

User Manual

for

T.E.A.R.S. - 1.0

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For more information, visit our web-page at:

https://www.math.lsu.edu/courses/capstone_course/Summer2015

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Introduction

The screenshot shows the T.E.A.R.S. software window. On the left, there is a 'General Information' panel with the following text:

- Number of Groups: 2
- # of participants in Group 1: 27
- # of participants in Group 2: 31
- Max Number of Trials: 72
- Percent of Answers Correct: 74.8563
- Average number of fixations: 81.3534
- Average trial Duration: 22.8998 seconds

The main window displays a table with the following columns: ID, IsExpert, Tester, Trial, Difficulty..., IsCorrect, Image, Ave_X, and ξ . The table contains 38 rows of data, with the first row highlighted in blue.

	ID	IsExpert	Tester	Trial	Difficulty...	IsCorrect	Image	Ave_X	ξ
1	1		2 c31km1	1	2	1	M10high....	627.3276	15
2	2		2 c31km1	2	3	1	Z17high....	859.6727	22
3	3		2 c31km1	3	3	1	Z10.png	901.4324	30
4	4		2 c31km1	4	3	0	Z06.png	896.2960	25
5	5		2 c31km1	5	2	1	M24high....	788.7034	22
6	6		2 c31km1	6	3	0	Z05.png	820.9056	27
7	7		2 c31km1	7	3	1	Z23.png	862.2863	23
8	8		2 c31km1	8	1	1	L13low.png	784.6350	28
9	9		2 c31km1	9	1	1	L11.png	844.1810	24
10	10		2 c31km1	10	3	1	Z24low.p...	593.2250	16
11	11		2 c31km1	11	2	0	M21low.p...	765.3425	30
12	12		2 c31km1	12	2	1	M18.png	825.2906	24
13	13		2 c31km1	13	3	1	Z09.png	827.6648	25
14	14		2 c31km1	14	3	1	Z02low.p...	866.6433	23
15	15		2 c31km1	15	2	1	M07low.p...	959.6143	18
16	16		2 c31km1	16	1	1	L18.png	824.5832	24
17	17		2 c31km1	17	1	1	L14high....	748.8000	20
18	18		2 c31km1	18	1	1	L24.png	860.0769	26
19	19		2 c31km1	19	3	1	Z15.png	811.7696	23
20	20		2 c31km1	20	3	1	Z08low.p...	826.9956	25
21	21		2 c31km1	21	2	1	M11high....	813.8983	24
22	22		2 c31km1	22	1	1	L05.png	842.7718	25
23	23		2 c31km1	23	2	0	M01.png	883.6565	23
24	24		2 c31km1	24	2	1	M20high....	741.5533	27
25	25		2 c31km1	25	1	1	L07low.png	862.7919	25
26	26		2 c31km1	26	1	0	L15low.png	829.9189	25
27	27		2 c31km1	27	1	1	L09low.png	617.7000	17
28	28		2 c31km1	28	3	0	Z13high....	753.4518	23
29	29		2 c31km1	29	2	1	M09low.p...	806.9467	24
30	30		2 c31km1	30	3	1	Z16low.p...	898.3367	24
31	31		2 c31km1	31	3	0	Z14.png	921.7480	22
32	32		2 c31km1	32	3	1	Z07low.p...	811.4306	23
33	33		2 c31km1	33	2	1	M12low.p...	829.4000	23
34	34		2 c31km1	34	3	0	Z12high....	762.3901	27
35	35		2 c31km1	35	2	1	M23high....	850.5167	26
36	36		2 c31km1	36	3	1	Z20high....	682.8222	24
37	37		2 c31km1	37	2	1	M08low.p...	746.7096	23
38	38		2 c31km1	38	3	1	Z11high....	746.1173	23

T.E.A.R.S. (Time-intensive Eye-tracking Analysis and Research Software) is a toolkit for analyzing arbitrary subsets of fixation data, and uses several powerful comparison algorithms to find patterns and recurrences within these data sets. In order to run **T.E.A.R.S.** the user must have a copy of **MATLAB 2014** and must also have the **Image Processing toolbox**. As the software name suggests, many of the processes run in **T.E.A.R.S.** are rather time consuming. To avoid possible errors, the programmers suggest reading and/or referring to this user guide when using any tool within **T.E.A.R.S.**

To run **T.E.A.R.S.**, first open **MATLAB** and select (double-click) the folder-path in the '**Current Folder**' panel on the left containing '**tears.m**'. Then, simply enter '**tears**' into the command window of **MATLAB**.

