

# **A Tool for Visualising Cell Model Results**

## **MInf Project Phase 2**

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### **Abstract**

This will be the abstract

# Acknowledgements

Acknowledgements go here.

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# Chapter 1

## Introduction

Biologists often use computer models to help guide their research as modelling is much cheaper than experimentation. There are a number of tools available for biological modelling. These tools typically require a certain level of numerical confidence to create and interpret. Not all biologists have this numerical confidence. Some researchers find writing and interpreting models a challenge; this can make them less effective in their research. It is therefore necessary for the tools they use to help them relate the data to their field by incorporating domain knowledge.

One such tool that can be used for modelling is Bio-PEPA [1], an extension of the PEPA process algebra. Bio-PEPA is currently implemented as a plugin for the Eclipse IDE. Bio-PEPA visualises the model results as time-series graphs. There is one team of Src researchers in particular, who use Bio-PEPA and do not have the numerical confidence, as described above, to be comfortable using Bio-PEPA. This team was the original focus for the project. The purpose of this project was to extend Bio-PEPA's visualisation capabilities to allow the previously mentioned team, and other similar users of Bio-PEPA, to more effectively analyse their results.

A significant problem with Bio-PEPA's visualisation capability is that it is difficult to represent spatial change on a time-series graph (as can be seen in Figure 1.1). In Bio-PEPA you can have a species at different locations in the cell, for example, next to the nucleus, next to the cell membrane and throughout the cytoplasm. The movement of this species through the cell can be modelled by seeing the population of it in each location over time. This is difficult to visualise on a time series plot as three lines are too abstract. It requires the use of a biological metaphor for spatial information to be easily interpreted. In this case using a visualisation based on a cell can more intuitively show how the species moves through the cell. It is this type of inference that the Src researchers find challenging to do with Bio-PEPA currently.

Over the course of the project the scope has been expanded. The original objective was to assess which forms of visual representation are most helpful and informative to laboratory science. At the end of the first project phase the objective changed (to reflect that the project was about delivering a finished program and not about the results of many experiments) to be to develop a tool to visualise the results of dynamic

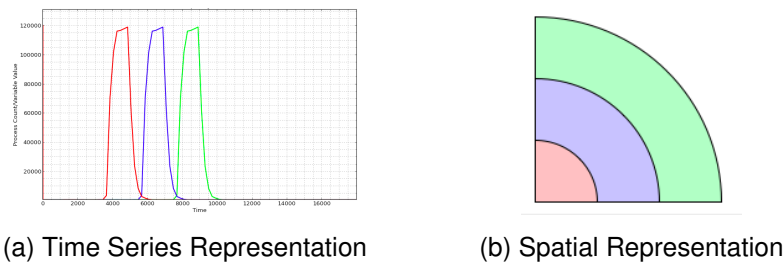


Figure 1.1: One species at three locations in the cell represented traditionally on a time series graph and also spatially in a cell

time-series models of intra-cellular behaviour based on biochemical reactions. This objective was focused on visualisation to aid those researchers who are not numerically confident. The second phase of the project has added to this objective to also aid interpretation and collaboration. This change in objective is to make the tool better for all users. To reflect these changes the focus on the original Src team has been generalised. Evaluations are performed with other potential users who are also biologists. Another factor in deciding to focus more on these other potential users was one of the key Src researchers going on maternity leave during the second phase of the project. This made it infeasible to evaluate the tool with them. It had been hoped to still evaluate the tool with them at the end of the second phase of development but they were still on maternity leave.

## 1.1 Where Does This Tool Fit In?

In the first stage of the project a review was performed of the features of a number of modelling and visualisation tools. This review included specialised software aimed at biologists and general software for anybody doing data visualisation.

As the project scope expanded to include goals not specifically related to visualisation it was prudent to perform another software review covering the new features, in particular software that allowed for collaborative editing. It is important to see what features are commonplace, which features are not commonplace but are useful, and which features are not useful. This analysis would then be used to guide development of the collaborative features of the new tool.

Four products were chosen for this second review: Google Drive [2], Pidoco [3], Lucid Chart [4] and SynBiowave [5]. Analysis of these software can be found below.

### 1.1.1 Previously Reviewed Software

The software packages reviewed at the start of the project were: Bio-PEPA, Uppaal [6], V-Cell [7], Cell-O [8], Copasi [9], Cell Designer [10] and WEAVE [11]. Bio-PEPA, V-Cell, Cell-O, Copasi and Cell Designer have been written for biological modelling.

Uppaal is modelling software for general use, and WEAVE is a general data visualisation tool. A review of these tools can be found in MPP1 report for this project.

cite this?

All offered some level of visualisation, some simply graphs, others more complex visualisations. Bio-PEPA offered only line graphs. Uppaal had visualisations that highlighted where in a finite state machine view of the model the current state is. V-Cell had visualisation of the model in a hierarchical set of circles. It could also display spatial elements of the results data by displaying a heat model view of the cell. Cell-O was aimed more at multi-cell models and was able to show them moving and splitting; it also had visualisation of the model as a finite state machine. Copasi only had graphs, although the user had more control over the display of the plots. WEAVE had the largest visualisation capacity being able to display a variety of standard graph types along with more interesting ones, such as geographical maps, but it did not appear to have anything specialised for biological models. WEAVE is also the tool that gave the user the most control over the visualisation.

The existing biological modelling software seems to be focused more on the ease of modelling. The visualisation features on offer are typically quite basic. They also lack the more innovative features that can be found in the newer general data visualisation software.

### 1.1.2 Google Drive

Google Drive which was previously Google Docs, is one of the most widely used collaborative pieces of software. Google Drive is used by a variety of user types. The focus of the review was on the word processor. Of the collaborative software reviewed this was felt to be the most user friendly. One of the nicest features was a cursor indicating where every user currently editing the document is typing. Each user is also given a different colour allowing you to know who is doing what in real time. Many people can edit a single document at once. As well as editing documents users can also leave comments on the document. Changes made by users appear near instantly to all users. The speed of editing is very important as it is frustrating as a user to have to wait to see changes others are making. It is also very easy to invite others to edit the document with you. Each document has a unique link and if a user visits that link they are taken to the document and can start editing. Different permissions can be granted to different users allowing some level of collaboration with people who you don't necessarily want to grant full write access. These users can then just look at it in real time and offer comments. Different parts of the editor have different levels of collaboration. All text that is changed is changed for all, but preferences like font choice are only changed for the user, unless another user edits any text. Also importantly from a User Experience (UX) perspective is that any conflicts that arise appear to be resolved without any user intervention. A history of what each user has done to the document is also provided and any unwanted changes can be rolled back.

### 1.1.3 Pidoco

Pidoco is a collaborative wireframing tool. It was not as user friendly for collaboration as Google Drive. Pidoco is much less instant. Sometimes manual refreshes were required to display the work the other users had done. Pidoco also supported multi user editing, however there was no way of seeing which users were editing a document. There was also a history of what changes each user had made. Pidoco has no messaging or commenting system which makes asynchronous collaboration more difficult. Sharing the work requires an email to be sent to the user: they cannot simply be given a link. Again different parts of the workspace have different levels of collaboration. Any UI widgets that are placed are shared, but if one user zooms in on a particular area other users are not forced to that zoom level.

### 1.1.4 Lucid Chart

Lucid Chart is a collaborative diagramming tool. Lucid Chart lies between Google Drive and Pidoco in terms of collaboration speed. It is not as instantaneous as Google Drive, but it does not require periodic manual refreshes like Pidoco. It also allows multi user collaboration and documents can be shared by link or email. Users can be granted read or read and write permissions on the document, like Google Drive, so you can collaborate with people you don't want to be able to fully edit. It has a chat system and users can comment on the document making asynchronous collaboration easier. Lucid Chart does not let you see what a user is doing in real time, but it does provide a full revision history so you can see what changes each user has made. Lucid Chart is different in that it appears to be fully collaborative. Even font preferences are synced across users: if one user clicks bold all users will start typing in bold.

### 1.1.5 SynBioWave

SynBioWave was a biological extension to Google Wave. It is the only example found of a platform for real time collaboration. Unfortunately it could not be made to work. The instruction page still refers to Google Wave. Google Wave has been discontinued since 2012 (some of its real time features found their way into Google Docs). An attempt was made to follow the instructions using Apache Wave (Google Wave was handed over to the Apache foundation) but the SynBioWave tool did not work. The user guide does seem to give a good indication of how the tool would work. The review is based on this. Wave was used to enable the collaboration. You would create a new wave and add your human collaborators. You would also add the SynBioWave agent as a collaborator. When the SynBioWave agent is added to the wave it adds a menu bar. Through this menu the human collaborators can call various functions; the agent then calculates the result and displays the visualisations. The agent and its results can be interacted with in real time. SynBioWave is modular and there are multiple agents available. Custom agents can also be created. The use of Google Wave allowed the biologists to focus on implementing domain functionality without having to worry



about the code infrastructure for real time collaboration. Unfortunately, by relying on a third party for the backbone of the system, SynBioWave appears to have become unusable after Google Wave shut down.

### 1.1.6 Findings

From studying the alternative modelling tools available and their visualisation capabilities it was clear that the visualisation capabilities of Bio-PEPA had to be increased. It was also clear that all the modelling tools were limited in their visualisation capability and that increasing the interactivity and customization available would be very useful.

We can see that there are a number of tools now that offer the ability to collaborate in real time. Only one, however, could be found that was focused on synthetic biology. This one tool appeared to now be obsolete. Given how useful tools like collaborative document editors have been more commonplace it seems like a good idea to try and bring this functionality into the systems biology domain. This would give Bio-PEPA a competitive advantage and hopefully encourage more researchers to use it.

## 1.2 Previous Work

Early on in the first stage of the project it was decided to separate this project from the Eclipse plug-in. It was felt that Eclipse is not the right tool to do data visualisation in.

The initial development stages were focused on getting the new tool from having zero functionality to matching the visualisation features of the Eclipse plug-in. This involved writing an early version of the UI, parsing the Bio-PEPA results data, and plotting it using matplotlib [12].

The next stage of the project was to extend the functionality. The first new feature was intensity plotting where the colour of the line increases in intensity/opaqueness as the population of the species increases. The next feature added was visualisation of the model. Model visualisation used a system of nesting circles and rings to build a hierarchical view of the cell from its model components. Finally the user was given control of the plot, allowing them to alter whether lines are plotted or not, what colour the line is and the thickness of the line. Figure 1.2 shows how the main screen of the program looked at the end of the first stage of development (MPP1).

Over the course of the first stage of work a number of evaluations were carried out with potential and actual users of Bio-PEPA. The findings from these evaluations were used to improve the tool.

## 1.3 Results

should this go after libraries used?

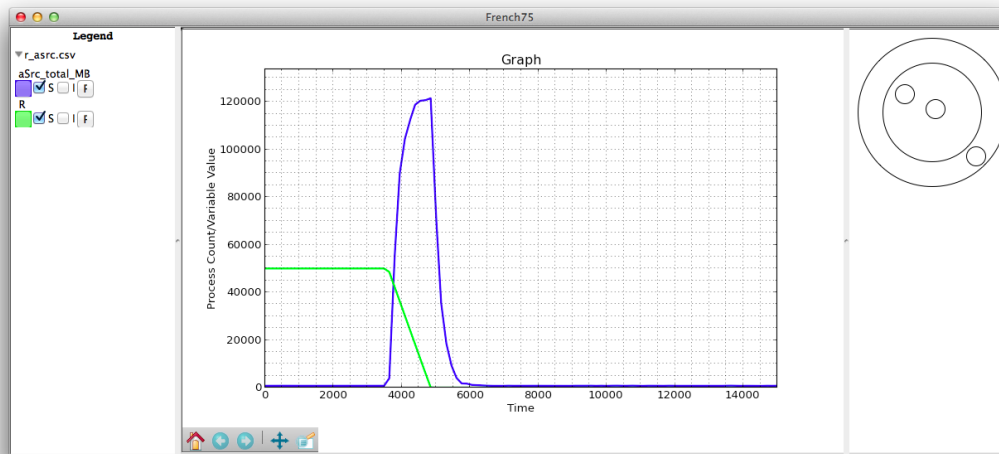


Figure 1.2: Main Screen of the Tool at the End of the First Stage of Development

The tool that has been developed for this project is a fully featured tool with innovative features. It has been positively received by potential users. The tool offers more visualisation and analysis capabilities than the Eclipse plug-in. Users can fully customize the appearance of the visualisation. There is also the ability to add supporting information to the visualisation with annotations and the ability to attach files. Sessions can be saved and loaded allowing users to spread their work over time.

