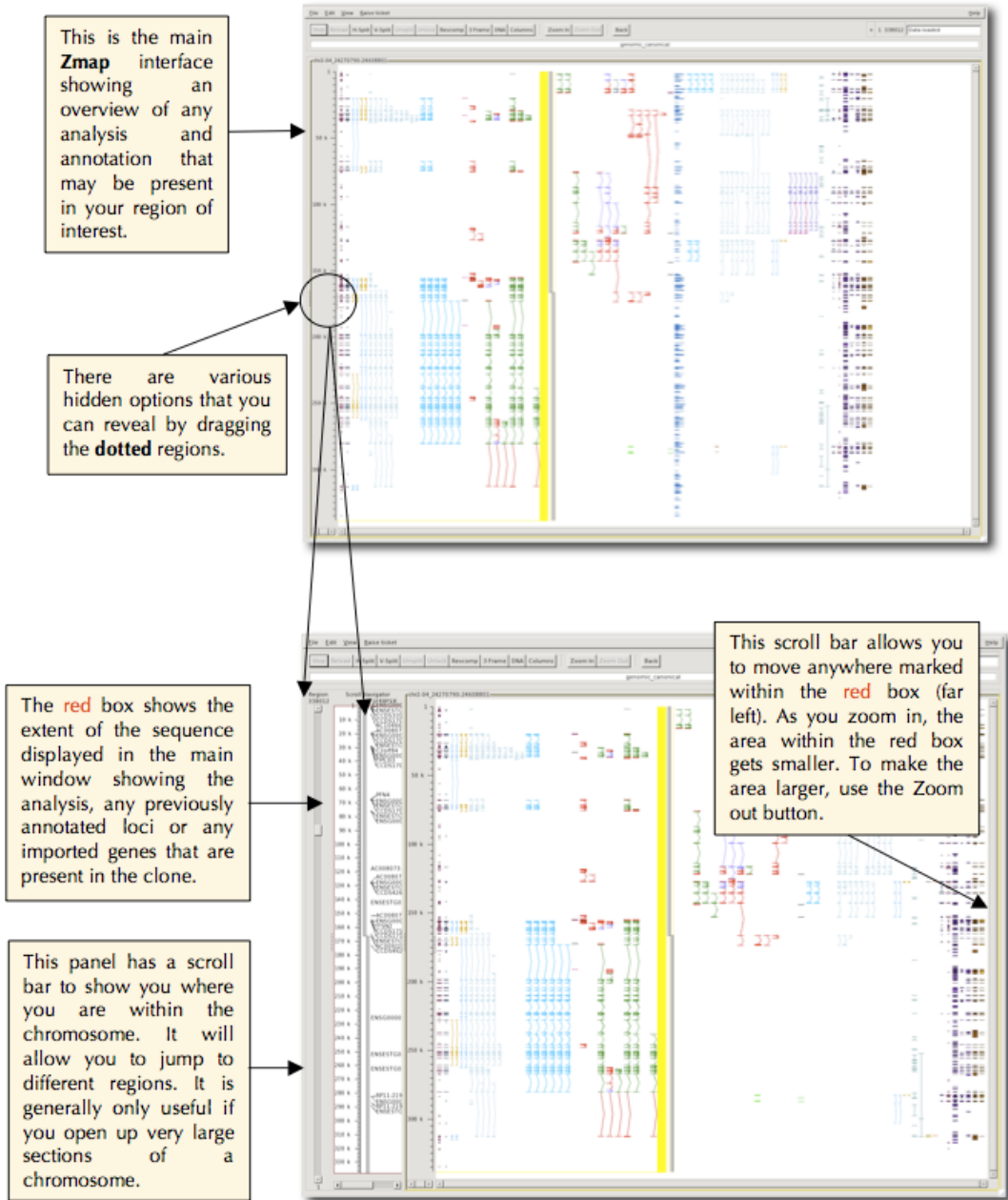


Figure 2: The main ZMap interface

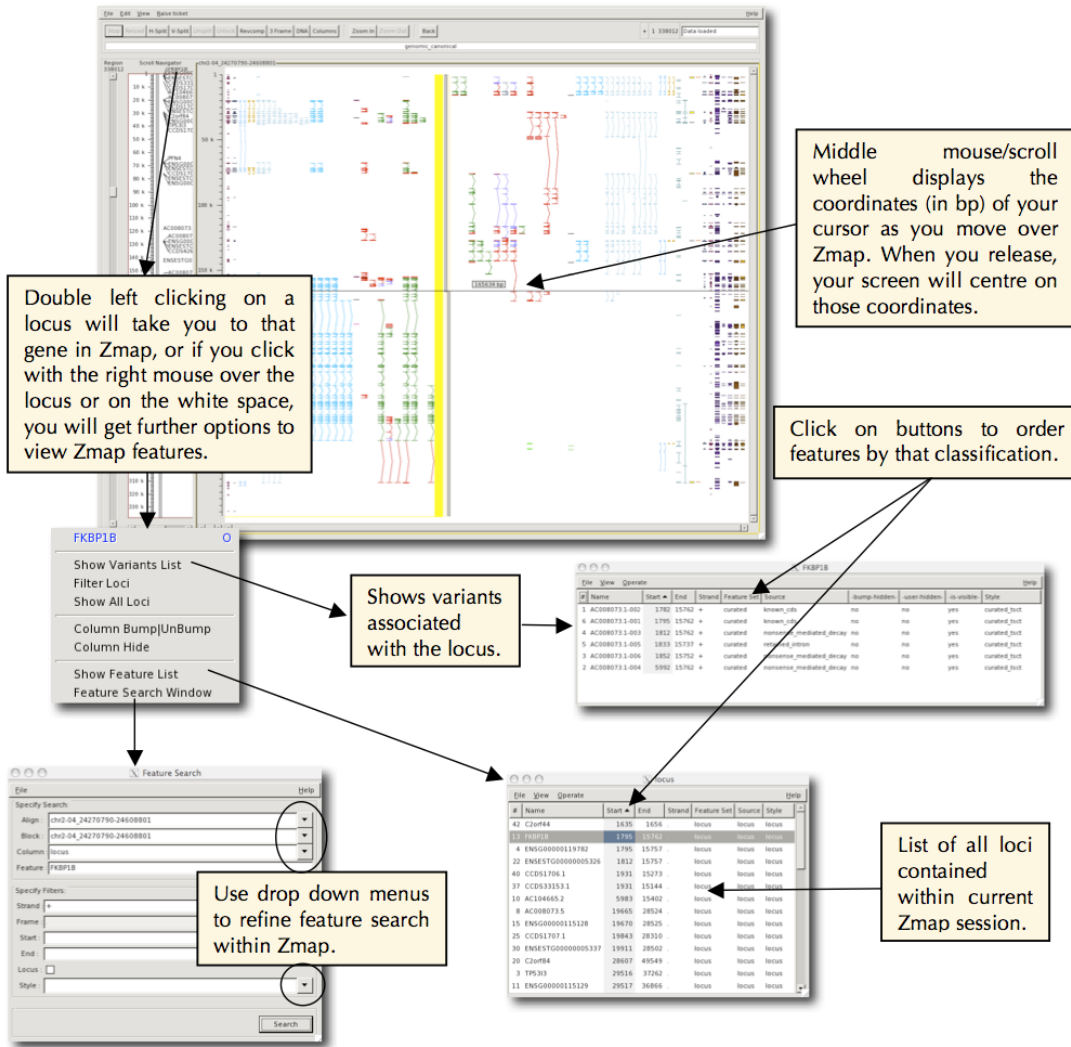


Figure 3: Navigating

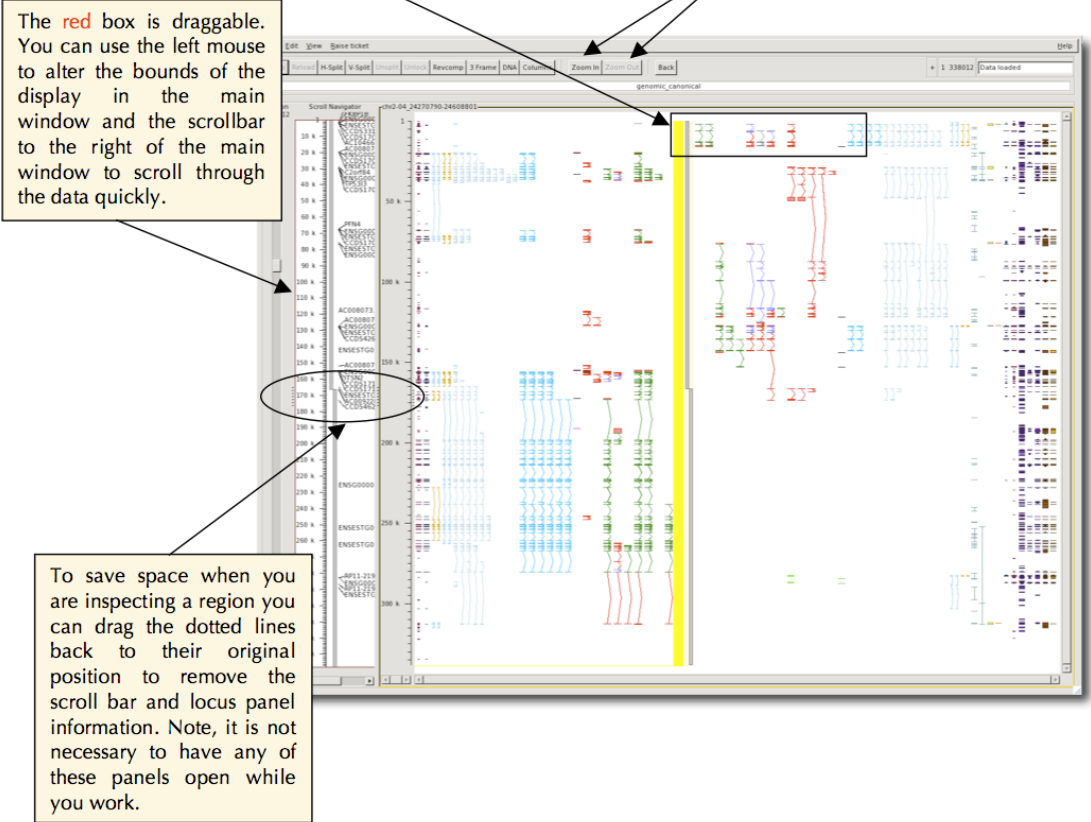


Figure 4: Zooming

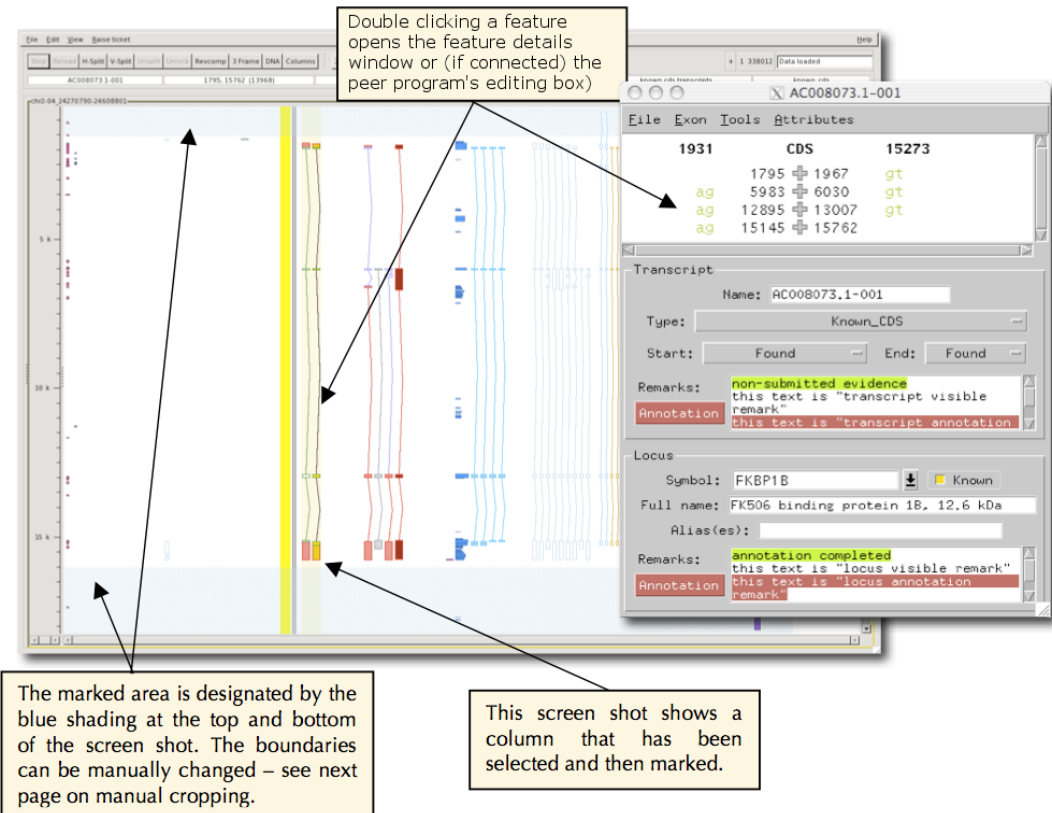


Figure 5: Focus and Mark

### The Focus Feature vs the Marked Feature

If you click on a column background then that column becomes the "focus" column and you can do various short cut operations on it such as pressing "b" to bump it. If you click on a feature then that feature becomes the "focus" feature and similarly you can do various short cut operations on it such as zooming in to it. (Note when you select a feature then its column automatically becomes the focus column.)

While the focus facility is useful, the focus changes every time you click on a new feature. Sometimes you want to select a "working" feature or area more permanently. To do this you can "mark" the feature or area and it will stay "marked" until you unmark it. "Marking" an area within ZMap to work on is essential, allowing you to work much faster. The "marked" area is left clear while the unmarked area above and below is marked with a blue overlay (see figure 5):

### Mark a feature

1. Select a feature to make it the focus feature.
2. Press "m" to mark the feature, the feature will be highlighted with a blue overlay.

Feature marking behaves differently according to the type of feature you highlighted prior to marking and according to whether you press "m" or "M" to do the marking:

1. If you press “m”, the mark is made around all features you have highlighted, e.g. a whole transcript, a single exon, several HSPs.
2. If you press “M” to do the marking around transcripts the whole transcript becomes the marked feature and the marked area extends from the start to the end of the transcript.
3. If you press “M” to do the marking around alignments all the HSPs for that alignment become the marked feature and the marked area extends from the start to the end of all the HSPs.
4. If you press “M” to do the marking around all other features: the feature becomes the marked feature and the marked area extends from the start to the end of the feature.
5. If no feature is selected but an area was selected using the left button rubberband then that area is marked.
6. If no feature or area is selected then the visible screen area minus a small top/bottom margin is marked.

### **Mark an area**

1. Select an area by holding down the left mouse button and dragging out a box to focus on that area.
2. Press "m" to mark the area.

### **Manual cropping of the marked borders**

You can manually change the borders of the marked area by putting your cursor over this area and using the cropping tool by clicking and holding with the left mouse button and dragging to make the area bigger or smaller.

### **Unmark a feature**

Press "m" or "M" again, i.e. the mark key toggles marking on and off.

### **General ZMap display features**

ZMap has rich feature display capabilities, which are highly configurable. Different features are displayed in distinct columns, and can be configured with many different styles as shown in figure 6.

1. The thick yellow line represents the genomic sequence; everything to the left represents the negative strand and everything to the right the positive strand. DNA matches (i.e. ESTs, mRNAs and RefSeq) and repeats have been configured so that they are all displayed to the right of the center, although they may align to either strand. Other features are configured to display in their relevant strand. The thin bar to the right is the clone that the genomic sequence is made up from. When connected to the Otter peer, double-clicking on this gives access to the DE editing window.
2. Annotated transcripts; green is coding (CDS), red is non-coding (UTR and transcript variants) and purple shows the "coding" region of NMD variants. Grey transcripts (see dotted line) contain exons outside the sequence slice being viewed and should not be confused with Halfwise hits.
3. Curated features, such as PolyA features are seen as horizontal black lines.