Please note that all of the data outputted by **T.E.A.R.S.** Is completely dependent on what is contained within the files inputted into the '**New Project**' menu. Blank cells and other unspecified inaccuracies in the data fed to our GUI may cause errors beyond the scope of this guide, and could possibly invalidate outputs given by any and/or all tools within **T.E.A.R.S.** Additionally, all fixation data should be scaled to a resolution of **740x580** for accurate results.

Excel files must be formatted in a particular manner, described in detail in the section dedicated to the '**New Project**' menu.

The comparison methods used in this GUI are as follows :
Levenshtien Distance, ScanMatch, MultiMatch, and Cross-Recurrence.

We hope your experience with our software is fantastic! If there are features you would like to see in future versions of **T.E.A.R.S.** please let us know. Thank you.

New Project

Project Name

of Expert Levels # of Difficulty Levels

Create Project

Load Data

Load File (.xls, .csv, .mat) Expert Level

Browse Load Data

Make Statistics Table

	1	2
1		
2		
3		
4		

The '**New Project**' menu requires first that you enter a name for your project and the number of groups being compared in your study. If applicable, you may also input the number of difficulty levels present in the trials. Once all of the information is added, select the '**Create Project**' option.

Now it is time to read in the data from your study. Hit the '**Browse**' button to search through your computer and add the relevant files either from the comma-separated-value (**.csv**) or Excel (**.xlsx**) file type. The labels necessary for reading the data are as follows, all fields case-sensitive:

RECORDING_SESSION_LABEL : The identifier of each subject

TRIAL : The number of the trial

CORRECT : Whether the subject gave a correct (**denoted by 1**) or incorrect (**denoted by 0**) response

CURRENT_FIX_DURATION : Fixation duration data
CURRENT_FIX_INDEX : The index of each fixation
CURRENT_FIX_X : The X coordinate value of each fixation
CURRENT_FIX_Y: The Y coordinate value of each fixation
DIFFICULTY_LEVEL : The difficulty of each trial

If your study uses image files, you will also need the column :

IMAGE : Identifier for the image used in each trial

Additionally, many of this project's functions REQUIRE that your fixation data is SCALED. The resolution of your fixation data, for accurate results, should be 740x580.

Additionally, if you are using images, you will be prompted to copy any image files (.png files give the best results of the common image file types) into the 'images' folder within the 'Project Name' folder.

Each data file you upload should contain the data of at most ONE group that you are testing. For example, if you are testing three different groups, you will need at least three different Excel files.

The 'Group Number' drop-down menu allows you to select the Group that each set of data is being added to. For example, suppose that all of the data for Group 1 is split amongst three Excel Files. You could load each of these into the Group 1 selection.

Every time you want to add a data file into the folder, select it along with the group number, then hit the 'Load File' button.

Now, you're all ready. Hit the 'Make Statistics Table' button, and go out and get a cup coffee; this feature can take a significant amount of time to complete, especially for larger data sets.

Once this process has finished, close the 'New Project' window. Your project is now saved and can be opened by the 'Open Project' function whenever you need it!

Information Gain

	Information Gain	Separating Value	Group 1 i...	Group 2 i...	Group 1 i...	Group 2 i...
Average of Y-Fixation	0.5141	468.4790	0.0833	0.9167	0.8707	0.1293
Reccurence Measure	0.3644	2.7625	0.7405	0.2595	0.0868	0.9132
Average of Fixation Duration	0.2252	257.6345	0.2481	0.7519	0.8040	0.1960
Std deviation of Fixation Duration	0.2051	107.8094	0.2712	0.7288	0.8110	0.1890
Reccurence Laminarity	0.1998	51.3044	0.9058	0.0942	0.3736	0.6264
Std deviation of Y-Fixation	0.1932	178.3085	0.1978	0.8022	0.7017	0.2983
Reccurence Determinism	0.1719	38.0289	0.7902	0.2098	0.3562	0.6438
Reccurence Center of Mass	0.0949	31.1323	0.3467	0.6533	0.5644	0.4356
Trial Duration	0.0647	37455	0.2366	0.7634	0.5588	0.4412
Average of Saccade Length	0.0644	100.1745	0.4100	0.5900	0.8392	0.1608
Number of Fixations	0.0306	96.5000	0.3366	0.6634	0.5464	0.4536
Growth Rate of Fixation Duration	0.0286	0.1484	0.4069	0.5931	0.6315	0.3685
Average of Angle between Saccades	0.0106	1.1400	0.4903	0.5097	0.3369	0.6631
Growth Rate of Saccade Direction	0.0105	-0.0057	0.4332	0.5668	0.5624	0.4376
Growth Rate of Angle between Saccades	0.0094	0.0037	0.5160	0.4840	0.4441	0.5559
Std deviation of Saccade Direction	0.0093	1.9380	0.3814	0.6186	0.5024	0.4976
Growth Rate of Y-Fixation	0.0082	-0.8857	0.4332	0.5668	0.5509	0.4491
Std deviation of Saccade Length	0.0072	291.0704	0.8286	0.1714	0.4594	0.5406
Growth Rate of Saccade Length	0.0067	-0.6369	0.4329	0.5671	0.5215	0.4785
Std deviation of Angle between Saccades	0.0062	1.0211	0.3954	0.6046	0.4897	0.5103
Growth Rate of X-Fixation	0.0036	-0.9161	0.4449	0.5551	0.5217	0.4783
Average of Saccade Direction	0.0031	0.5645	0.3265	0.6735	0.4725	0.5275