There are also features aimed at making it easier for biologists who aren't comfortable analysing graphs to analyse their work. This has been implemented as animated cell segments simulating the movement of species through the cell.

Innovative features include using plots as queries to find similar plots allowing for discoverability from within the tool, and real time collaboration that allows two collaborators to work together and see each others changes in real time.

There is focus on ease of use in the finished product with all features made as easy as possible to use with little guidance. Relying on recognition not recall.

By implementing more advanced visualisation features this tool should make BioPEPA a more attractive modelling tool. The visualisation features have been brought into line with the features in alternative modelling tools.

### 1.3.1 This Document

The rest of this document details the work that was undertaken during the second phase of this project (MPP2). This is split into separate sections.

Chapter 2 discusses the goals that had been identified and the new goals that have been added. This discussion includes what goals have been refined and why certain goals have been dropped.

Chapter 3 discusses the implementation work that has been performed as part of this project phase. The discussion includes challenges faced and solutions for those challenges.

Chapter 4 discuss the evaluation of the project. This is spread over multiple evaluation sessions and includes evaluations with potential users as well as self evaluations.

## 1.4 Libraries Used

This section details the libraries that were used to build this project. They are all Python libraries. Some libraries have been more important than others. This is detailed below.

### 1.4.1 matplotlib

matplotlib [12] is a graphing library for Python. The graph visualisation is built on top of matplotlib. The graph visualisation acts a wrapper around matplotlib. The user does not have to write scripts that make calls to matplotlib.

Features built on top of matplotlib are:

- Intensity Gradient Plotting
- Annotation infrastructure
- Data normalisation
- Line preferences

### 1.4.2 wxPython

wxPython [13] is the Graphical User Interface (GUI) library. The GUI and the animation visualisation are built on top of wxPython. The animation visualisation uses wxPython's drawing api. This is detailed further in Sections 3.3 & 3.4.2

Features build on top of wxPython are:

- The UI
- Animation visualisation
- Animation annotations

### 1.4.3 SimpleXMLRPCServer

SimpleXMLRPCServer [14] has been used to enable communication between instances of the program. SimpleXMLRPCServer was used to avoid having to write client and

server code that would send and receive messages over sockets. This allowed the development to focus on implementation of the Application Programming Interface (API)

Built on top of this library is

- Real time collaboration

#### 1.4.4 Singleton

talk about the singleton

#### 1.4.5 Miscellaneous

A number of other Python libraries were utilised in the project. They are all part of the standard Python library. They did not provide any significant relevant behaviour, instead they were used to aid implementation of the features in this project. More significant discussion of how any of these libraries were used can be found around the discussion of the features that utilised them. The libraries are: uuid, re, platform, os, time, threading, subprocess, copy, random, numpy, itertools, collections, math, pickle, sys, inspect.

# Chapter 2

## Goals

To help guide development during the project a number of goals were identified. Over the course of the project the goals have been refined, new goals have been added and some goals have been dropped.

### 2.1 Previous Goals

Through research into the problem a number of goals were identified. After performing a review of the existing software, the goals were refined. The initial user evaluations also fed into the refinement of the goals of the project. Some of these goals were completed in the first stage of development, the other goals were carried forward into the second phase of development.

The goals that were completed in the first stage of development are:

- Improve existing capabilities – The purpose of this goal was to add to Bio-PEPA's visualisation capabilities and bring these more into line with the other modelling software. This goal was completed in the first phase of development. The user was given extra control over the appearance of their graphs. Intensity plotting was also implemented.
- Visualise the model more intuitively – The purpose of this goal was to help the biologists to increase their understanding of what the model and the graph is showing them. This goal was completed in the first phase of development with the implementation of the hierarchical drawing of the model components.

The following goals were to be completed during the second phase of development:

- Provide visualisations that take into account the domain – The purpose of this goal was also to help the biologists further understand what the model and the graph are showing them. This has been implemented in the second phase of the project. This goal was merged with animation of the data.

- Animation of the data – The purpose of this goal was to make it easier for the biologists to interpret spatial data by showing the species in their results moving through the cell. This has been implemented: the user can now visualise species moving through a cell.
- Investigating which visualisation combinations work best – The purpose of this goal was to try and work out whether there were any combinations of visualisations that the program could offer that the biologists found particularly useful. This was originally planned to be performed in the second stage of the project, but due to the project focusing on non visualisation features the goal has been removed.
- Add the ability to annotate the visualisations – The purpose of this goal is to allow the biologists to add extra supporting information directly onto the visualisation. This goal has been implemented: the user can add annotations to the visualisations.
- Allow the ability to save and load the program state – The purpose of this goal was to allow the biologists to complete their work in multiple sessions. This goal has been implemented: users can save and load the program state into files, these can then be emailed to other users who can modify them.
- Provide a full session history to the user – The purpose of this goal is to allow the biologists to recover from any mistakes they might make. This has been implemented: the user has a full undo and redo history within the session allowing them to easily correct mistakes.
- Data Mining – The purpose of this goal was to have the program provide some level of automatic annotation and setting optimisation to reduce the workload for the biologists. Due to other features being added that reduce the need for this goal, the goal has now been dropped. The research for this goal was not lost however as a similar goal was added. Data mining to identify features in not a goal that should be abandoned either, instead it would be implemented if the development continued.
- Making meta data accessible – The purpose of this goal was to The plan for this has not changed since the first phase. This goal has not been completed in this phase of development. It was pushed back to allow development on real time collaboration and plots as queries features. It has not been abandoned, instead it would be completed if development was to continue.

## 2.2 New Goals

During the first half of the second phase of development new goals were identified.

### 2.2.1 Data Manipulation and Export

This goal was added after a meeting with the maintainer of BioDARE [15]. This web based tool is aimed at results from laboratory experiments, specifically experiments relating to circadian rhythm.

BioDARE is an online biological data repository. It allows its users to upload their experiment data. This is then available to all other users. BioDARE will generate static graphs of the uploaded data. These plots have little ability to be customized. Instead users can download the data and customize it in a separate application.

The maintainer explained that he had found that researchers often visualise their data after it has been normalised. This removes any issues where different species have different scales and makes it easier for the users to interpret the data.

BioDARE can also export the raw data allowing the researchers to use it in other applications if they desire.

This seemed like a very useful feature for users and it was added as a project goal.

### 2.2.2 Time Series Data as a Query

There has been a lot of research in recent years [16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28] into using time series data as a query against a database of other time series plots through some similarity measure.

This would be a very useful feature to add to the tool. By allowing researchers to use a plot as a query they would be able to discover other plots exhibiting similar behaviour. This could be the same species reacting the same to different species, or a species that is exhibiting similar behaviour. This could help the biologists discover alternative candidate species for research.

This feature would be most effective if there was an online repository of plots for the biologist to search. This would also allow them to discover potential collaborators. Online repositories of data do exist: BioDARE is one of them. These repositories are unlikely, at the moment, to store the data in a suitable way to allow for efficient search.

Given the potential usefulness of this feature, and the fact that none of the other modelling software had this capability it has been added as a goal.

### 2.2.3 Real Time Collaboration

An important area where all the reviewed software were lacking is collaboration, real time or not. The current work flow using the existing software is: perform your analysis, save it, email it to a colleague with additional files and notes, have them open it, they attempt to work out the visualisation is showing. It is a disjointed conversation.

It is not just the modelling software that did not offer collaboration. None of the visualisation software reviewed offered real time collaboration either.

Incorporating real time collaboration was added as a goal to offer researchers a better work flow.



# Chapter 3

## Phase 2 Development

This chapter details the development work that has been performed in the second phrase of the project. This includes conceptual problems faced and their solutions.

### 3.1 Design Decisions

During this phase of development a number of design decisions had to be made. The decisions broadly fell into two categories: UI decisions and architectural decisions. Ideally there should be a method behind such decisions. The decisions and the reasoning behind them are below.

#### 3.1.1 UI Design

During the planning phase of the project (MPP) wireframe designs of the UI components were developed. The purpose of these wireframes was to allow for the UI to be evaluated, at least once, before development started. It is cheaper to fix problems early. Fixing the problem before any development is particularly cheap.

Should I talk about whether wireframing was useful or not?

During the second phase of development (MPP2) the UI was refined and new elements were added to it. It would have been desirable to again first create wireframes and evaluate them with the users before starting development. This approach was not feasible in this stage of development. It would have been necessary to hold more frequent user evaluations, which would have placed a greater strain on the user's time. The alternative would have been to keep the evaluations at the planned frequency and have the development bottlenecked by evaluating the wireframes. It was decided to not use wireframe prototypes. Instead the UI was developed and then users evaluated it. The UI was then changed as required. This iterative approach allowed for the UI to be developed without being bottlenecked by waiting for wireframes to be evaluated and

without requiring a greater commitment of time from the evaluation group. The improvement of the architecture outlined in Section 3.1.2 made changes to the UI easier. After the improvement the UI was contained much less information about the state of the tool. The UI is now effectively just calling API. Having the UI separated means that that a change to the UI no longer requires a refactoring of program logic.

### 3.1.2 Architectural Design

After the first stage of development there was little architectural design. The three main elements of the tool were the UI, the graph visualisation, and the model visualisation. Each component of the UI contained different parts of the visualisation data. Some parts of this data were duplicated across different parts of the program. As you can see in Figure 3.1 each component was passing information to the other components. Some of the items being passed around had references to other components. At the start of this phase of development (MPP2) the features that were to be implemented were too complex for this ‘lack of architecture’ architecture.

The program was refactored to focus on an architectural model where all the data was centralised in a single location. This can be seen in Figure 3.2. If the original architecture was continued, with the new components included, the interaction would look like Figure 3.3. Trying to debug and add existing features would have been non trivial. The visualisation and UI components select what they need to display from this central location. The simplified architecture in Figure 3.2 greatly reduced the amount of information that was having to be passed between all the components. A refresh of the UI components is then all that is needed to display the most update information. This is similar to the Model View Controller (MVC) architecture. A simple diagram for MVC can be seen in Figure 3.4. In this case the controller and the view are not entirely separated. The model in this project is the session state that contains the information to be displayed. The distinction between the view and the controller is currently blurred with some aspect of control being handled by the view. Separating these would be a priority for future work.

A singleton was used to store the session state. A singleton only allows one instance of itself. This prevented the accidental creation of multiple session state objects. Using a singleton helped guaranteed that all session data would be kept together.

Figure 3.5 shows a more detailed architectural diagram of the system.

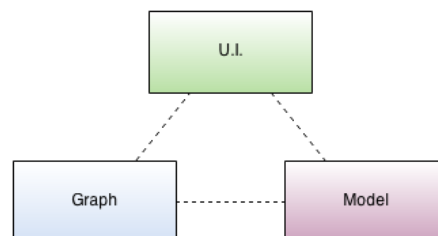


Figure 3.1: Architectural layout at the start of MPP2

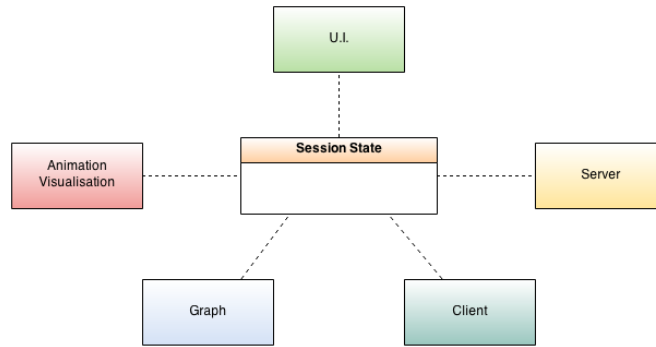


Figure 3.2: Architectural layout at the end of MPP2

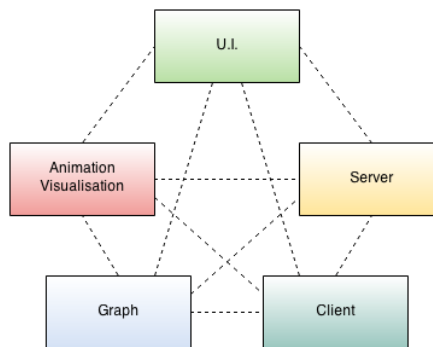


Figure 3.3: Architectural layout at the end of MPP2 if initial architecture was continued.