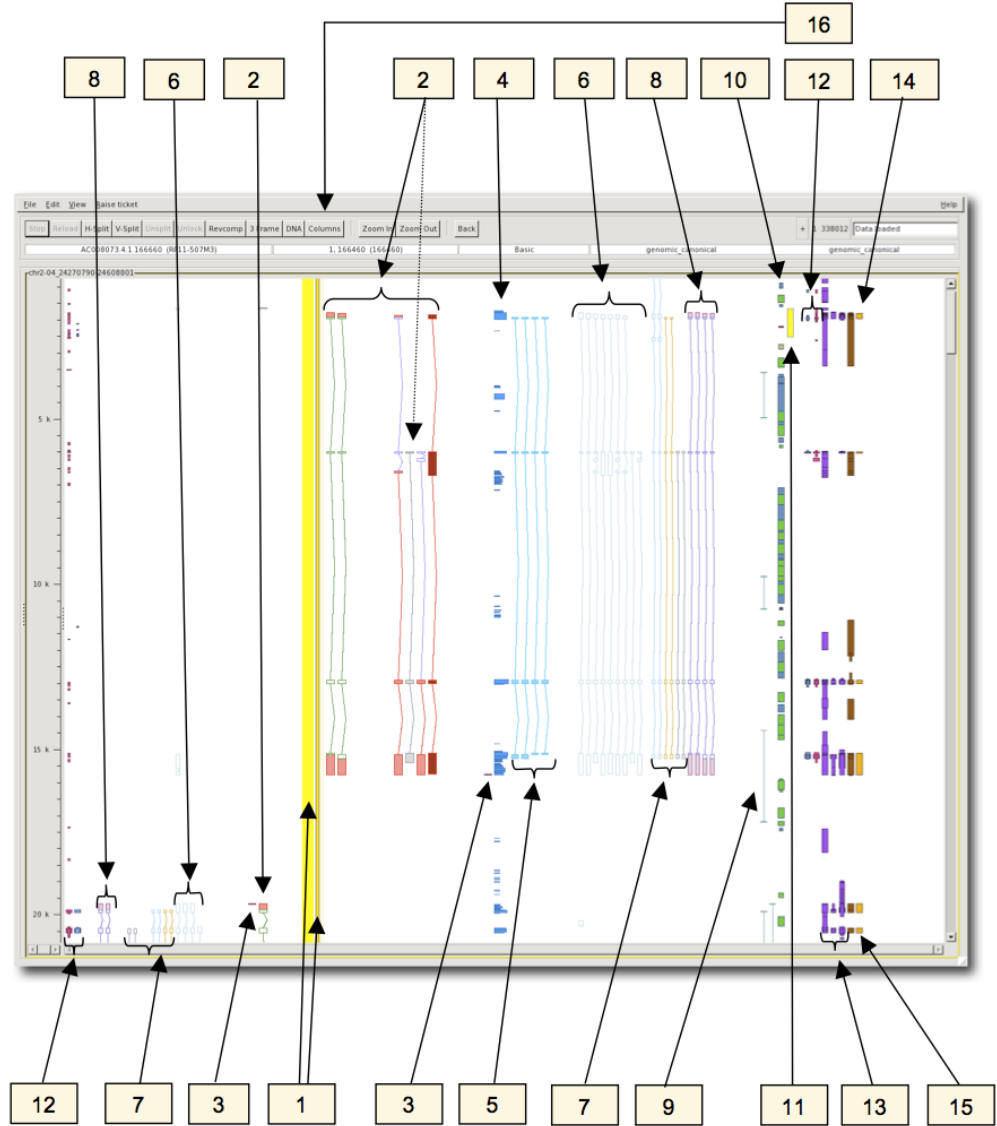


Figure 6: Features

## 16) Features and analysis available

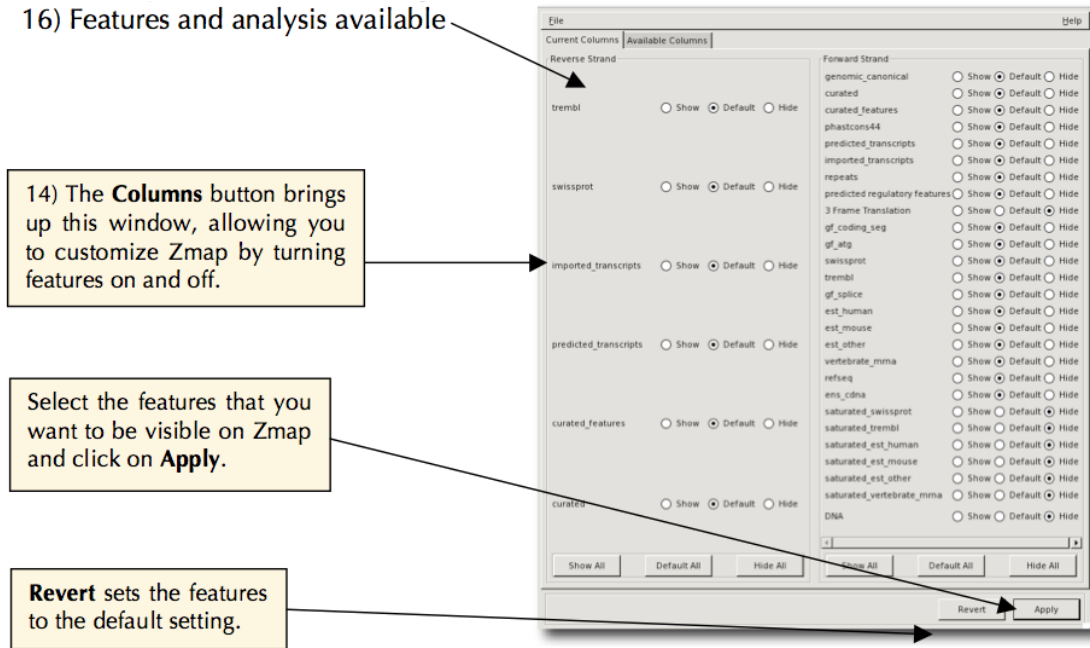


Figure 7: Columns dialog

4. Phastcons44 - conserved regions detected using multiple sequence alignments of 44 organisms.
5. Imported annotation from CCDS (human and mouse only).
6. Imported transcripts via DAS source. Here PASA\_ESTs are shown.
7. Predicted transcripts such as Genscan (pale blue), Augustus (gold) and Halfwise predictions of Pfam (grey).
8. Imported annotation from Ensembl.
9. gis\_pet\_ditags and chip\_pet\_ditags are indicators of transcript boundaries.
10. Repeats ( blue=Line , light green=Sine , gold=other ), tandem repeats are red.
11. CpG islands appear as yellow boxes.
12. Protein matches are strand specific - SwissProt are light blue and Trembl pink.
13. EST matches are displayed as purple blocks and are broken down into human ESTs, mouse ESTs, and other ESTs from other organisms. 5' reads are on the left and 3' on the right.
14. mRNA matches contains all species and are displayed as brown blocks,
15. RefSeq matches are the orange blocks.
16. Features and analysis available (see figure 7).

## Toolbar

Many functions are available from the toolbar section in ZMap. See figures 8 and 9.

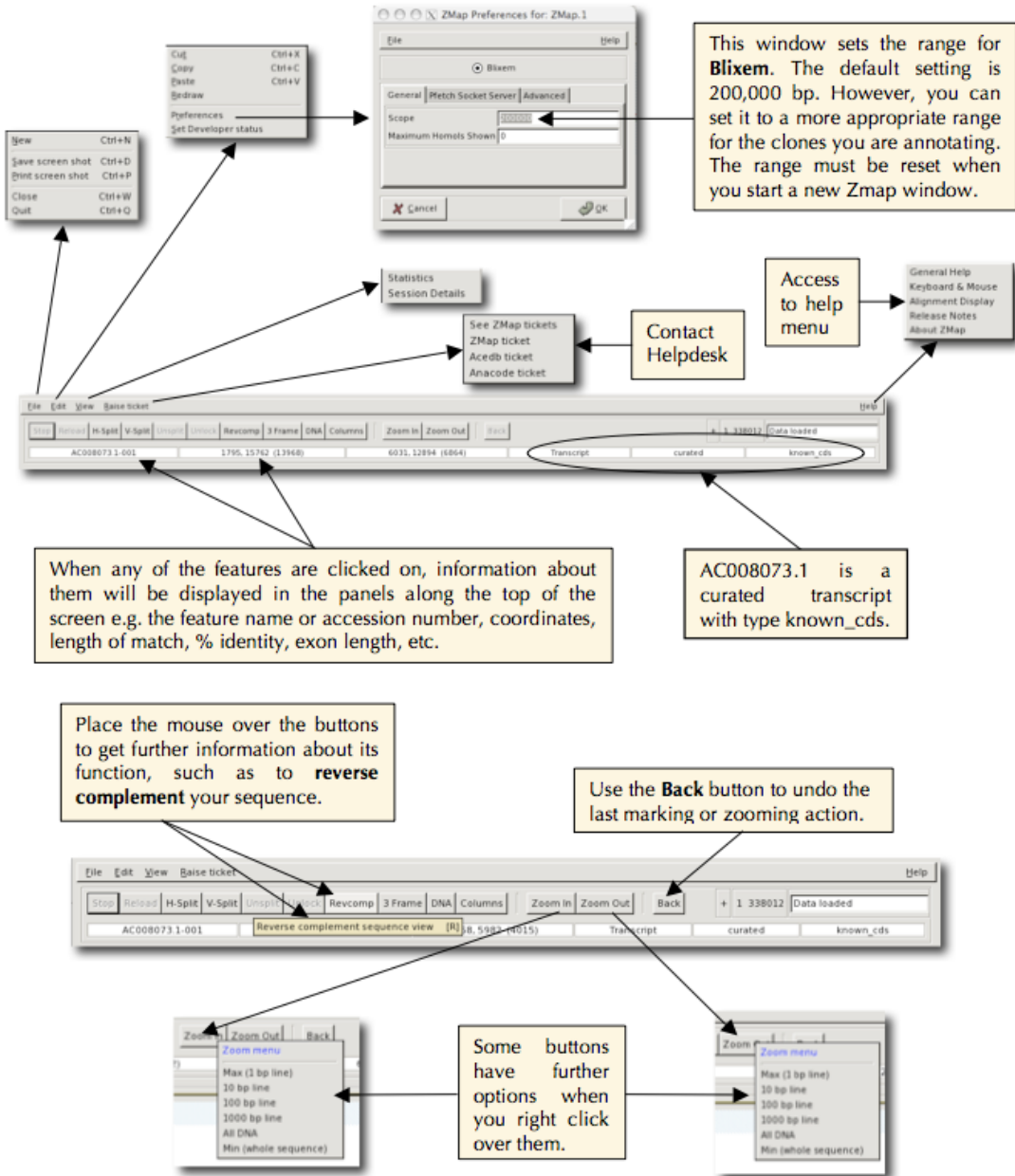


Figure 8: Toolbar feature details

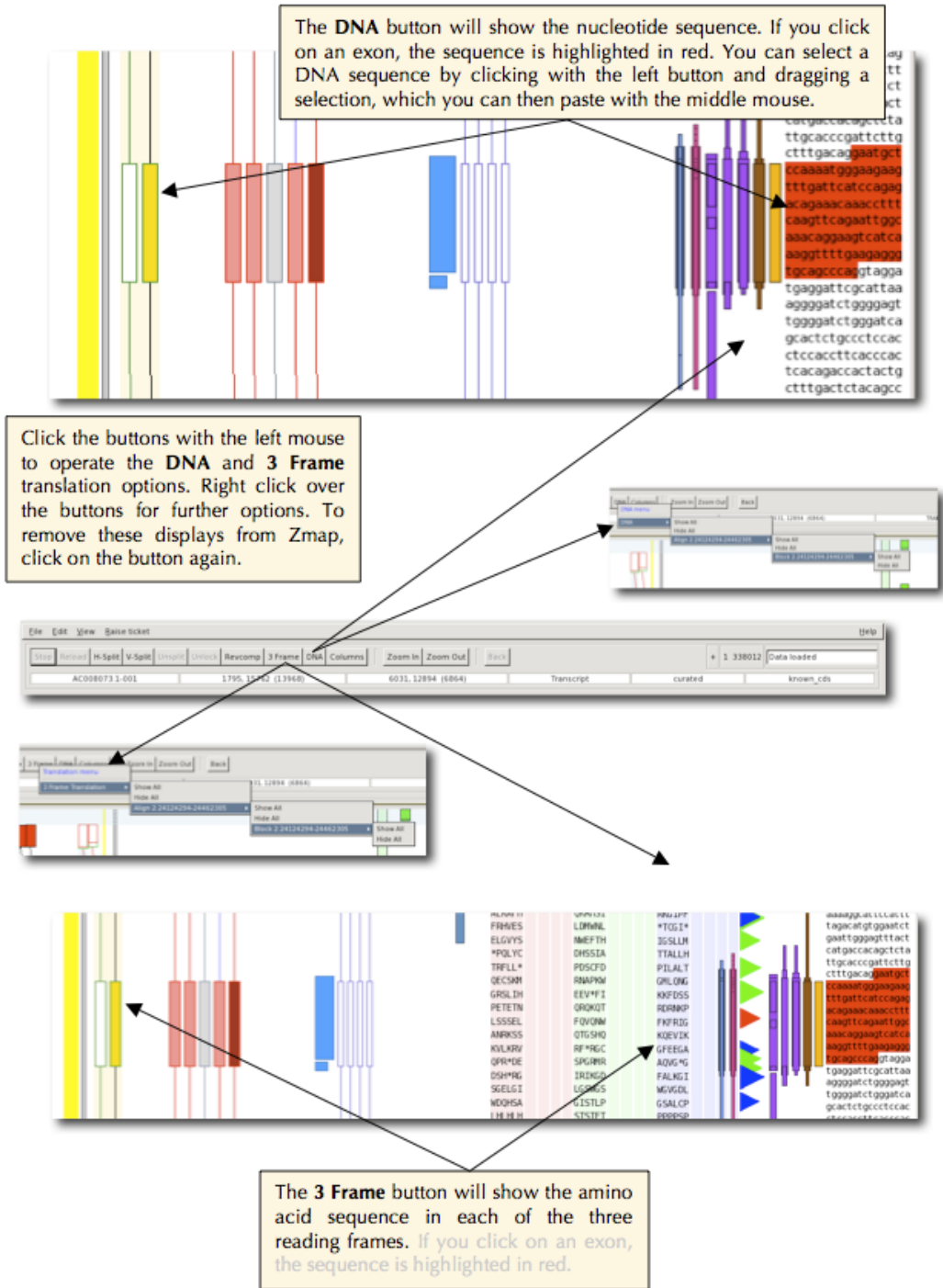


Figure 9: Feature display details

## Show feature details

Right click on a gene object or 'o' key when highlighted to see information on otter IDs and Ensembl IDs. For BLAST hits, double click on the HSP to get the feature interface where you will find details on alignment and on what gene object the HSP has been assigned to as evidence, if any (see figure 10).