The '**Information Gain**' window calculates the information gain with respect to **Groups** (Level of Expertise) or **Correctness**.

Group Number is assigned during the creation of a project for each uploaded Excel file. It is aimed to reflect the Expertise Level. For instance, Group 1 may refer to Novices while Group 2 refers to Experts. *Correctness* reflects whether or not the trial is correct (referred as Group 1) or incorrect (referred as Group 0).

If you open the window at the first time, you need to choose one of the options '**with respect to Groups**' or '**with respect to Correctness**'. Then click on '**Calculate**' button. The calculating will take several minutes, after which you will see a table similar to the one on the figure. The result is saved into a file named '**gainWrtGroups.csv**' or '**gainWrtCorrect.csv**'.

The screenshot shows a window titled "Information Gain" with two radio buttons: "with respect to Groups" (unselected) and "with respect to Correctness" (selected). There are "Calculate" and "Close" buttons. Below is a table with 7 columns: Information Gain, Separating Value, Group 1 i..., Group 0 i..., Group 1 i..., and Group 0 i... (repeated). The table lists 25 characteristics of a path.

	Information Gain	Separating Value	Group 1 i...	Group 0 i...	Group 1 i...	Group 0 i...
Trial Duration	0.1770	60000	0.0140	0.9860	0.8172	0.1828
Growth Rate of Fixation Duration	0.1398	2.5663	0.9922	0.0078	0.6411	0.3589
Number of Fixations	0.1211	41	0.6393	0.3607	0.9736	0.0264
Growth Rate of Saccade Length	0.0739	-1.8429	0.6887	0.3113	0.9800	0.0200
Reccurence Laminarity	0.0692	26.7661	0.6762	0.3238	0.9105	0.0895
Reccurence Determinism	0.0668	5.4887	0.7001	0.2999	0.9772	0.0228
Growth Rate of Angle between Saccades	0.0629	0.0105	0.9689	0.0311	0.6937	0.3063
Reccurence Measure	0.0606	5.5261	0.9702	0.0298	0.7015	0.2985
Growth Rate of Y-Fixation	0.0594	4.2213	0.9780	0.0220	0.7001	0.2999
Reccurence Center of Mass	0.0589	22.8458	0.6737	0.3263	0.8671	0.1329
Growth Rate of X-Fixation	0.0505	10.7312	0.9924	0.0076	0.7131	0.2869
Std deviation of Y-Fixation	0.0501	137.2285	0.7099	0.2901	0.9831	0.0169
Growth Rate of Saccade Direction	0.0498	0.0288	0.9774	0.0226	0.7083	0.2917
Average of Saccade Length	0.0291	140.4444	0.8569	0.1431	0.6826	0.3174
Std deviation of Saccade Direction	0.0204	1.7164	0.7263	0.2737	0.9365	0.0635
Average of Angle between Saccades	0.0175	1.7405	0.9656	0.0344	0.7336	0.2664
Std deviation of Angle between Saccades	0.0174	0.7647	0.7370	0.2630	0.9945	0.0055
Std deviation of Fixation Duration	0.0167	288.3385	0.9340	0.0660	0.7301	0.2699
Average of Saccade Direction	0.0160	-0.3983	0.7301	0.2699	0.9280	0.0720
Average of Y-Fixation	0.0142	600.5003	0.9712	0.0288	0.7369	0.2631
Std deviation of Saccade Length	0.0104	111.7049	0.8240	0.1760	0.7143	0.2857
Average of Fixation Duration	0.0032	459.8686	0.9008	0.0992	0.7440	0.2560