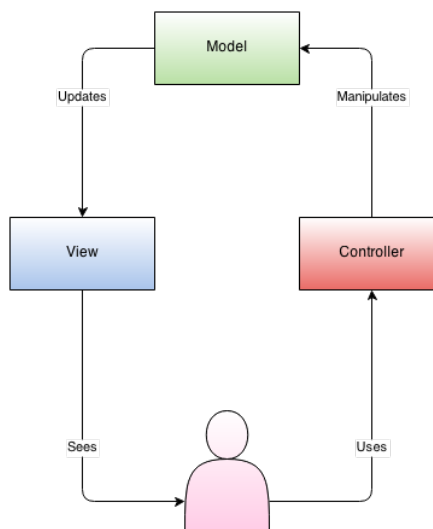


Figure 3.4: Basic MVC architecture

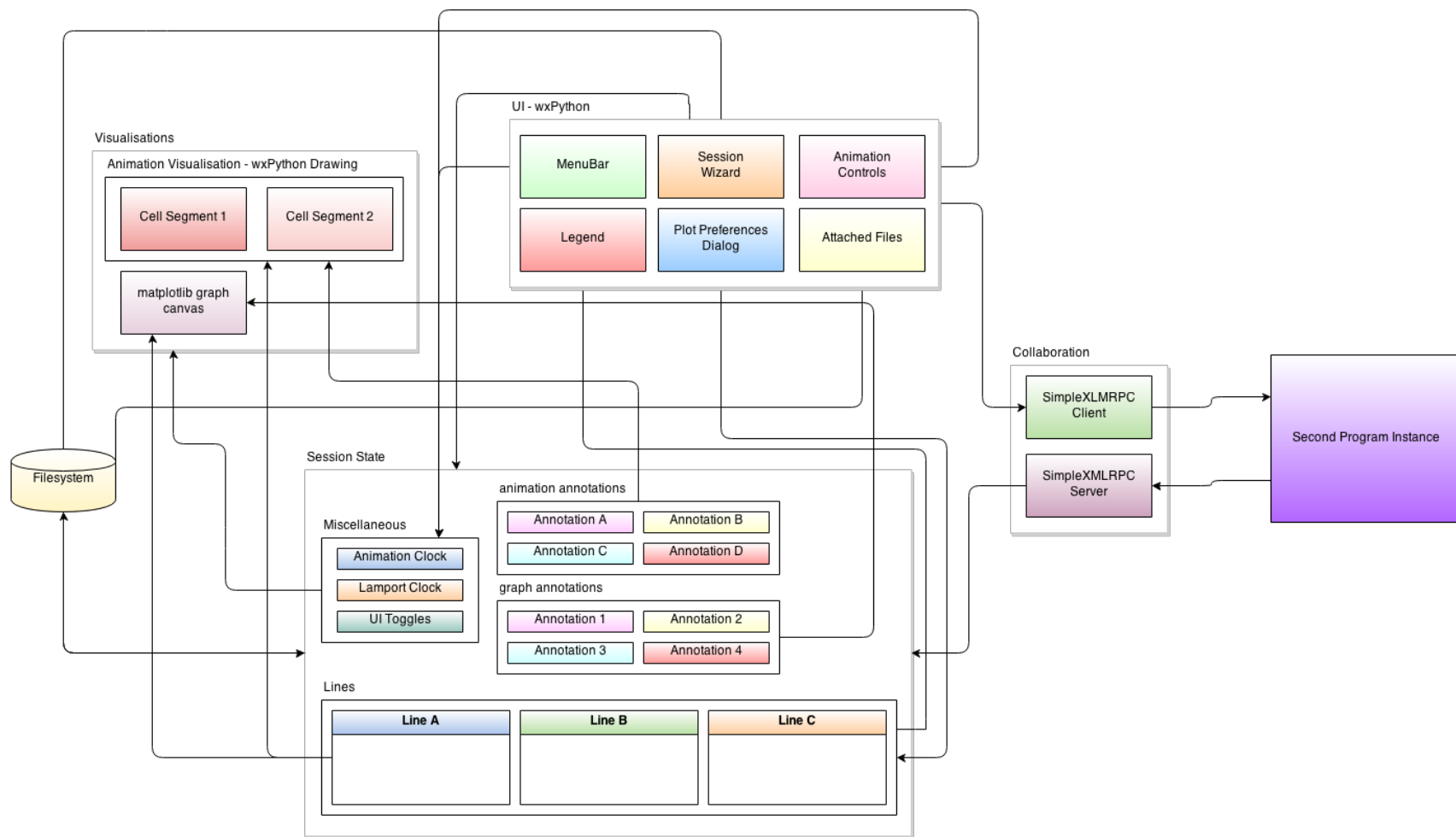


Figure 3.5: Detailed breakdown of program components and how they interact

## 3.2 Finished Product

Not sure how necessary this is. The plan would be a screenshot by screenshot walkthrough - may fit better in the self evaluation. it will possibly just be duplicating screenshots that exist elsewhere

## 3.3 Animation

Animation was a key goal of the project. The core aim of the project was to help biologists who are not comfortable with traditional time-series graphs. The goal was to provide them with visualisations closer to what they see in their domain. This has been accomplished by displaying spatial data in the shape of a cross-section of a cell.

The animation is ideally used to display species moving through a cell. This collapses what would be multiple different lines in a graph, that give no indication of their real position in the cell, into a single image. This can be seen in Figure 1.1.

Animation is just a sequence of static images which, when viewed frame by frame at a sufficient rate, will appear to be a dynamic image. There are two basic ways that animation can be implemented: rendering the entire animation before it is required, or generating a static image as a function of time and every frame calculating what the image should be and displaying that to the user. Both approaches have their positives and negatives. The most appropriate depends on the situation.

We must also decide how the animation will be displayed. wxPython does not have built in animation support but it does, however, have built in drawing support through the paint device context. This allows for arbitrary drawing on UI panels. wxPython also provides some drawing primitives. Various primitives can be handled such as squares, circles, triangles and text. There is also support for drawing arcs. These primitives can also be combined to form regions to allowing more complex shapes to be drawn. During the planning phase (MPP) it was decided to draw cell-cross sections. These cell cross-sections were going to be arcs. wxPython providing support for arcs was perfect for this. This approach would be most suitable for generating images as a function of time.

Using wxPython it would also be possible to render the animation beforehand into a video format and then play it. wxPython has a media control widget. This widget could be used to play the rendered video.

matplotlib does have support for animation with the matplotlib.animation library. This implements animation as a function of time. To display animations matplotlib requires: a function that generates the data to be plotted (which will change depending on the time value) and the interval of time between frames. matplotlib also has support for shapes. Each cell cross-section could then be treated as a separate matplotlib animation.

Rendering the animation before it is needed is not the correct way to do animation in this circumstance. One of the purposes of this tool is to allow for interactivity and customisation. This would potentially lead to the animation having to be rendered multiple times. The animation that was planned to be included in this tool is also not complex enough to warrant pre-rendering. Pre-rendering would be more suitable for more realistic animations. One would be doing this after the insight has been learned from the experimental data. The purpose of the animation here is to help the user to find this insight. For these reasons treating animation as generating static images as a function of time was chosen.

wxPython and matplotlib's animation capabilities are quite similar, both have primitives for drawing the necessary shapes and both are implemented as drawing static images as a function of time. Given that the capabilities of each were similar wxPython was chosen for animation to avoid the scaffolding code that matplotlib would have required. The scaffolding code is the code required to bind graphs to UI components and set up the correct axis size.

The animation visualisation will usually be one or more cell cross-sections. Each cross-section is split into compartments. These compartments are parsed from the model file. In the model file compartments are stored hierarchically. The structure of the cell in the model can be inferred from this hierarchy. Each compartment in the model relates to one segment in the cell cross-section. wxPython's in built arc drawing was used to draw the cross-sections. The cross-section has to be drawn from the edge of the structure into the centre. Each arc is drawn on top of what is already there. If the larger outer arcs were drawn last then they would cover the inner arcs. The division of the cell cross-section is such that every visible segment has the same width.

Over the course of the animation the colour of the segment changes to reflect the colour in the intensity plot version of the line. This allows the researcher to compare the two visualisations and will hopefully help build their confidence understanding the graph plot.

Animation takes advantage of the data centric model. When the play button is pressed a thread is created whose job it is to increment the animation clock and refresh all the UI elements. When refreshed the UI elements pick up on the change of clock and update what they display appropriately.

The first step towards animation was calculating at what points the line changes colour. This is done during the interpolation phase that was implemented in the first stage of development. To do the interpolation the original plot is split into multiple separate plots with the results padded with null values to allow matplotlib to plot them as one. This is detailed further in . The number of these null values is counted and that gives the time at which the colour should change. This is stored as an array of time, colour tuples. Each line also has an internal counter to say how many elements into this array the line is. When the animation clock is updated and the refresh triggered, each cell segment finds the appropriate colour change point in the line by going through the array and comparing each time to the animation clock until the time is found and updates the segment colour. This position is then used as the new starting point on the next refresh. This removes the need to look through past entries unnecessarily.

The user can control the animation clock through the time slider. Moving the time slider updates the animation clock to be the value of the time slider and triggers a refresh of all the cell segments. This ensures that the user is seeing up to date information. As described above, each line has a list of times at which it changes colour. These are sorted according to time. During ‘normal’ animation the previous position in the array is tracked so that the whole array is not searched every time unnecessarily. When the time slider is used this previous position might become a future position. To solve this the previous position is set back to the start of the array. On the next refresh the array will be searched from the start for the appropriate time and colour.

A nice UX feature that has been added is a line on the main graph that indicates the current time in the animation. This can be seen in Figure 3.6. When the animation is running the line moves along the graph. This again helps the user build a correspondence between the two visualisation types and may help with their understanding of what is being displayed on the graph.

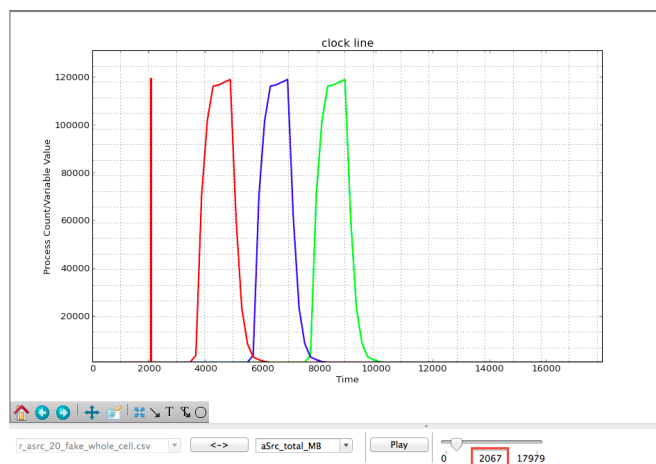


Figure 3.6: Vertical line that is drawn on the graph to indicate current animation clock time.

The user is also given control over what is displayed. The cell segments can be drawn in a file focussed manner or a species focussed manner. In the file focussed manner a cell cross section is drawn for every species in that file. The user can switch between all results files that have been added to the session. In the species focussed manner a cell cross section is drawn for every file that contains that species. The user can select between all species found across all the results files. These are illustrated in Figure ??

Figure 3.7 shows some example cell segments. Figures ?? & 3.7b show the cell segments being drawn in the file focussed manner. Both files contained results relating to the the species “aSrc\_total\_MB” In Figure 3.7a this species was present at all locations in the cell. In Figure 3.7b the species is missing from the middle location in the cell, so that segment has been left white. Figure 3.7c shows the cell segments drawn in a species focussed manner. One cell cross section has been drawn for each file which contains the species “aSrc\_total\_MB”. We can see that in one of the files the species wasn’t present at all locations.