## Exporting features for gene objects

Figure 11 shows how you can view and export an annotated sequence to your home directory in various different ways, such as dumping features directly.

1. In the main ZMap window, right click on an annotated gene object.
2. From the drop down menu select **Export Feature DNA** and choose sequence required from CDS, transcript, unspliced and with flanking sequence.
3. Alternatively, select **Export Feature peptide** and choose either CDS or transcript.
4. Figure 11 also shows the results of **Show Feature DNA** for annotated gene object AC008073.1-001 in FASTA format; firstly, the section of the transcript that corresponds to the CDS and secondly the whole transcript, including the untranslated region (UTR).

## Bumping features

This section describes how to select a feature, mark it and then zoom in to it and examine evidence that overlaps that feature. The default setting for ZMap is to show HSPs drawn on top of each other. This saves space on the canvas making it easier to see the general features of the region of interest. The bump option allows you to see the HSPs as multiple alignments.

1. Click on the feature you are interested in (perhaps a transcript)
2. Mark it by pressing "m"
3. Zoom in to the feature by pressing either "z" or "Z" (as described previously).

Now when you bump an evidence column to look at matches that overlap the feature you will find that bumping is much faster because only those matches that overlap the feature get bumped and you also have fewer matches to look at. The quickest way to bump a column is:

1. Click on the column to select it.
2. Bump it by pressing "b" (if you press "b" again the column will be unbumped). If you have marked a feature then bumping is restricted to matches that overlap that feature, otherwise bumping is for the whole column.

If you use the default bumping mode (i.e. you pressed "b") then you will find all matches from the same piece of evidence are joined by coloured bars, the colours indicate the level of colinearity between the matches (see figure 12).

1. **Green:** the matches at either end are perfectly contiguous, e.g. 100, 230 → 231, 351
2. **Orange:** the matches at either end are colinear but not perfect , e.g. 100, 230 → 297, 351. Matches may also be this color when there are extra bases in the alignment, e.g. around clone boundaries.