It had been hoped to allow the user to save the animation. This could have either been



Figure 3.7: Example cell segment animations

allowing the user to save individual frames or allowing them to save an animated gif of the whole animation. Saving to a gif was chosen as it was felt to be more useful to a user to be able to export the entire animation rather than a single frame of it. Allowing the user to save a single frame would be added in at a future date. To generate the gif we need to save all frames. As these frames are generated as needed, not prerendered it was required to run through the whole animation to save each frame. This meant that saving all the frames took as long as the animation took to run. At this step hundreds of static images have all been saved. Then a shell script is required to convert the saved frames into a gif. Given that this was a very slow and user unfriendly approach the decision was made to remove the ability to save animation from the project. It would be implemented if future work was to be performed.



## 3.4 Annotation

Annotation was an important feature to add. It is one of Grinstein's features that data visualisation software should have [29]. This is because the purpose of a visualisation is to aid analysis. If you have a print out of a visualisation you are able draw on it and highlight areas of interest. Users need to be able to do this on digital versions of their visualisations. As well as helping users analyse their data, being able to annotate also means that images for presentations can be prepared without having to save the visualisation and open it in an external program. If you did this and then wanted to change the visualisation you would have to re-annotate it. This is frustrating for a user and wastes their time. Being able to do it from within the visualisation solves this problem as the raw data and the annotation data are together. Another benefit of being able to annotate is having another way of attaching additional information to the visualisation when sending it to a colleague. Having this information on the visualisation saves them from flicking back and forth from an email or other supporting documentation.

Implementing annotation of the visualisations on offer in this tool had to be done in two parts. This is because there are two parts to the annotation: the graph and the cell level view. One is a static visualisation and one is a dynamic visualisation. Each of these required a different approach. These are detailed below.

### 3.4.1 Annotation of the Graph

Annotations of a static image are extra elements that are added by the user. We need to decide what extra elements we are going to allow the user to add. We also need to decide how to render the annotations.

For this project there are two options for drawing the annotations on the graph: with matplotlib and with wxPython. matplotlib has support for annotations. In matplotlib annotations have two forms: one form uses the `annotate()` and `text()` functions which adds an arrow with optional text to the graph and adds text without an arrow to the graph respectively. The second way would be drawing with wxPython. As in Section 3.3 we can use device contexts to draw primitives on UI elements with wx. The primitives offered by the device context do not include arrows which are a key annotation type. In Section 3.3 wx was chosen over matplotlib partially because using matplotlib would have involved more scaffolding code. For annotating on the graph this scaffolding code is already there from the creation of the graph. For these reasons it was chosen to implement annotation on the graph using matplotlib's support.

matplotlib's support for elements on the graph is quite thorough. Text and arrows are obvious choices as they allow the user to indicate and explain areas of interest. matplotlib then allows for more arbitrary drawing. So as to not overload the user with annotation choices it was decided to limit the number of annotation elements that they could use. In addition to the text and arrows the user can also add circles as an annotation.

Users of the new tool are provided with four annotation types: arrow, text, arrow with text and circles. Buttons for each of these annotation types have been placed on the matplotlib toolbar. This can be seen in Figure 3.8. These four annotation types come from different libraries within matplotlib. An annotation class was defined to attempt to hide their differences. This would also make it easier were more annotation elements added in the future.

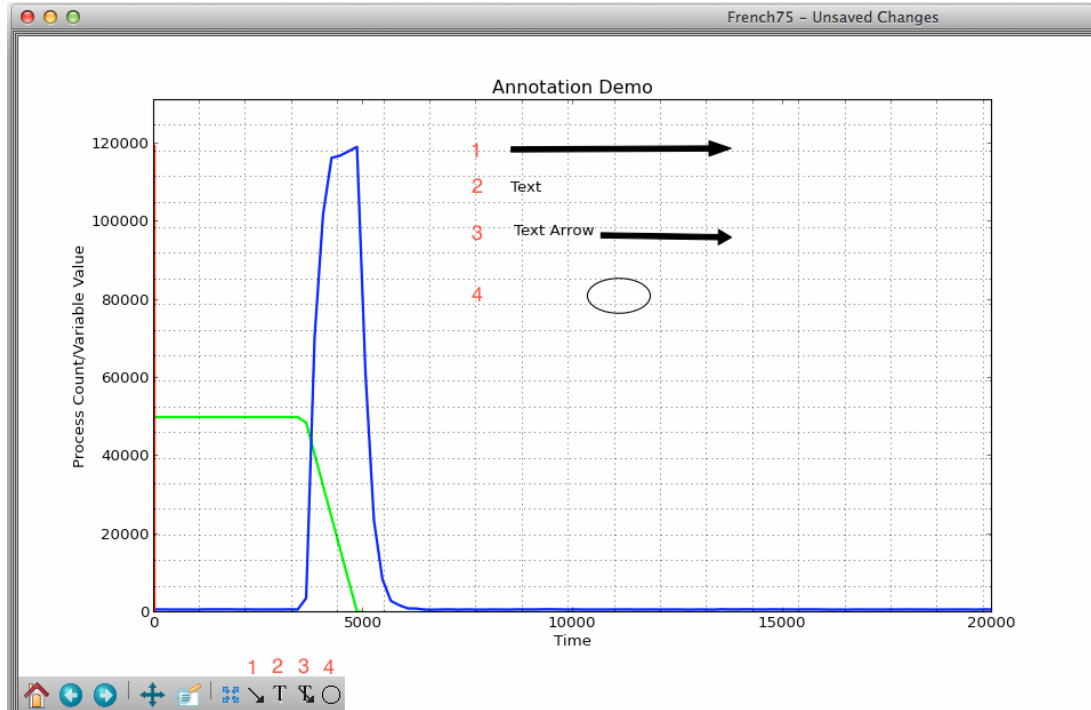


Figure 3.8: Example of each of the four graph annotation types and the toolbar buttons that correspond to them.

There were problems in making the annotation creation process user friendly. For all of the annotations the user needs to first press the appropriate button on the toolbar. The next actions depend on the annotation type. Text and circle annotations were intuitive as all they require is one click. The user clicks where they want the annotation to be placed and it will be drawn on the graph there. However the two arrow annotation types required two clicks. The first click marks the start point (the tail of the arrow) and the second click is the finish point (the head of the arrow). This was not obvious, user evaluation revealed that the users did not know that it required two clicks and didn't know whether the arrows would be drawn head to tail or tail to head. The technique for placing arrows was subsequently changed so that the first click still fixed the position of the tail of the arrow. However the behaviour after the first click changed, so now a temporary annotation is continuously redrawn that has the head of the arrow wherever the cursor is. This allows the user to see the arrow they are drawing. To indicate to the user that they need to click on the graph the cursor is changed after pressing one of the annotation buttons on the toolbar. The use of different cursors is a common technique to help guide users to perform actions.

It was important that annotations could be edited or deleted. The annotations can not

just be clicked as they are not a UI widget like a button. The solution to this was to have an array of annotations. When a user right clicks on the graph it searches through all annotations and selects the annotation that was closest to the click (if that distance was below a certain threshold). The selected annotation is then highlighted red, and a context menu appears to give feedback to the user that they have successfully selected an annotation. This can be seen in Figure 3.9. The context menu then gives the user the option to edit or delete an annotation. Editing an annotation only allows for editing text. For changing position, the annotation has to be deleted and redrawn.

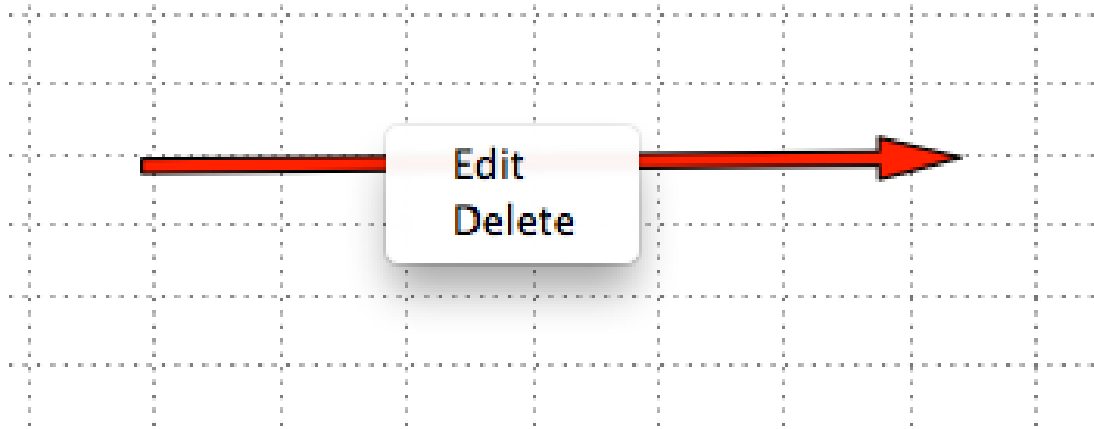
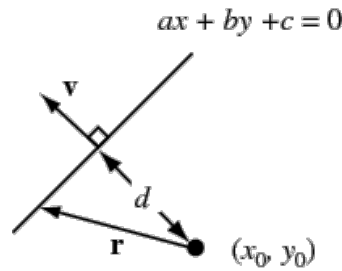


Figure 3.9: Context menu on selection of graph annotation.

To calculate the distance from the location of the mouse click to the annotation two problems had to be solved. The first was how to calculate the distance from a point to a line. An equation for this can be seen in Figure 3.11. Equation 3.1a is the equation for a line in two dimensions. This represents the annotation. The equation can be calculated from the start and end points of the annotation. Equation 3.1b is the position of the mouse click. Equation 3.1c shows the equation for calculating the distance from the point to the line. The second problem that had to be overcome was the difference in scale. In effect we have two coordinate systems. The results coordinate system and the visual coordinate system. These difference are illustrated in Figure 3.13. The annotations in Figure 3.13a are further away from each other in the results coordinate system than the annotations in Figure 3.13b. However they are closer in the visual coordinate system. When selecting an annotation the user is using the visual coordinate system but the distance is calculated using the result coordinate space. This could lead to the wrong annotation being selected. The solution to this was to transform one coordinate system into the other. When calculating the distance from the point to the line, the values are scaled by the size of the graph.

A similar issue to the difference in scale when calculating the distance from the mouse to annotation was encountered when switching the graph to normalised mode. When normalised the graph in the y axis only goes between 0 and 1. The coordinates of the annotation are likely to be very much outside this range and the annotation would not appear on the graph. The solution, if the graph was normalised, was to normalise the positions of the annotations when drawing them. A side effect of this technique is that the lines are potentially going to rescale themselves. This could leave an annotation point to nothing. This is unavoidable as the annotation has no concept of what it is



(a) Diagram of the point to line distance problem

$$y = -\frac{a}{b}x - \frac{c}{b} \quad (3.1a)$$

$$(x_0, y_0) \quad (3.1b)$$

$$d = \frac{|ax_0 + by_0 + c|}{\sqrt{a^2 + b^2}} \quad (3.1c)$$

(b) Equations for calculating the distance from a point to a line.

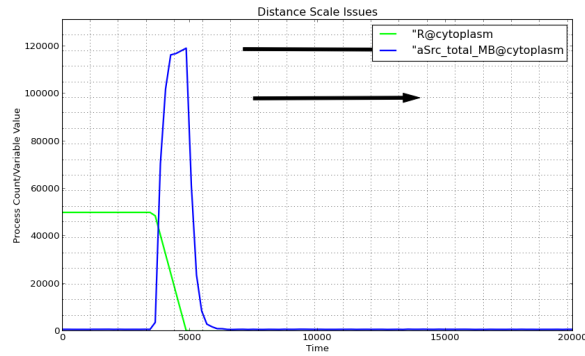
Figure 3.11: Equations and diagram for calculating the distance from a point to a line (taken from WolframMathWorld [30])

pointing at. It may be pointing to the intersection of many lines; in this case it would be impossible to keep the annotation pointing at what it was originally pointing at after rescaling the axis. During user evaluations, when the annotation was not visible during data normalisation, the user was very confused as to why their annotation had disappeared. It was therefore decided to have the annotation be visible but most likely incorrect as it is clearer to the user what has happened than just not displaying the annotations.