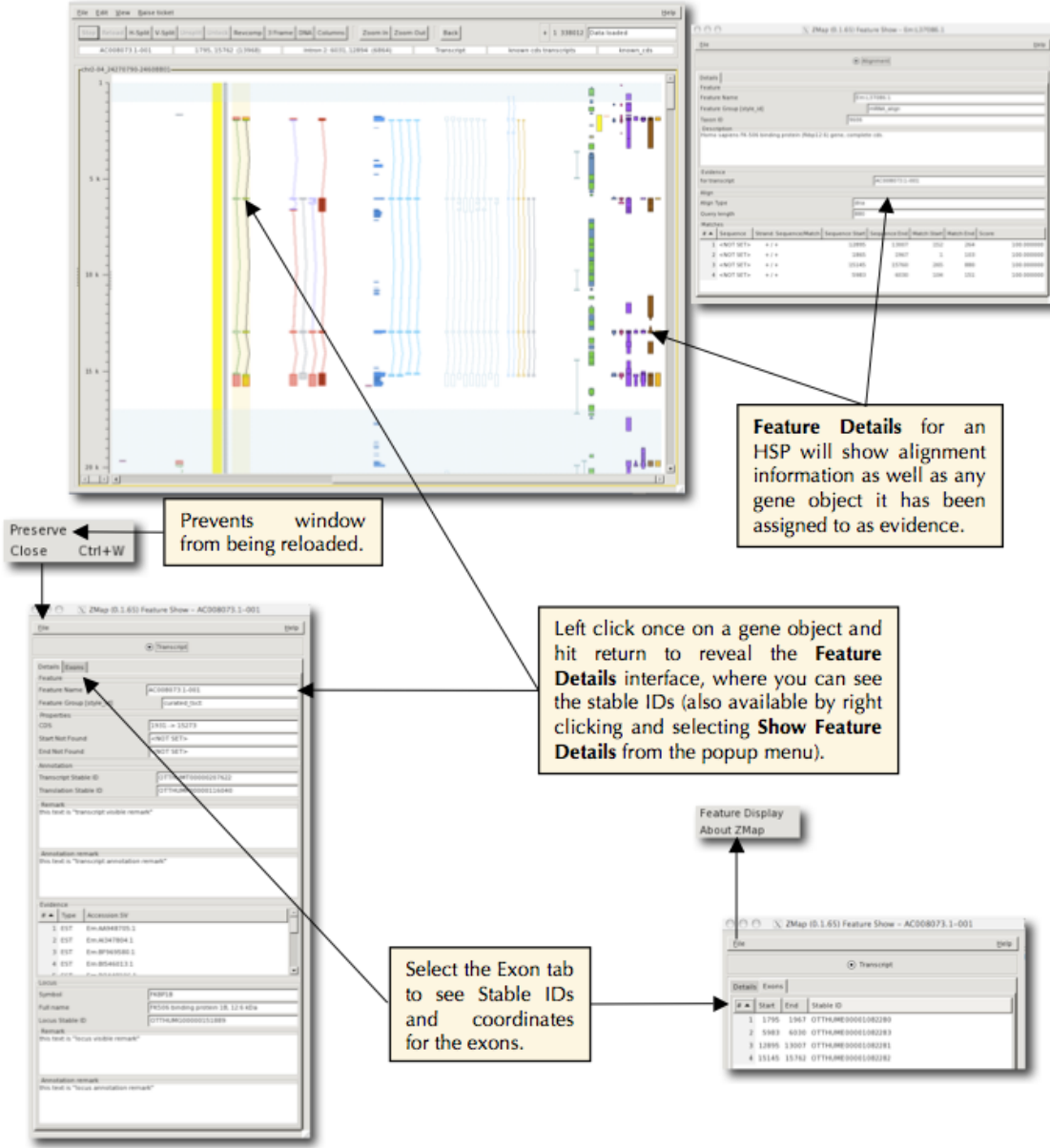


Figure 10: Feature details dialog

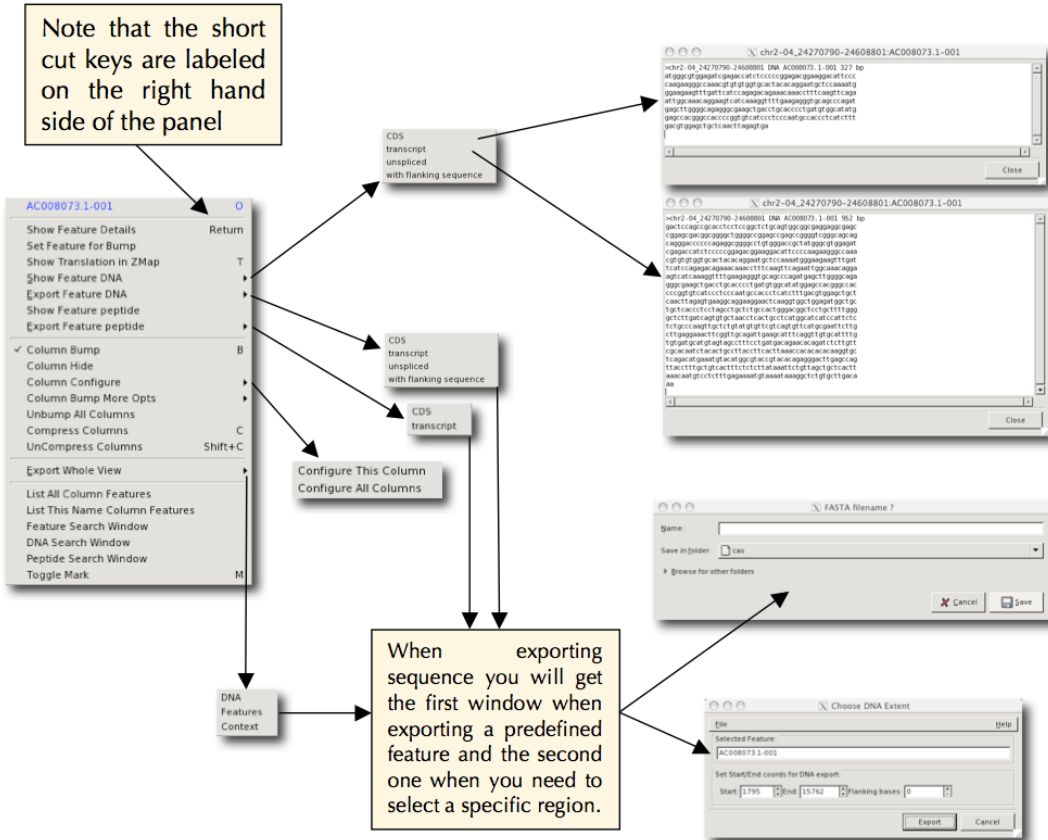


Figure 11: Exporting features

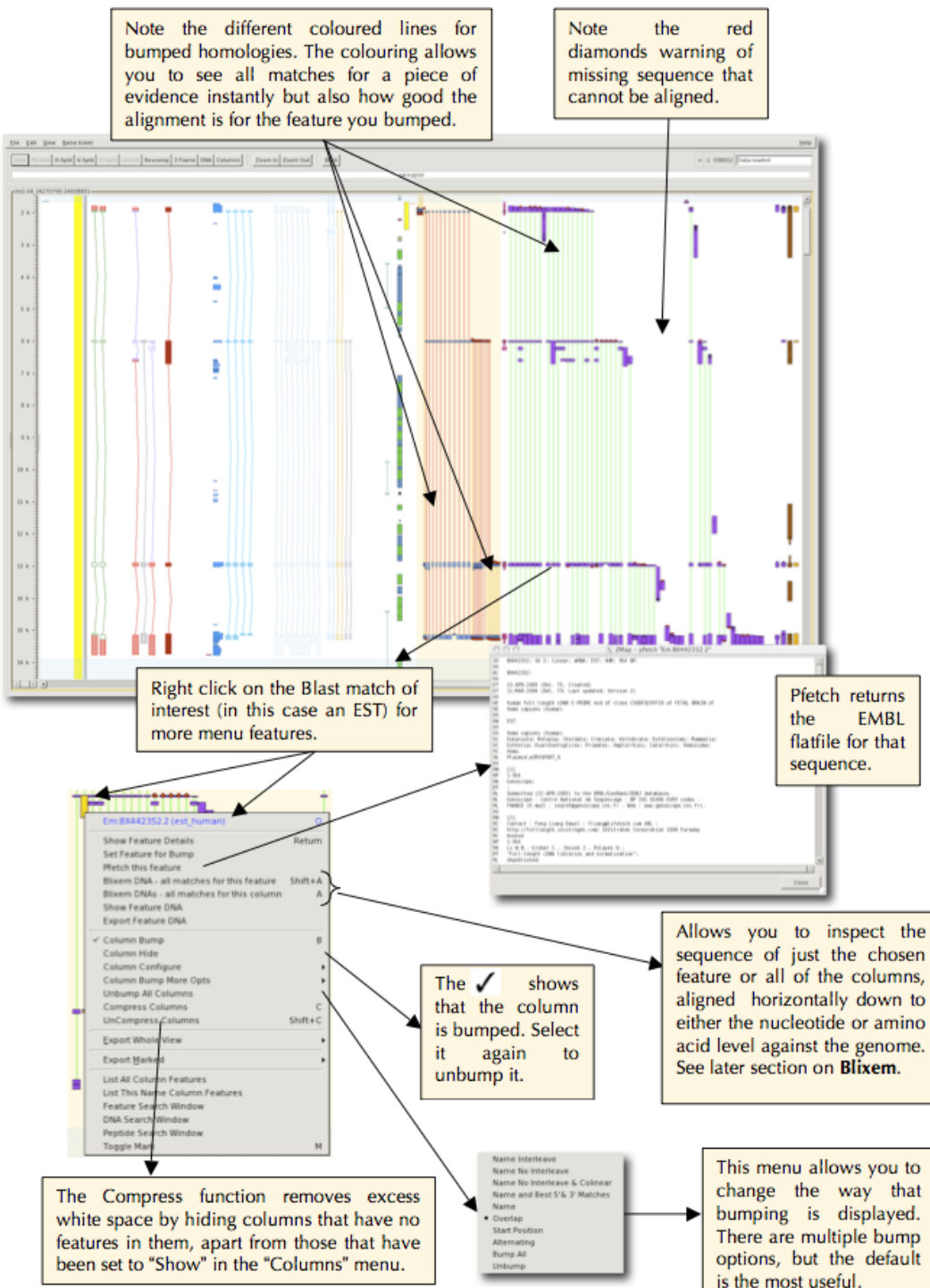


Figure 12: Bumping features

3. **Red:** the matches are not colinear, e.g. 100, 230 → 141, 423

Alignment quality of the HSPs is depicted by the width of every alignment displayed since the width is a measure of that HSP's score. Therefore, the wider it is the closer the score is to 100%. The precise score is displayed in the ZMap details bar by clicking on the alignment. If HSPs are missing either the first or last Blast alignments in the set, they are marked with a red diamond at their start/end respectively. This indicates if they do not start at the first base/amino acid and/or do not end with the last base/amino acid of the alignment sequence. The screen shot below shows what options you get when you right click over a homology - note that you can also select an HSP and type "o". You also get further options such as retrieving the EMBL file for that homology using pfetch and starting **Blixem**, see later section (note, HSPs do not need to be bumped to use Blixem).

### Searching for a sequence in ZMap

DNA and peptide search windows are provided from within ZMap and can be accessed by right clicking on ZMap space and selecting the option at the bottom of the menu. Both search windows are shown in figure 13.

### Searching for a feature in ZMap

This option allows you to list all the features contained in a column in one window. There are further options for you to search within these results to find a specific feature. The list of column features can be exported as a GFF file via the File menu. See figures 14 and 15.

### Selecting single or multiple features and hiding/showing them

1. If you left click once on a feature in ZMap, you will highlight all of its exons, the coordinates of which are now stored in the paste buffer and can be copied elsewhere, such as into the transcript editing window in Otter.