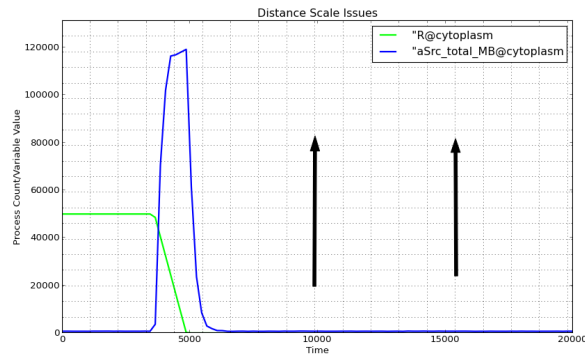
### 3.4.2 Annotation of the Animation

After completing annotation of the graph it was important to expand this to the animation panel, as this is the other visualisation option available to a user. Annotating animations posed more of a challenge than annotation of the graph and there were a number of issues to overcome.

1. How to implement the annotations? For the graph matplotlib has built in annotation support. wxPython does have drawing support but not built in annotation support. Annotating on the animation will need manual handling of the drawing on top of the animation visualisation. Manual drawing means that the automatic layout functionality that wxPython provides cannot be used.
2. When to display the annotations? When an annotation is drawn on the graph it is displayed at all times. The appearance of the graph does not change over time. However the appearance of the animation visualisation does change over time. The problem faced when annotating is whether to have annotations available at



(a) Annotations with distance of approximately 20000



(b) Annotations with distance of approximately 5000

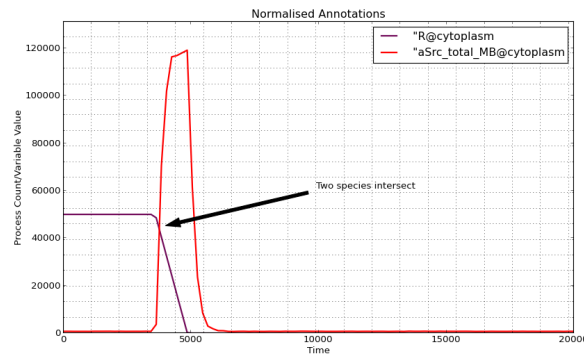
Figure 3.12: Two sets of annotations illustrating the problems when calculating the distance to an annotation

only specific times in the animation, or to have them there the whole time; and if they are going to appear and disappear, how can it be done without being distracting?

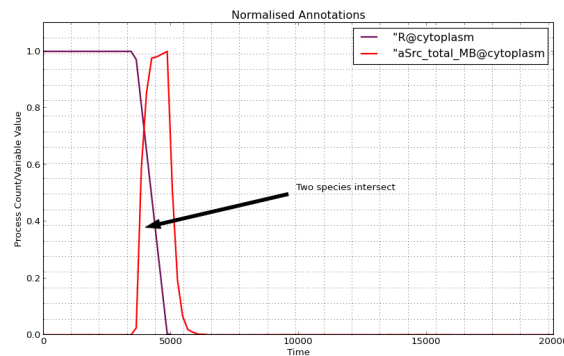
3. How to give the user control over the annotations? When a user wants to edit or delete an annotation on the graph it is always there. However on the animation panel if the annotation is temporal, then it is not always visible for the user to edit or delete and it would be frustrating for a user to constantly have to search through the animation to look for annotations to change them.

A platform where users are able to annotate an ‘animation’ is YouTube [31]. YouTube’s annotation interface can be seen in Figure 3.14. Youtube’s approach is to allow the user to set what period of time an annotation is visible for. There is also a management panel where the user can see all of the annotations and edit or delete them. This approach solved issues two and three and was adapted for this tool.

On the right hand side of the tool there is a panel which lists all annotations that have currently been added to the tool, this can be seen in Figure 3.15. This provides the user with the persistent view of what has been added. When the user creates an annotation they are shown a dialogue that allows them to enter the annotation text and to choose a duration. This indicates to the user that the annotation will only be visible at certain



(a) Annotation pointing at the correct intersection of species



(b) Annotation pointing at nothing after data normalisation

Figure 3.13: Graphs illustrating what happens to annotations whilst data is normalised

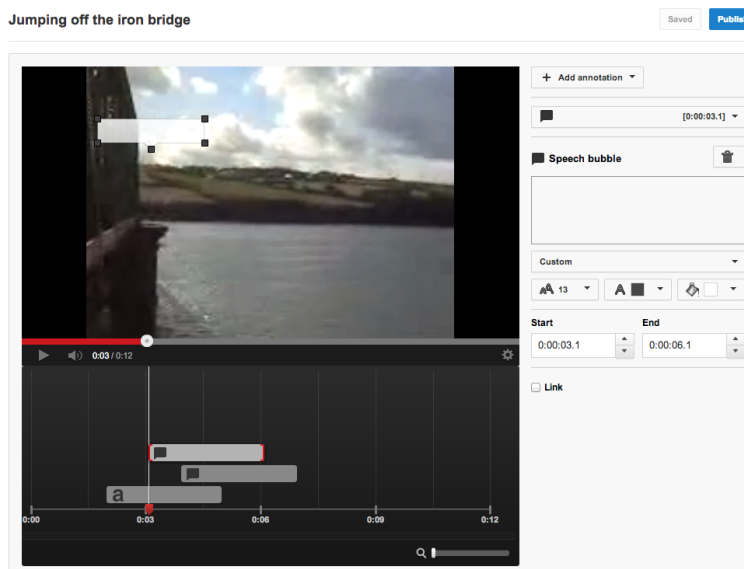


Figure 3.14: YouTube video annotation interface

times. The start and end times are given in model time. The duration displayed to the user is given as the real time that the annotation will be visible for. This can be seen in

Figure 3.16.

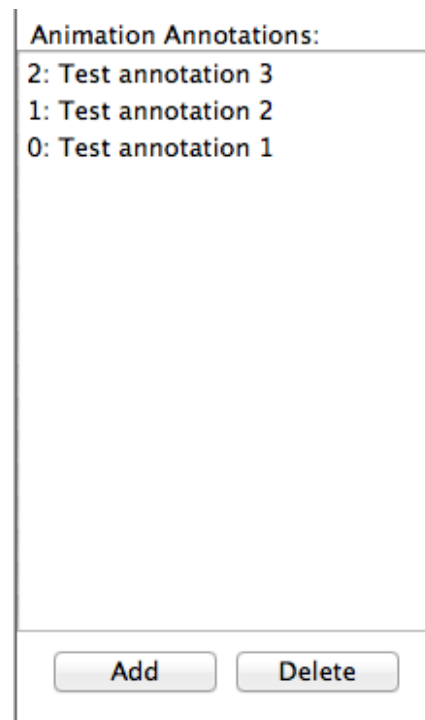


Figure 3.15: Animation annotation list

The first problem – how to display the annotations – has also been solved. Drawing in wxPython takes place on panels. Each cell cross section is placed on its own panel. This drastically limits the available space. The panels are not wide enough to have the annotation text drawn on. The solution was to assign each annotation a number and that number is what is used to annotate the cell. The number is then displayed in the annotation list box allowing the user to read the appropriate annotation. This can be seen in Figure 3.17

## 3.5 Search

Begin able to use a time series graph as a query against a database of time series data was one of the new goals added to the project. It was felt it would be of great use to the biologists as it would allow for discoverability within the tool. It had been identified as a potential goal whilst researching data mining in time series data. It was added as a definite project goal after the user group backed it as a feature they would very much like to have. It is also a feature in the tool that incorporates cross domain knowledge in an area of active research.

Some problems needed to be overcome before using plots as a query was useful:

- How to cope with different scales?
- How to cope with events happening at different times?

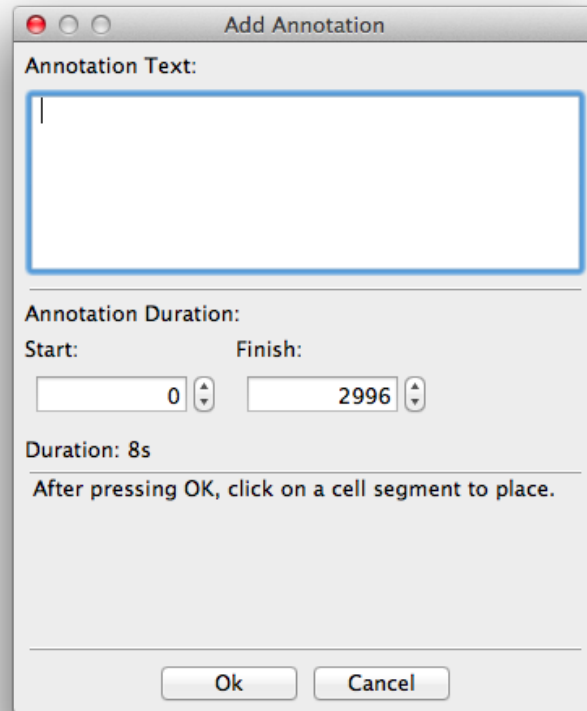


Figure 3.16: Animation annotation dialogue

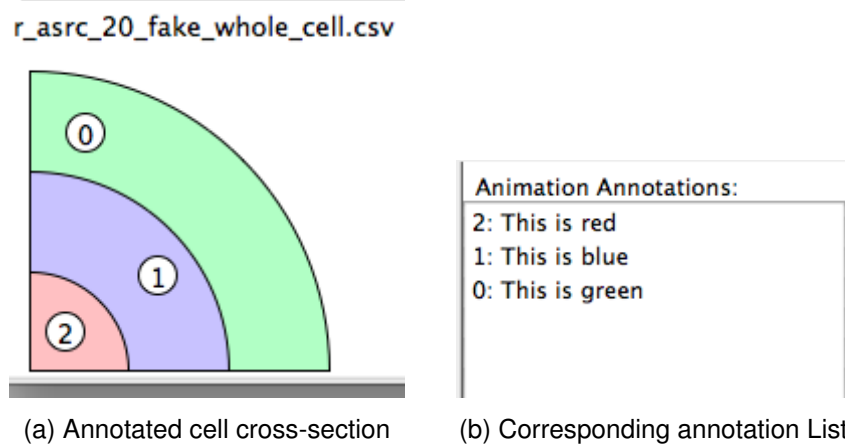


Figure 3.17: Animation annotations

- How to represent the plot to allow for efficient search?
- How to determine similarity between two graphs?

Early techniques used simple similarity measures such as Euclidean distance, but these gave poor results [17]. The reason for the poor results is that Euclidean distance does not take into account scale or time. Figure 3.18 illustrates a case where Euclidean



distance would give a poor similarity measure. The two lines are identical, but the green line has been shifted in time. Euclidean distance will not recognise that they are similar. The same problem would happen if one line had been shifted in the y-axis above the other.

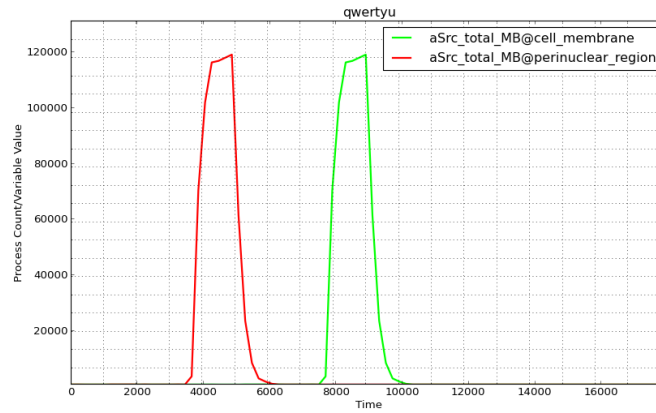


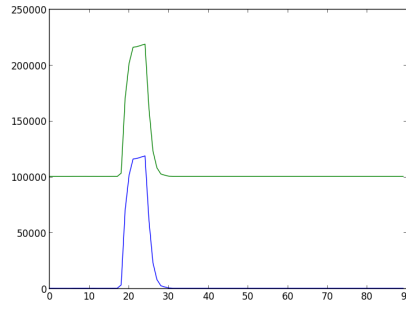
Figure 3.18: Identical plots shifted in time

It is important to define what it means for two lines to be similar. It is obvious that in a case such as Figure 3.18 that the two lines are similar. Both lines are the same shape, they are exhibiting the same behaviour. When determining similarity of plots we should take a time invariant approach. Perhaps less obvious is the case where we have offset in the Y-Axis as shown in Figure 3.19. Figure 3.19a shows two identical lines where one line had an initial 100000 population level. Figure 3.19b shows two lines where one line is double the other line. Figure 3.19a is effectively the same case as Figure 3.18. Figure 3.19b is different, but it is clear looking at the graph that they are exhibiting the same behaviour. When determining the similarity of plots we should take a scale invariant approach [16].