2. You can select multiple features by holding the Shift key down and left clicking with mouse (same as for multi select on the Mac, Windows etc). This option will highlight a single exon at a time for each feature, but the accession numbers of each feature and the individual exon coordinates are held in the paste buffer. This is a particularly useful way of selecting ZMap hits to use in the OTF alignment tool in Otter, as all selected homologies will be held in the paste buffer and automatically pasted into the OTF accession window. Each of the exon coordinates can also be pasted into the transcript editing window in Otter (see figure 16).
3. You can remove selected features in ZMap by pressing **Delete** on the keyboard and restore them by pressing **Shift-Delete** (note on the Mac you need to press **Fn-Delete** and **Shift-Fn-Delete**). This is a particularly useful way of removing evidence that you have already assigned to a transcript object.

### Creating/editing features

ZMap can be used to create new features or edit existing features. ZMap can generate variant objects quickly - existing transcript objects can be used as a template for a new object while a ZMap HSP (or any other feature, nucleotide or peptide) can be used to provide the coordinates for the new variant.

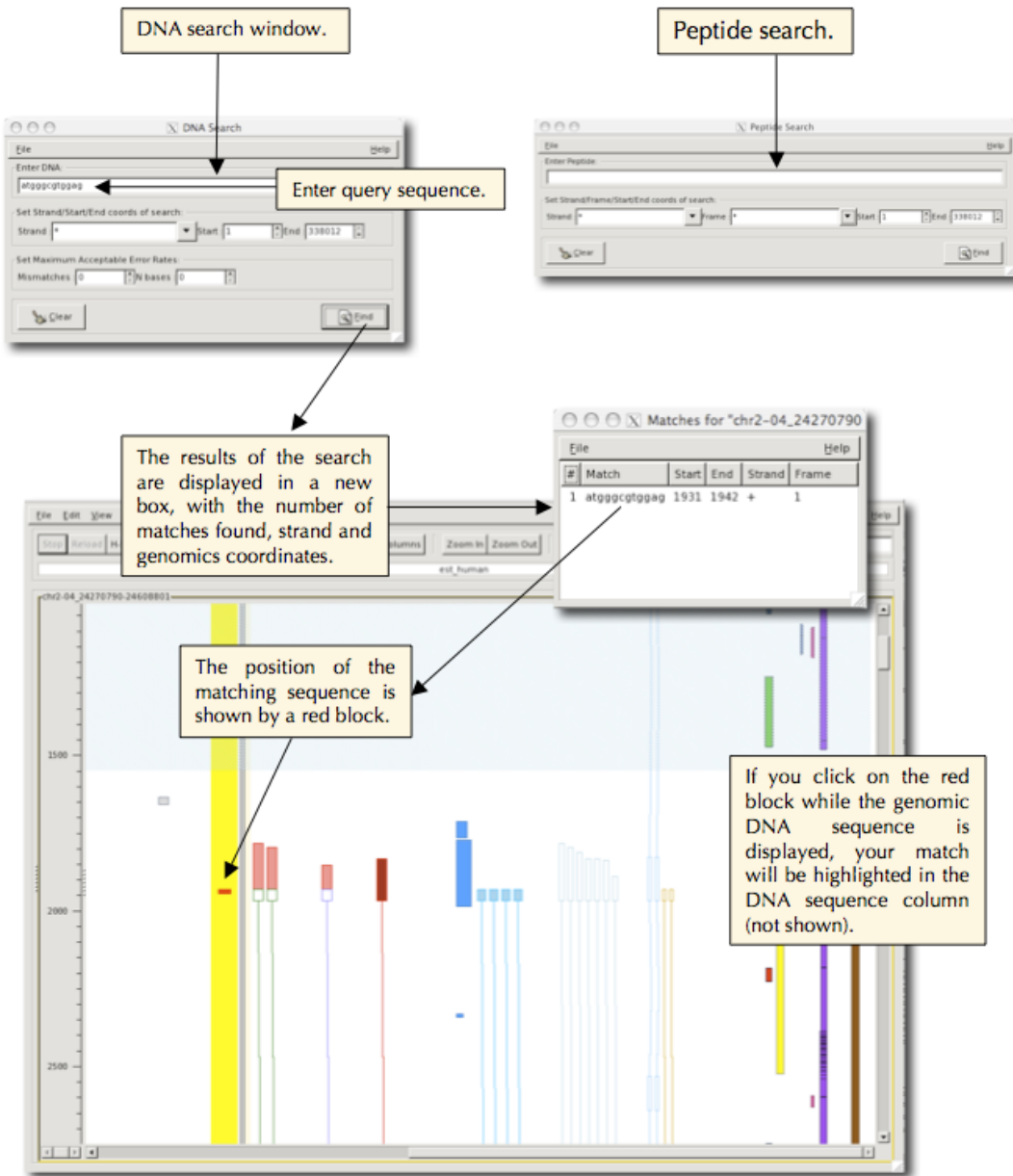


Figure 13: Searching for a sequence

vertebrate\_mrna

- Column Bump
- Column Hide
- Column Configure
- Column Bump More Opts
- Unbump All Columns
- Compress Columns
- UnCompress Columns

Shift+C

Export Whole View

Show Feature List

Feature Search Window

DNA Search Window

Peptide Search Window

Toggle Mark

Show Style

Blixem DNA Alignments

Blixem DNA Alignments - All Columns Shift+A

M

vertebrate\_mrna

#	Name	Start	End	Strand	Query Start	Query End	Query Strand	Score	Feature Set	Source	Mode
9	Em1U11671	131013	134286	-	745	4112	+	99.699997	vertebrate_mrna	vertebrate_mrna	mRNA_align
1	Em1U11671	138209	139400	-	554	745	+	100.000000	vertebrate_mrna	vertebrate_mrna	mRNA_align
4	Em1U11671	138877	139873	-	457	553	+	99.000000	vertebrate_mrna	vertebrate_mrna	mRNA_align
5	Em1U11671	208709	208930	-	335	456	+	100.000000	vertebrate_mrna	vertebrate_mrna	mRNA_align
7	Em1U11671	209017	209062	-	289	334	+	100.000000	vertebrate_mrna	vertebrate_mrna	mRNA_align
8	Em1U11671	209425	209581	-	122	286	+	100.000000	vertebrate_mrna	vertebrate_mrna	mRNA_align
2	Em1U11671	208447	208511	-	57	121	+	100.000000	vertebrate_mrna	vertebrate_mrna	mRNA_align
3	Em1U11671	209988	210093	-	23	56	+	100.000000	vertebrate_mrna	vertebrate_mrna	mRNA_align

Click over a column with the right mouse to activate this menu. Select Show feature List.