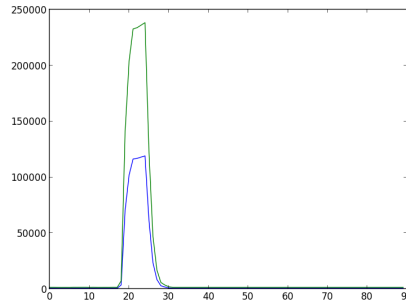
Approaches that have been researched to determine similarity between time series data have included shape based approaches and dynamic programming techniques. Not all of these approaches have included dimensionality reduction. The dimensionality reduction transforms the input data into a vector space with fewer dimensions. Similar values in the data space will, in theory, be transformed to the same value in the vector space. The necessity for dimensionality reduction is explained later in this section. After researching alternatives an approach was adapted from the work of Lin & Li [21].

Their approach uses normalised sub sequence data to reduce the dimensionality of the dataset. The reduced data can then be used to efficiently find the most similar plots in the database. The input data is converted into sub lists to ensure that all features will have been captured. These lists are then normalised and have their dimensionality reduced. This is where Lin & Li's work left the topic. Work from the field of information retrieval was then applied to give the similarity ranking. This approach is explained in further detail below.

The first step is to take the input data and return all the sub lists of this data. To do this a sliding window was used. The size of the sliding window will determine the



(a) Identical plots shifted in population level



(b) Identical plots scaled in population level

Figure 3.19: Graphs illustrating different ways lines can be offset in the y-axis

feature size in the reduced vector. Lin & Li use a size of 8 for the sliding window. The same value was chosen in this implementation. If we have input data  $[1, 2, 3, 4, 5]$  and windows size  $n = 3$  (for illustrative purposes only) then we get the sub lists as  $[[1, 2, 3], [2, 3, 4], [3, 4, 5]]$ .

The second step is to normalise each sub list. Each sub list is normalised to be zero mean and unit variance. This places each sub list onto the same scale. This removes the effect of the offsets seen in Figure 3.19. To normalise to zero mean and unit we need the mean ( $\mu$ ) and the standard deviation ( $\sigma$ ). We then have:

$$\left[ \frac{X - \mu}{\sigma}, \forall X \in sublist \right]$$

The third step is to reduce the dimensionality of the data and build the fingerprint. Lin & Li's approach also involves converting the data from a continuous representation into a discrete representation. By normalizing the sub lists to be zero mean and unit variance we allow a Gaussian distribution to be fitted to the data. This can be seen in Figure 3.20. This approach does assume the Gaussian distribution is a suitable distribution. Lin & Li found it to be a suitable distribution but does note that other distributions could be investigated: this was outside the scope of this project. For each data point we use the Gaussian function and map it to a letter. This can be seen in Figure 3.20 which shows a line formed of eight datapoints, our sublist, that have been normalised to be zero mean and unit variance. Each datapoint has been mapped to one

of the three sections of the distribution divided by the breakpoint and has been replaced with the letter that represents that section. Breakpoints are used to divide the Gaussian distribution into sections with equal probability. To split the distribution into three sections two breakpoints are needed. The values of these breakpoints can be found in statistical lookup tables. If our breakpoints divide the distribution into three sections then each datapoint can be mapped to one of these three sections. Lin & Li divide their distribution into three sections and represent each section with a character. They found three sections to be suitable, again alternative numbers of breakpoints could be used but this is outside the scope of the project. This reduces the dimensionality of the dataset as multiple real values are mapped to a single letter.

After this stage each sublist has been transformed into an eight character long string. The plot is represented as a list of strings. These strings represent the features of the plot.

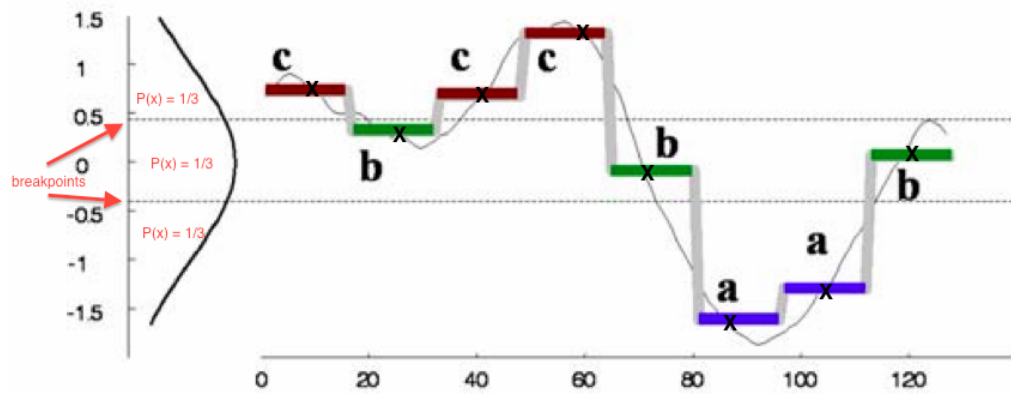


Figure 3.20: Fitting a Gaussian distribution to time series data, taken from [21].

Lin & Li now call for the removal of local duplicates of within the list of words. If we have runs of a certain word then that should be reduced to a single instance of the word. For example if we have the fingerprint “AAA”, “AAA”, “BBA”, “BBB”, “BBB”, “AAA”, “AAA” it should be reduced to “AAA”, “BBA”, “BBB”, “AAA” not “AAA”, “BBA”, “BBB”. We are not removing all duplicates of the word. This reduces the number of entries in the fingerprint and provides an efficiency benefit. It will also remove the effect seen in Figure 3.19b where Euclidean distance would fail if one line is a scaled version of the other.

The plot is now represented as a list of strings without local duplicates. Our strings all have a length of eight characters, the strings are composed from a three letter alphabet. This gives us a vocabulary of  $3^8$  possible words. The index of the plot is a  $3^8$  element vector where each entry in the vector contains the count of the word in the line. This is a very wasteful representation as most of the entries in the vector will be zero. We can replace this with a sparse representation of the where we store what words make up the line and their count.

This representation of the plot also solves the problem of features being temporally

located in the graph. In the plot index there is no ordering of the features, in effect all the features happened at the same time. This is a common representation in information retrieval and is referred to as bag-of-words. A more suitable name in this domain would be bag-of-features.

Now that we have a representation of the plot that is scale and time invariant and allows for efficient comparisons we can find a method to calculate the similarity. Lin and Li leave similarity as future work and use their representation to perform clustering, classification and anomaly detection. It was therefore necessary to find a technique, potentially from a different domain, to find the similarity.

We could simply just use Euclidean distance again, using the two vectors as the input, but this does not do anything to weight rarer features. If two plots have rare features in common then that is much more significant for similarity than if they share very common features. A similarity measure that takes into account the importance of a word is therefore desirable.

It was decided to implement tf.idf weighted cosine as the similarity measure. This takes into account term frequencies to provide a higher weight to rarer vectors. tf.idf is very well researched in the field of information retrieval and text mining. Given the representation is equivalent to a bag-of-words it seems sensible to apply tf.idf to this domain. A description of how tf.idf weighted cosine works can be found in Section 3.5.1.

The tf.idf weighted cosine similarity [32, p. 243] can then be calculated between the query plot and each individual plot in the database. These similarity scores can then be ranked.

### 3.5.1 tf.idf

term frequency-inverse document frequency (tf.idf) is a technique for calculating how important a word is to a document. tf.idf requires a corpus of document vectors. With the corpus data tf.idf can take as input a query vector and transform that into a vector of weights, where the weights are the tf.idf scores of each word.

Why do we care about weighting words

Equation 3.2a is the equation for finding the similarity between a short query term and a longer document. This equation would be applied between the query term and every document in the corpus. This would then give a similarity ranking. Below is an explanation of the components of the equation and how they affect the weighting of a word.

- $tf_{w,Q}$  – The number of times word  $w$  appears in query  $Q$ . If a word is repeated in the query it is likely to be important.
- $tf_{w,D}$  – The number of times word  $w$  appears in document  $D$ . If a word is repeated in the document it is likely to be important.

- $k$  – A squashing factor. The first occurrence of a word is more important than repeats.
- $|D|$  – Length of the document
- $|C|$  – Number of documents in the corpus
- $df_w$  – Number of documents in the corpus that contain word  $w$
- $avg|D|$  – Average length of all documents in the corpus.
- $tf_{w,D} + \frac{k|D|}{avg|D|}$  – Normalising factor, repeated words are less important unless the document is longer than average.
- $\log \frac{|C|}{df_w}$  – Increases the weighting of words that are rare across the corpus.

Equation 3.2b is the equation for calculating the **tfidf!** (**tfidf!**) weighting of a single word in document vector. This equation would be applied to every word in the document vector and transform the document vector into the document's weight vector.

Equation 3.2c is the equation for **tfidf!** weighted cosine similarity. This gives a better measure of similarity when comparing the query is a document as is the case here

Citation for this is tts lecture slides is this ok?

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$$s(Q, D) = \sum_w tf_{w,Q} \times \frac{tf_{w,D}}{tf_{w,D} + \frac{k|D|}{avg|D|}} \times \log \frac{|C|}{df_w} \quad (3.2a)$$

$$D_w = \frac{tf_{w,D}}{tf_{w,D} + \frac{k|D|}{avg|D|}} \times \log \frac{|C|}{df_w} \quad (3.2b)$$

$$S(D_1, D_2) = \frac{\sum_w D_{1,w} \times D_{2,w}}{\sqrt{\sum_w D_{1,w}^2 \times \sum_w D_{2,w}^2}} \quad (3.2c)$$

tf.idf allows for efficient search through the use of an index and an inverted index. The index is a mapping from document to words and the inverted index is a mapping from words to documents. If we look up a document we find what words are in it. If we look up a word we find what documents contain it. When a new document is added to the database these two indexes are updated. With these two indexes all the components of **tfidf!** can be calculated for very little cost. For example  $tf_{w,D}$  would involve looking up document  $D$  in the index, this will return to us all the words in document  $D$  and their counts. We can then quickly find the count of word  $w$  in  $D$ .

## 3.6 Real Time Collaboration

More and more programs are allowing their users to collaborate in real time. Some of these pieces of software are discussed in Section 1.1. Only one piece of software could be found that was biology focused. Implementing real time collaboration in this tool is therefore a great advantage.

There are two broad ways that real time collaboration could have been implemented: in a distributed manner or by having a single centralised instance. A single centralised instance is the approach taken by all the software reviewed. The approach taken in this tool is the distributed approach.

The single centralised approach would have been simpler. With the distributed approach you have to ensure that each instance of the program has the same ordering of events and is displaying the same state. In the centralised approach there is a single repository which has all history of what has happened. The order that the single program instance thinks events happens in is the order that those events happened in.

Ideally the collaborative model used would have been the centralised model. This would however have required a significant reworking of how the tool worked. All the software reviewed was browser based. Given the time constraints on the project and the fact that it had been developed up until this point as a standard desktop application it was felt it would be easier to implement distributed collaboration.

The collaboration that has been implemented in this project is only between two program instances.

The first stage in allowing distributed collaboration was enabling the different program instances to talk to each other. There are a number of techniques to allow for this. Data could have been sent directly through sockets, another option would be to create a web server and send get requests and pass parameters. It was decided to use an existing library to handle the communication. The library chose was `simplexmlrpc`. This allows each instance to run a client and a server. The server has an api and this can be called.

On initial connection the ‘master’ sends its entire session state the the ‘slave’ uses this as its initial state. After this whenever either client performs an action it issues a call to the other clients server to perform the same action.

This approach was not without its problems. A fundamental problem with having the collaboration follow a distributed model is how to ensure that both program instances have the same ordering of events. To solve this problem Lamport clocks were used. Any action that affects the session state, an action that does this is one that will need to be sent to the collaborators, increments the Lamport clock. The Lamport clocks are then sent with every message. When a message arrives the program checks the Lamport clock and works out where in the history it should go. After reordering the history all the visualisations are redrawn. This also has to be done in the instance of the program that is sending the change.