Export results as GFF file.

To search for a feature, enter your query here and click on search.

Note, the format needs to be correct for Zmap, so use \* as a wild card. For example accession numbers may have a database prefix and version suffix such as Em:U61167.1, so use the following format \*accession\_number\*, if you are not sure about the database and version.

This lists all the accession numbers and associated information for the column "vertebrate\_mrna". The results can be ordered using the buttons at the top.

The result lists all the exons and associated match information for query accession Em:U61167.1.

Figure 14: Searching for a feature

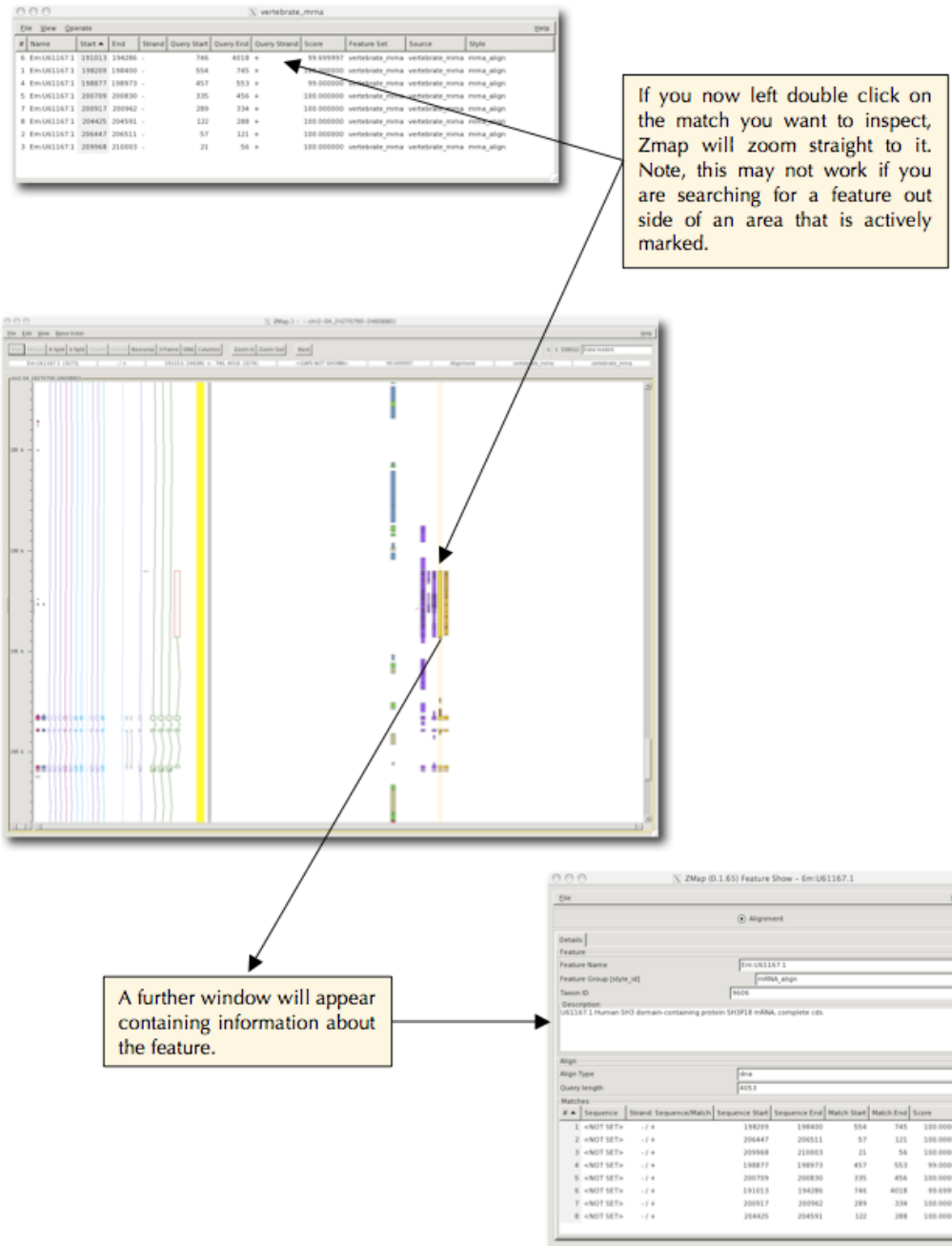


Figure 15: Feature search results

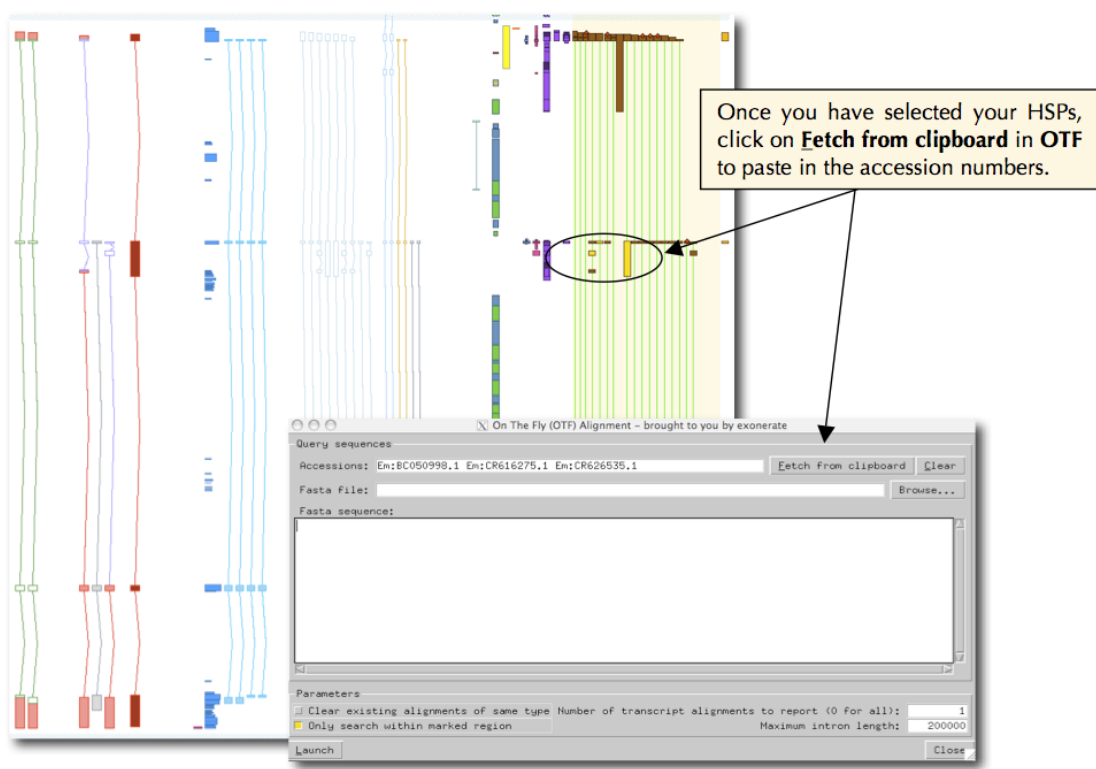


Figure 16: Pasting selected features to Otter's OTF dialog

## Standalone ZMap

Features are edited by copying them to a temporary feature in the **Annotation** column (see figure 17). The temporary feature can be based on any existing feature(s). Multiple features or nucleotide coords can be used to adjust feature/exon coordinates. The coordinates and other attributes can also be edited manually. When finished, the temporary feature can be saved to the relevant column - either as a new feature, or to overwrite an existing one.

The Annotation menu in the right-click menu also offers other options such as: clear the Annotation column; delete an exon/intron; undo/redo the last operation.

Right-click the temporary feature and select **Highlight evidence** to highlight the feature(s) that were used to construct it.

To create a new temporary feature, select the transcript (or other feature) you wish to create a variant from and press **Ctrl-K** (or right-click and select Annotation -> Copy selected transcript(s)<sup>1</sup>). Select other features and use Ctrl-K to copy those in to adjust the coordinates of the original feature. ZMap will do its best to make a sensible merge of the new coordinates, e.g. extending the feature extents if a coordinate lies outside the current range. If ZMap cannot automatically merge a coordinate it will ask for more information, e.g. if a coordinate lies within an intron, ZMap will ask whether it should form a 5' or 3' intron boundary.

**Double-click** the temporary feature to open the **Edit Feature** dialog to edit coordinates and other attributes manually. You can specify the new feature's final name and feature set. If you wish to save these attributes to the temporary feature (without saving the feature to the feature set) then click **Save Attributes**. To go ahead and save the feature to the named feature set, click **Create Feature**. Note that to save the feature back to file you also need to **Save/Export** the relevant feature(s) by going to the File menu and selecting one of the Save or Export menu options.

## Variant construction with Otter

When running ZMap under Otter, variants are constructed in a different manner. See figures 18 and 19.

## Splitting windows in ZMap

Use the **split** window function to effectively reduce the size of the window when looking at homologies. This is of particular use when you have to deal with very large introns because you can essentially reduce the introns to whatever size you wish, or when there are very many HSPs, because you can keep your gene object in view and static, but still scroll across the evidence. See figure 20.

## Launching in a ZMap from Otter

This function allows you to open two or more sequences alongside each other (such as a human region and the syntenic region in mouse, or two haplotypes), so that simultaneous investigation can be carried out. To do this you will need to open both sets of clones in the same Otter session. To open both ZMap windows in one window as shown below, you need to select "Launch In A ZMap" option in one clone set. These clones will open to the left of the already open Otter session. This screen shot shows human gene SF3B14 and the syntenic region in mouse. The gene copy and paste function (referred to in the Otter section) is of much use here, saving time when building gene objects. See figure 21.

<sup>1</sup>Note that you need to enable the Annotation column from the Preferences dialog for the Annotation menu to appear. Ctrl-K will automatically enable the Annotation column if it is not already enabled.

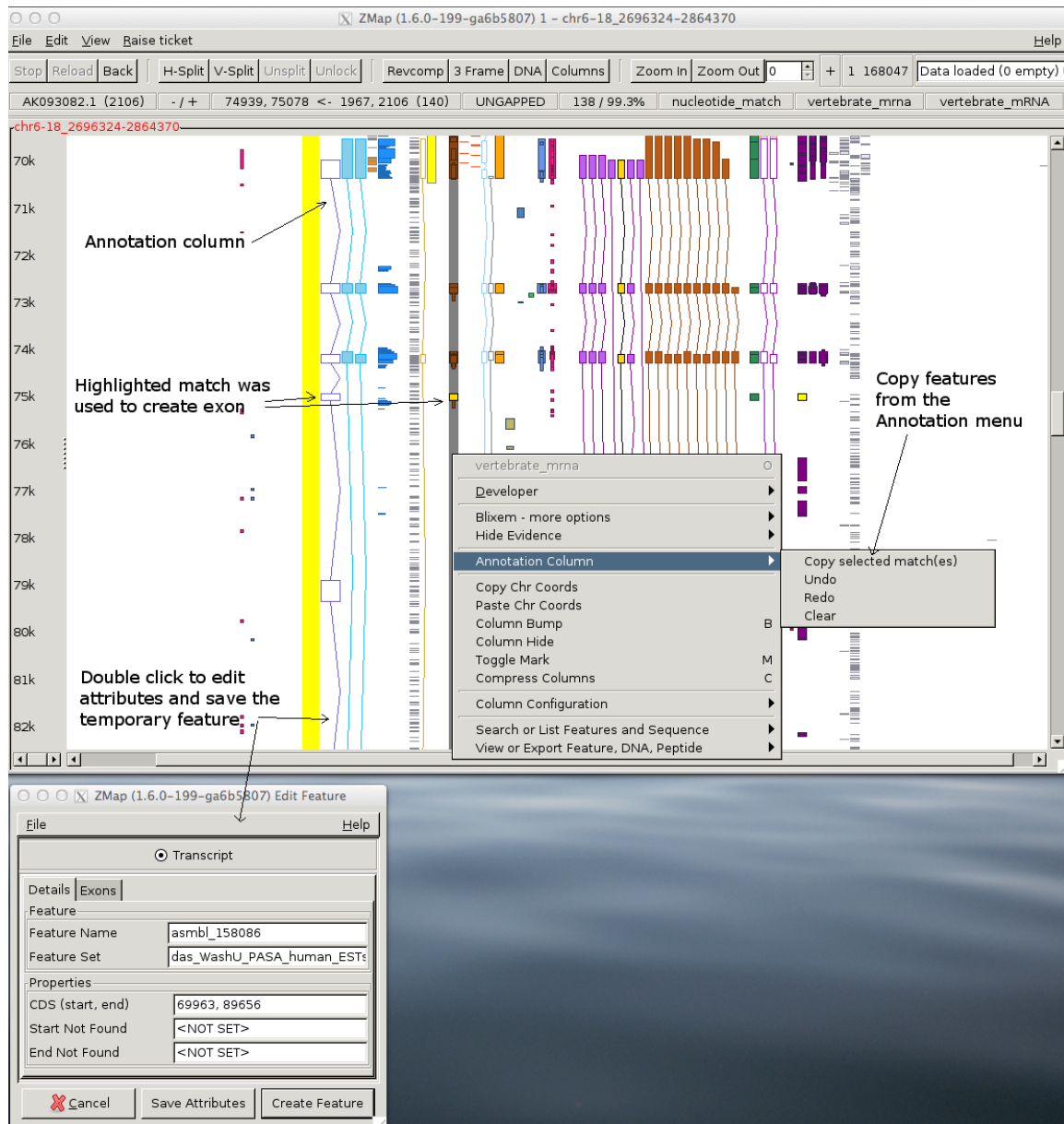
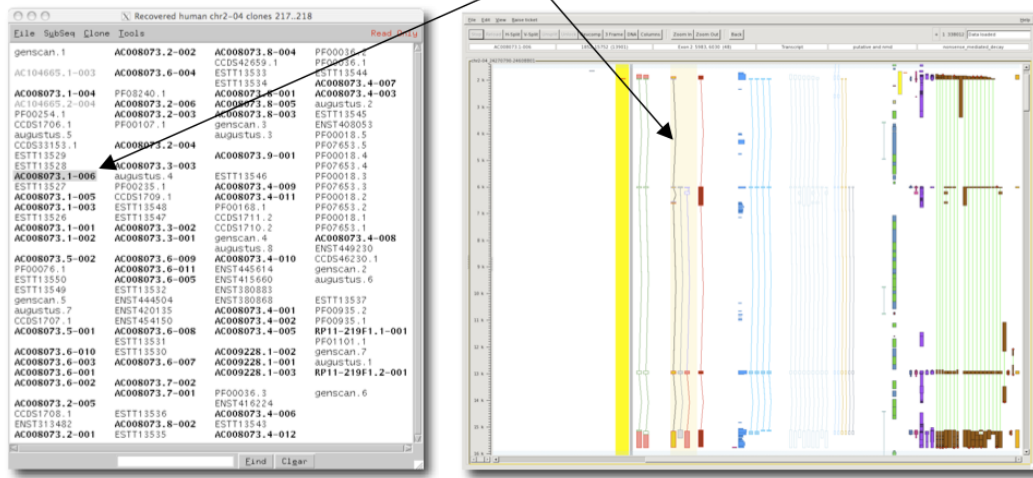


Figure 17: Feature editing in ZMap

1) Select the object that will form the foundation to the new variant, either by highlighting the object in Otterlace or clicking on the object in Zmap.



2) Click on the HSP that will give its coordinates to the new variant object.

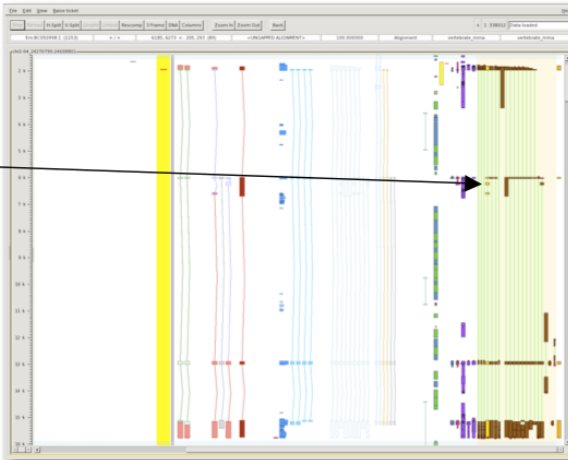


Figure 18: Variant construction in Otter (1)

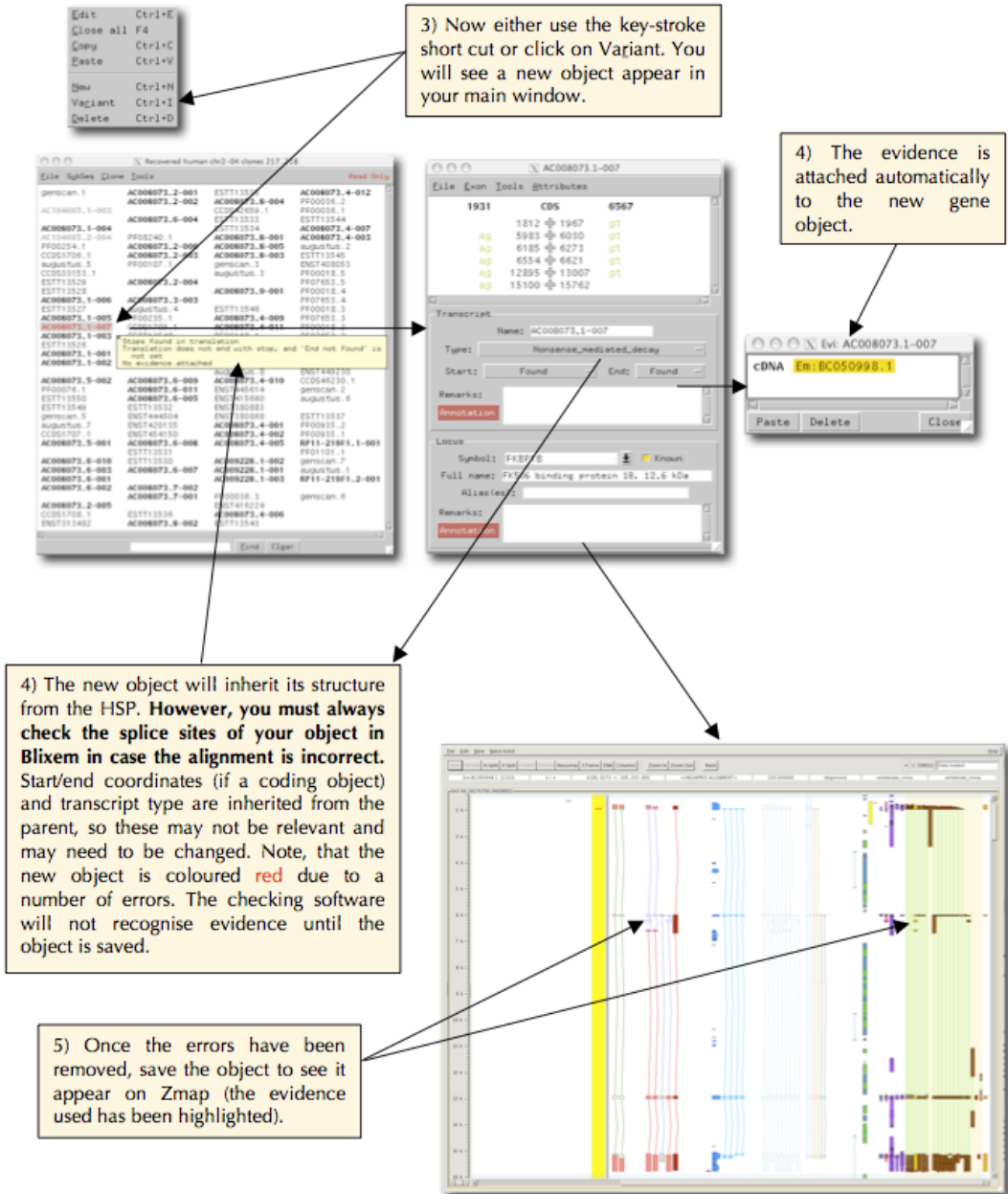


Figure 19: Variant construction in Otter (2)