Another issue that was encountered was to do with how the program was architected. The session data was often directly modified by methods that were bound to UI ele-

ments. These methods are also the ones that called the client to send the message to the collaborator, if the server on the other end then called the same method to perform the action it would have resulted in an infinite loop. These methods would also often do more than just modify the state, for example when modifying an annotation the method will find the appropriate annotation and edit it. The client would then send a message saying delete annotation 1. The server would not be able to call the original method as it doesn't take a parameter. The solution to this was to split the methods up more. Now the more typical flow is the UI bound method calls another method that modifies the state. The server is able to call the same method. This removes the need for duplicated code.

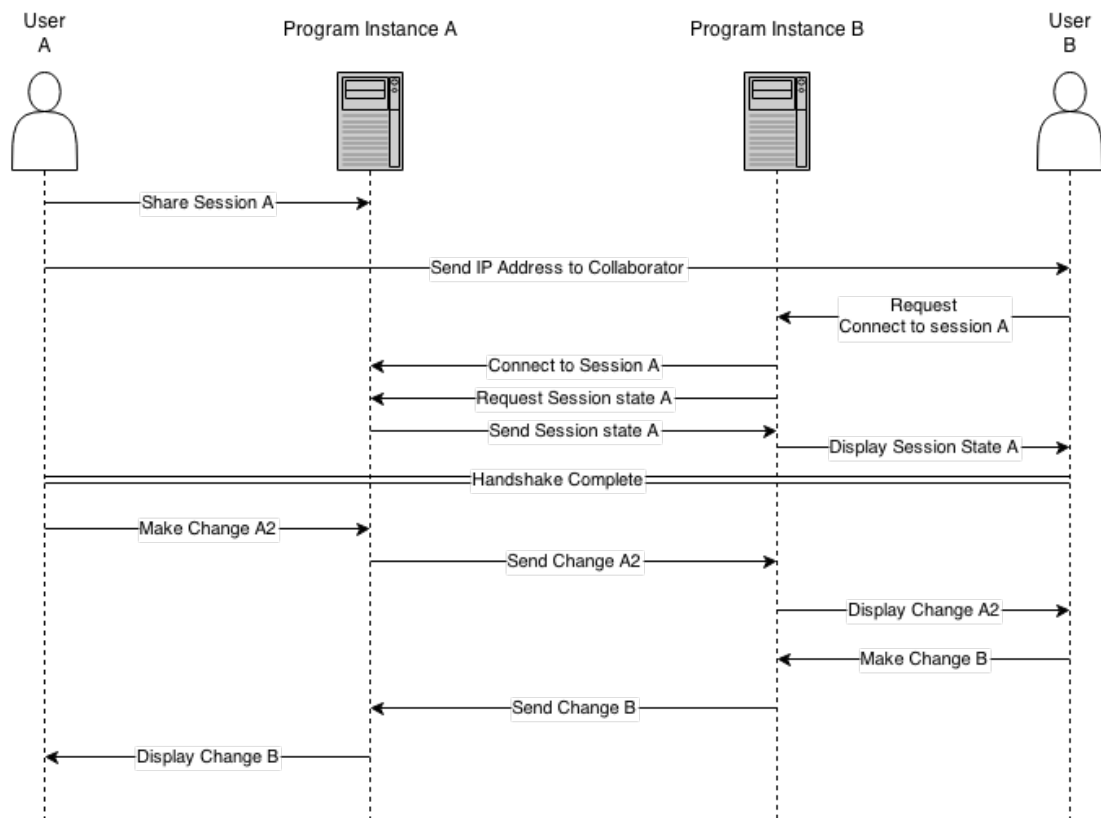


Figure 3.21: Collaboration handshake stage

## 3.7 Usability

In all the evaluations of the project users have commented on the difficulty of using parts of the tool. Action has been taken to make it easier to use. Many of the changes have been guided by Shneiderman, Norman & Nielson's guidelines. Specifics are detailed below.

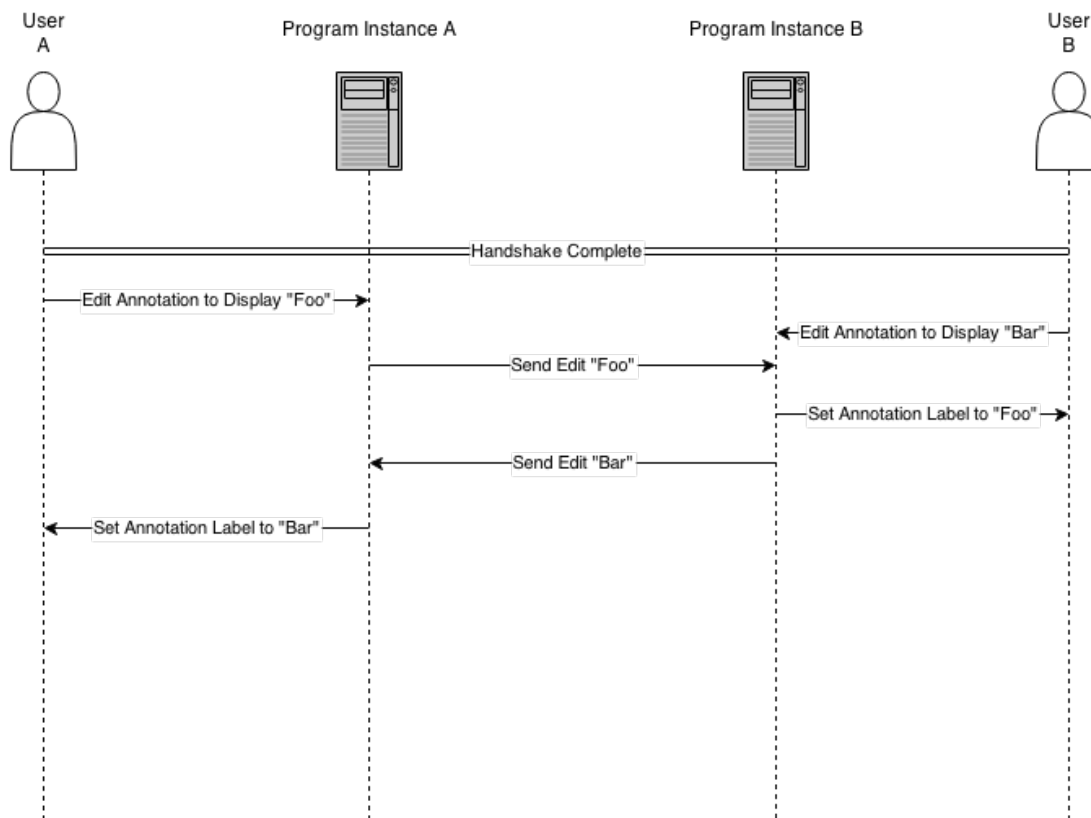


Figure 3.22: Out of sync collaboration

### 3.7.1 Undo & Redo

Shneiderman calls for easy reversal of actions and Nielson calls for user control and freedom – an emergency exit from an unwanted state. To address this, an undo/redo functionality has been added. This required refactoring of the project code, so that the session data is in one location, inside a singleton. Any changes to this data are picked up during the next UI update and are reflected in the visualisations. The session data is stored as a dictionary. To implement undo and redo, copies of the data dictionary are pushed and popped onto the stack. Copies are pushed onto the undo stack on any atomic change the user makes. This gives the user a full session history to go back through and this was one of the early goals from the first project phase.

A problem was encountered when trying to copy the dictionary onto the stack. When just pushing the dictionary onto the stack it would not put a new copy of it onto the stack, so any changes to the dictionary after it has pushed onto the stack are also there in the stack. Python dictionaries have a copy method. Copy only does a shallow copy – any objects in the original dictionary will have their reference placed in the new dictionary. This was fine for some parts of the session dictionary, but for others it was not. In particular, lines and annotations, which are custom objects presented problems. This was solved by using deep copy. With deep copy a new copy is made of objects as well. Some elements of wxPython and matplotlib were unable to be deep copied, but this was fixed whilst focusing more around the data – so the UI elements use the data,



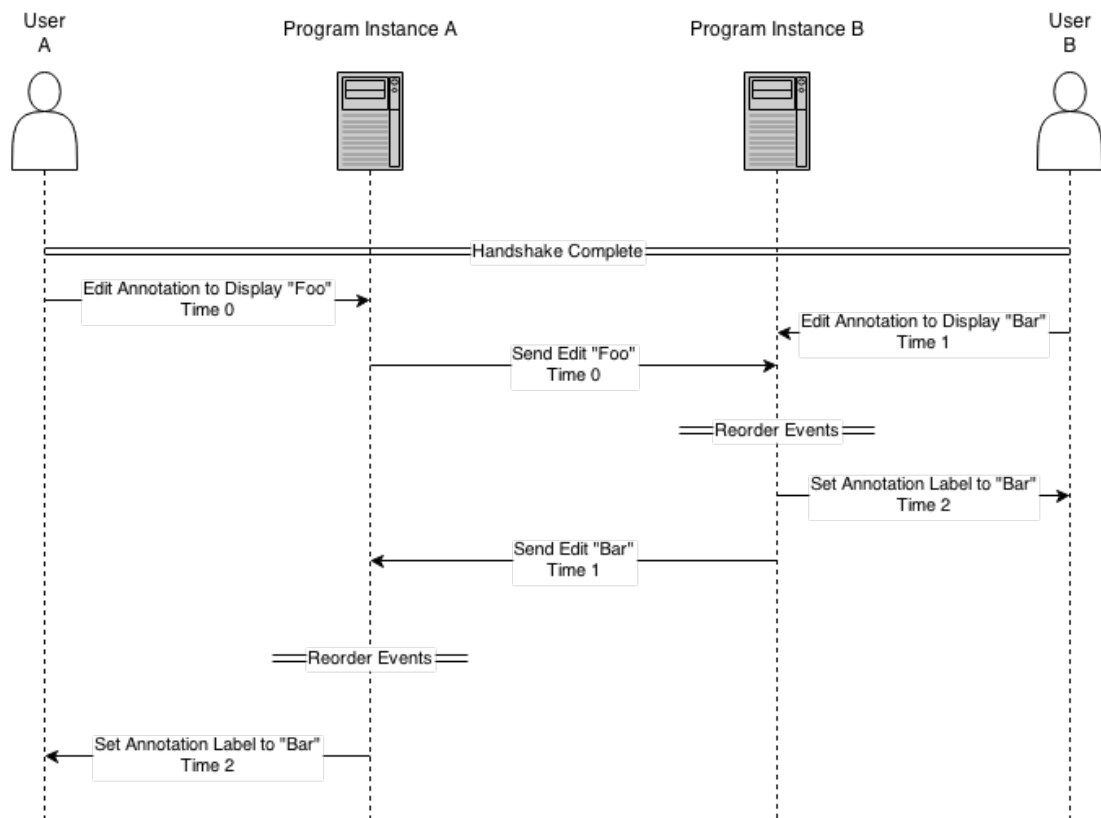


Figure 3.23: Use of Lamport clocks to keep collaboration in sync

not the other way around.

### 3.7.2 Saving

It is important that a user is able to save and load the visualisation session as they may not be able to complete all their analysis in one sitting and may want to come back to their work in the future. Without the ability to save and load the user would have to repeatedly add annotations and change preferences and attach files. It was possible to add Saving and loading by building on the work done to implement undo & redo, although further work was required. Python has a module called pickle to serialise and deserialise data. When saving, the dictionary containing the session data is pickled and written to a file and when loading the reverse happens. Because the program is now focused on the data model, once a previous session has been loaded, a UI refresh is triggered and the visualisation reflects the loaded data.

Saving the data also enables limited collaboration. The user can customize the appearance and add annotations. They can then save the state to a file and email that file to a colleague. The colleague can then load the file and see the user's work. The colleague can then correct any issues and add their own work. The colleague can then save this and email it back to the user. This is useful and is better than no collaboration, but it is entirely non realtime.