The screen can be split horizontally or vertically (as shown) multiple times. An active window must be selected for **splitting**.

**Unsplit** will remove the last split window.

The windows will be locked together when you first open them. To scroll independently within each window, use the **Unlock** button.



Figure 20: Splitting windows

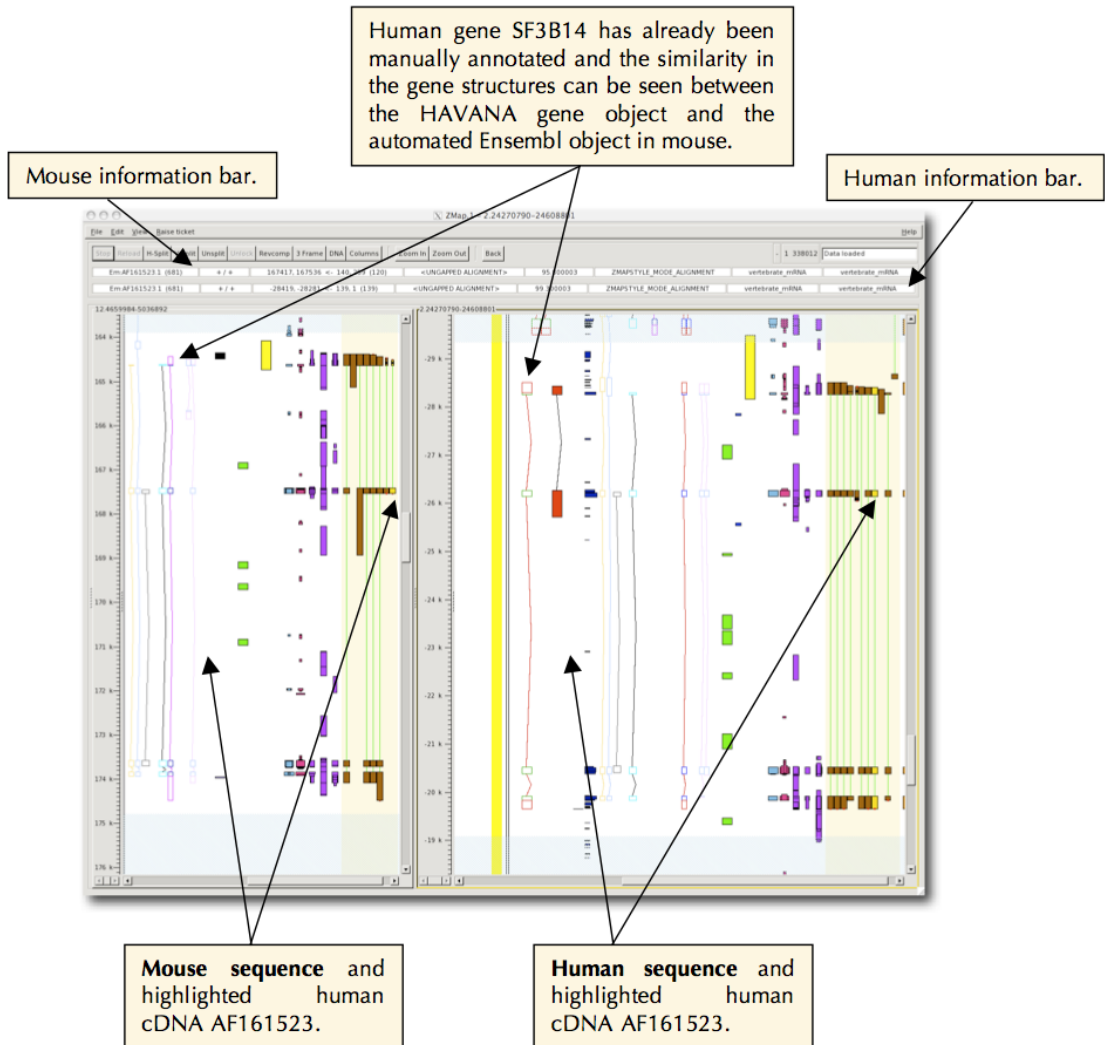


Figure 21: Launch in an existing ZMap

## Keyboard and mouse shortcuts

In general ZMap will be faster for zooming, bumping etc if you make good use of the built in short cuts. These can often avoid the need for ZMap to redraw large amounts of data that you may not even be interested in. For example, click once (highlight) on a feature and a carriage return will bring up evidence. Another example is to press T for translation.

### All windows

Short Cut	Action
Cntl-W	close this window
Cntl-Q	quit ZMap

### ZMap Window

Short Cut	Action
Control keys	
+ (or =), -	zoom in/out by 10%
Cntl + (or =), Cntl -	zoom in/out by 50%
up-arrow, down-arrow	scroll up/down slowly bit
Cntl up-arrow, Cntl down-arrow	scroll up/down more quickly
left-arrow, right-arrow	scroll left/right slowly
Cntl left-arrow, Cntl right-arrow	scroll left/right more quickly
page-up, page-down (Mac users should use fn and up/down arrow)	up/down by half a "page"
Cntl page-up, Cntl page-down	up/down by a whole "page"
Home, End (Mac users should use fn and left/rights arrows)	Go to far left or right
Cntl Home, Cntl End (Mac users will have to configure their keyboards for this)	Go to top or bottom
Delete, Shift Delete	Hide/Show selected features.
Enter	Show feature details for highlighted feature.
Shift up-arrow, Shift down-arrow	Jump from feature to feature within a column.
Shift left-arrow, Shift right-arrow	Jump from column to column.

### Alpha-numeric keys

Short Cut	Action
a	Blixem all sequences in column
A	Blixem only highlighted sequence in column
b	Bump/unbump current column within limits of mark if set, otherwise bump the whole column.

B	Bump/unBump current column within limits of the visible feature range.
c	compress/uncompress columns: hides columns that have no features in them either within the marked region or if there is no marked region within the range displayed on screen. Note that columns set to "Show" will not be hidden.
C	Compress/unCompress columns: hides all columns that have no features in them within the range displayed on screen regardless of any column, zoom, mark etc. settings.
h	Toggles highlighting (good for screen shots).
m	mark/unmark a range which spans whichever features or subparts of features are currently selected for zooming/smарт bumping
M	Mark/unMark the whole feature corresponding to the currently selected subpart (e.g. the whole transcript of an exon or all HSPs of the same sequence as the highlighted one) for zooming/smарт bumping
o or O	show menu Options for highlighted feature or column, use cursor keys to move through menu, press ESC to cancel menu.
r	reverse complement current view, complement is done for all windows of current view.
t or T	translate highlighted item, T hides Translation.
w or W	zoom out to show whole sequence
z	zoom to the extent of any selected features (e.g. exon/introns, HSPs etc) or any rubberbanded area if there was one.
Z	Zoom to whole transcript or all HSPs of a selected feature.

### ZMap Mouse Usage

Left	Middle	Right
<i>Single mouse button click</i>		
highlight a feature or column Plus drag: draw a rectangle around an object for zoom	horizontal ruler with sequence position displayed, on button release centre on mouse position. Release mouse outside ZMap window to prevent recentering.	show feature or column menu - for options such as pfetch, show feature DNA, show peptide, export peptide
<i>Double mouse button click</i>		
display details of selected feature. Double click on object to get edit window	same as single click	same as single click
<i>Shift + mouse button click</i>		
highlight a subpart of a feature (e.g. a single exon or alignment match) OR multiple highlight	same as single click	same as single click

### Tips for a speedier ZMap

1. Specifically: zoom and mark within ZMap early on after launching. Either select a gene object and press 'z' to zoom OR select a rectangle to zoom in by dragging the left mouse button around it. Reverse complement now if necessary, then press 'm' to mark the region.
2. The quickest way to zoom out of ZMap again is to right mouse click on the 'zoom out' buttons

at the top of zmap and choose one of the options (this is definitely much quicker than doing individual 'zoom outs' with the left mouse button). Likewise for 'zooming in' again (or use keyboard equivalents).

3. Bump within a marked region only. Bumping without marking is slow and removes the lines connecting Blast matches.
4. When you have finished working within a marked region, unbump the evidence you have been working on (e.g. ESTs) and unmark that region before you go on to select the next region to mark and bump - or you could miss visualising the evidence in the new region.
5. If you want to get rid of some white space try the compress 'c' function or alternatively toggle off some of the columns. **Warning - this may hide features as well.** If a column (e.g ESTs) is bumped and you want to lose it temporarily, it is quicker to turn the column off (when you turn it on again it will still be bumped when it reappears) than unbump then rebump again later.
6. Jumping to genes/objects: If you expand the left hand 'scroll navigator' overview' you can jump directly to genes and objects by double-clicking on them.