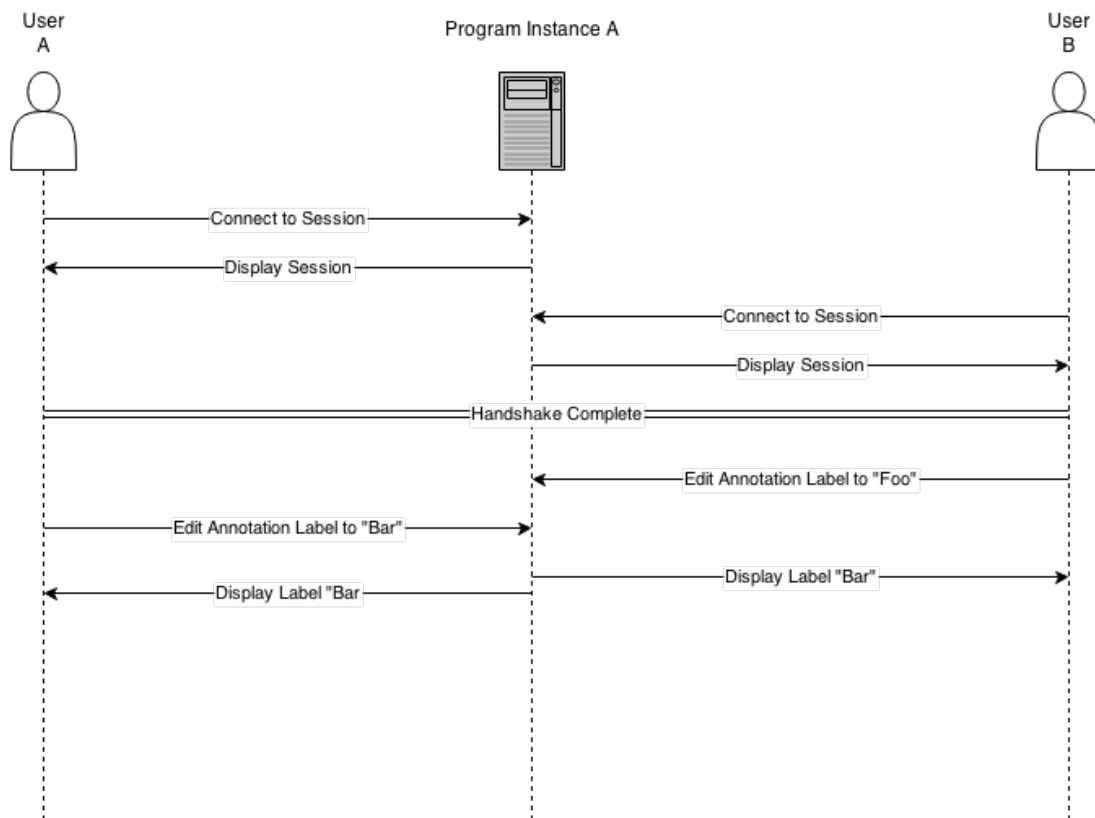


Figure 3.24: Communication flow in a centralised collaboration model.

### 3.7.3 Feedback

Norman and Shneiderman both call for feedback to be given to the user so that the user can be sure that an action has been accomplished. This feedback can come in a number of different forms and was in place in some parts of the project already.

Existing feedback in the project was a natural byproduct of some of the features; for example, when loading a results file the feedback that the load operation has been successful is that a graph appears on the screen. If the graph does not appear then something has gone wrong. Additional feedback has been added to the project:

- When adding annotations the cursor changes to indicate to the user that they can interact with the graph in a different way.
- The title bar text changes to display “unsaved” when the user makes a change and then changes back to “saved” when a successful save has been performed.

### 3.7.4 Guiding the User

The first evaluation of the second phase of the project unearthed that the users struggled to choose the correct action as there were multiple ways of doing the same action that had slightly different use cases. There was also confusing language in the menu

options. These multiple paths have been removed. For example, now there is only one way to open results files initially. To help guide the user further UI elements are enabled and disabled as appropriate. Now when the program is first loaded the only action a user can perform is to load a session or start a new session, or join a session. Afterwards other UI elements are enabled to allow the user to start using the tool effectively.

The new session wizard would typically be the entry route in the first time a user runs the program. It is therefore important that it helps them understand what it is they are doing. The process of setting up the session should also be as easy as possible so that the user does not dislike the prospect of using the program. The session wizard can be seen in Figure ??.

Form validators are used on the title fields and results fields. The validators ensure that a title and at least one results file are included. This stops the user from creating an invalid session. There is also help text on each page of the wizard instructing the user on what actions they should take.

Put the figure here for the session wizard

The requirement of the model file for parsing animation has also led to animation replacing the previous model visualisation. In the set up phase after the the model and the species have been parsed the user is presented with a cell segment, similar to what is seen in the animation panel. The cell segment is split into different regions, one for each region of the cell. These segments are then coloured if the selected species is present in them. The user can select between all species in the selected results files. This has a number of benefits. First, they can sanity check that they have matching results and model files. Second, they can see how the model is structured.

To make setting up the animation user friendly to control required the model file so that the location hierarchy could be parsed. Before this there was an awkward system where the user had to input where in the cell a species is. This was time consuming, awkward and quite brittle. At the time it assumed that there would be three compartments in the species, which is a terrible assumption to make.

## 3.8 Data Manipulation and Export

In a meeting with the developer of BioDARE it was discovered that biologists found the ability to export their data very useful. This allows them to use their data in external applications as required.

Bio-PEPA's CSV format is non standard. The first lines of a results file contain meta-data about the experiments. In the worst case these lines have the potential to cause an external CSV parser to parse the data incorrectly. In the best case the user has to delete them. Allowing them to export the data makes it easier for them by removing potential errors and unnecessary steps.

Another feature that the developer of Bio-DARE recommended was allowing the biologists to normalize their data. Biologists will often normalise their data in the y-axis. This puts all of the data on the same scale. This makes the data much easier to analyse if there is a significant difference in the scale. Without it the user would be required to swap between different zoom levels. Normalising the data makes the user's analysis much easier.

There are different normalisations that can be applied. The data can be normalised to be zero mean and unit variance as it is for Section 3.5. Another normalisation is to normalize all the data to be between zero and one, this can be calculated using the minimum and maximum data values from the species in the results. BioDARE allows the user to choose what type of normalisation they would to use on their data. For this project zero to one normalisation is the only normalisation that the user can have displayed on the graph. This has been done to make the choice simpler for the users. They do not have to think about which normalisation they should apply. In the future if desired more normalisation options could be added. Currently if they want to use a different normalisation the user can export the data and normalise that using their desired technique.

Give an example of normalised graphs?

The zero to one normalised data is also exported along with the original data. This allows the user to manipulate the normalised data in external applications if they desire.

# Chapter 4

## Evaluation

### 4.1 Evaluation

#### 4.1.1 First Evaluation

The first evaluation in the second phase of the project occurred in November 2014. The user group was made up of two people. One who had taken part in the first evaluation meeting and one person who had no knowledge of the project.

##### 4.1.1.1 User Group

As I did not have a domain expert available I was not able to do insight based evaluation. I took a more traditional approach. Before the evaluation I prepared a typical scenario that a user might encounter. The task was to open a file, annotate it and run the animation visualisation, and attach supporting documentation. The task was prepared at two levels of instruction. The first level was a paragraph of text that described what was to be done. The second level was a step-by-step list of instructions to perform. I observed the user group as they attempted the task and offered assistance when required. Afterwards the user group was given a questionnaire to fill in about their experience. After filling in the questionnaire we went through and discussed their answers and any further thoughts that they had.

The task was prepared at two levels to try and gauge how easy the program is to use. The users were first presented with the textual description and if they had been unable to complete the task with this then they would have been given the step-by-step instructions instead. The users were able to complete the task from the textual description alone. This is a good sign that the new tool is usable.

Some issues were encountered:

- The users were unfamiliar with MacOS – Both users were unable to locate the menu bar as it is not attached to the program as in Windows. Future evaluations will use Windows.

- The users were unclear as to what was going to happen when annotating. When annotating the graph with an arrow the user has to click twice to place it but there is no indication of this, nor was it clear to them which way the arrow would be drawn. This has now been fixed. Different cursors are used to give feedback to the user that they should click, and rather than just relying on two clicks with no information as to where the arrow is going to point, after the first click (which places the tail of the arrow) an arrow will be drawn that follows the cursor until the second click placing the annotation.
- Lack of ability to edit, move, or delete annotations – Once an annotation was placed it was there permanently. The ability to edit annotations was always planned, but had not been implemented in time. But the amount of frustration it gave the users was very high. It was a principle in all three of Norman, Neilson and Schniederman's lists that a user should be able to fix mistakes. Since the evaluation, editing and deleting of annotations have been implemented. This means any mistakes can be corrected.
- Initially they were confused by what all the buttons on the matplotlib toolbar did. After discovering the tooltips and seeing what effect the buttons had they were comfortable with them. If a user were to do something they did not intend they are able to undo it. All the matplotlib built-in buttons on the toolbar can be undone and redone from the toolbar. Any buttons implemented for this tool are covered by the undo and redo functionality implemented across the whole program. Being able to recover from their actions on the toolbar means no hindrance to discovery and so needs no further action. It would be desirable to have the two undo methods unified but a way to do this could not be found.
- The users were confused by some of the terminology. In particular "save graph" and "save model". These items in the menu have now been grouped more carefully to help the user distinguish them. A related issue was worrying that "save graph" was going to override the results file. To rectify this the menu items that create new files have been renamed "export ...".
- The users struggled to start a new session. When asked for a title they did not know what the title was going to be used for. When trying to add files, rather than use the add files button in the dialogue, they tried to use the file menu. Having two routes into the visualisation seemed to be confusing them. Now the file menu open file has been removed. To create a visualisation the user has to go through the new session wizard.
- When placing species in the cell one of the users did not understand what they were being asked to do. One of the users did understand. To fix this user input has been removed from the equation. This has required the model file to also be chosen, but then species locations are parsed automatically.
- They liked the animation feature and thought it would be very useful (One of the users did their PhD in transport and expressed a desire to have had this feature during the PhD). They did feel that it wouldn't be useful directly for papers, but that it would be useful when deciding what to include in a paper.

- One of the users asked if there was a map of the cell. When presented with the model visualisation they thought that it did look nice, but they were unsure of its usefulness. The model viewing has since been merged into the animation visualisation.
- The results from the questionnaire indicated that both users thought the tool's appearance was good. The tool was average in difficulty to use – neither easy nor difficult. The annotation buttons on the toolbar were clear as to what they did. It was obvious how to attach supporting files. Both users thought that it is very useful to attach files to the session so that they can be easily emailed to a colleague. They thought it would be useful to have the graph automatically annotated, but they wanted the ability to disable any automatic annotations. Both users found the animation useful.

#### 4.1.1.2 Personal

At this stage in the development the program was in a state where some existing functionality had been broken and gone unnoticed during the implementation of the new features. This highlighted architectural flaws in the code. There were multiple paths through the program that data was taking, and duplicated code in places. Since then a majority of these bugs have been ironed out and the duplications removed. The code much better architected. At the time of the evaluation with the users not all the features could be tested with them – mainly the plot preferences dialogue. These features have now been fixed and they will be evaluated by the users at the next meeting.

Having the users use the program has also highlighted a number of usability problems: menus being badly organised and named, features such as annotation rely on assumed knowledge to work them. All this created an unfriendly environment for the user. This was due to losing sight of the need for usability during development and when testing new features not removing the knowledge of the code from my mind. Since this evaluation the three lists of usability have been refocused on and the code gone through and the principles applied.

I am pleased with the positive feedback on animation and annotation – two of the core new features.

## 4.1.2 Evaluation 2 - Start of Second Semester

### 4.1.2.1 User Group

### 4.1.2.2 Personal

## 4.1.3 Evaluation 3 - End of Second Semester

### 4.1.3.1 User Group

### 4.1.3.2 Expert User

### 4.1.3.3 Personal

## 4.1.4 Overall Self Evaluation

Scribble over lots of stuff talk about changes

talk about the architecture

talk about perhaps being more strictly mvc

## 4.1.5 Search Results

move this into evaluation?

Initial results are very promising with the tf.idf weighted cosine providing much clearer results than with simple euclidean distance. There is not a large enough truth set of plot similarities to be able to confidently say that it is effective. However these early promising results seem to indicate that further research would certainly be worthwhile, but outside the scope of the project.



# **Chapter 5**

## **Conclusion**

### **5.1 Conclusion**

#### **5.1.1 Comparison to Objective**

#### **5.1.2 Challenges Faced**

#### **5.1.3 The Future**

#### **5.1.4 Final Remarks**



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