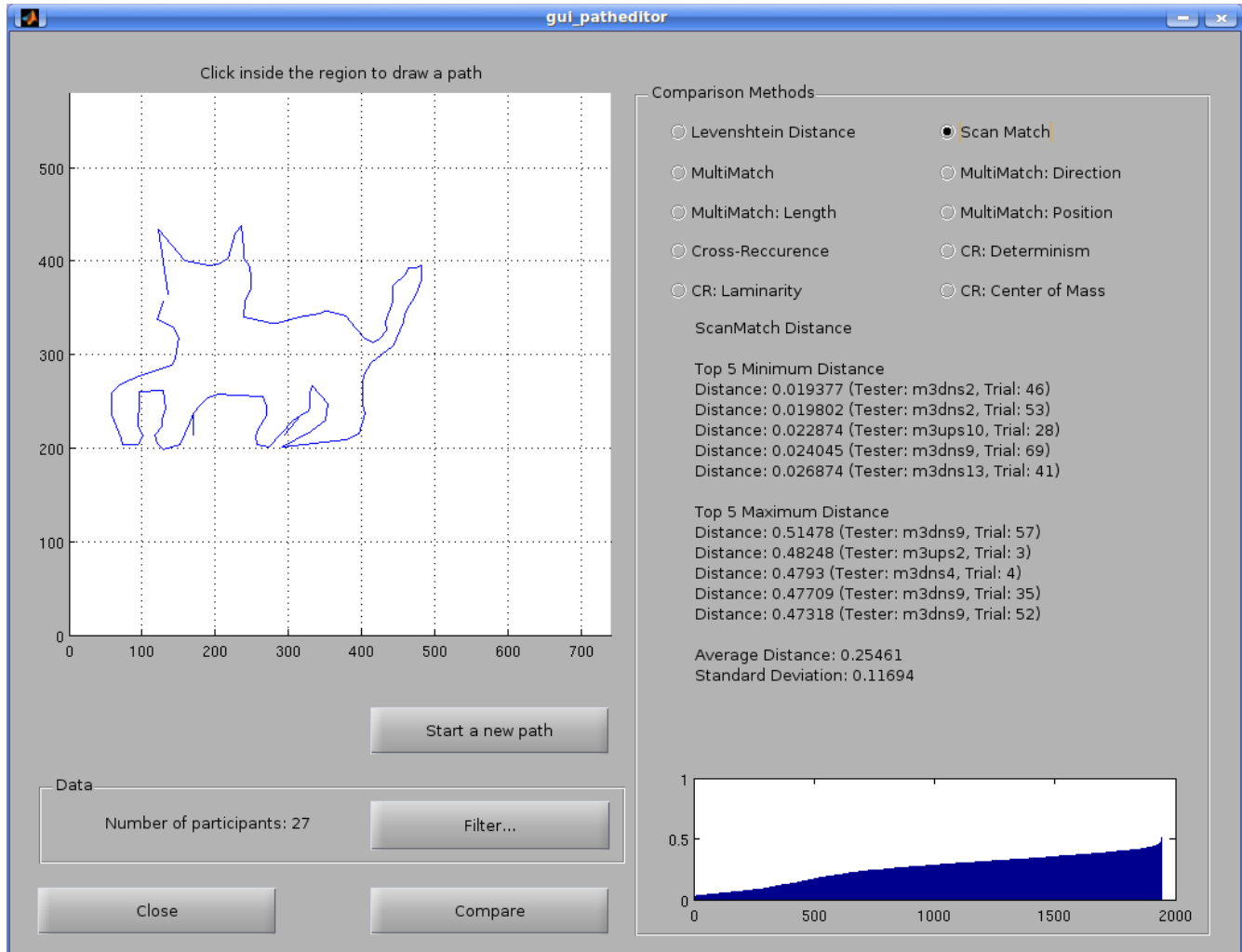
In the table, each row corresponds to one of the characteristics of a path. The meaning of the columns is following:

- **Information Gain:** (takes values from 0 to 1) the bigger information gain the more significant the characteristic in distinguishing trials between Groups or between Correct and Incorrect answers.
- **Separating Value:** while calculating Information gain, whole range of values of a characteristic are divided into two intervals. For instance, all values of **Average of Y-Fixation** are divided into those with the average greater then **468.4790** and those with the average less then **468.4790**.

- Meaning of the next column will depend on the selection of **Information Gain**:
- If '**with respect to Groups**' is selected
 - **Group 1 in (C > V)**: fraction of the trials among the total number of trials with the characteristic greater then the separating value, that are performed by participants from Group 1. For instance, **8.33%** of the trials with **Average of Y-Fixation** greater then **468.4790** are performed by participants from Group 1.
 - **Group 2 (C > V)**: fraction of the trials among the total number of trials with the characteristic greater then the separating value, that are performed by participants from Group 2. For instance, **91.67%** of the trials with **Average of Y-Fixation** greater then **468.4790** are performed by participants from Group 2.
 - **Group 1 in (C < V)**: fraction of the trials among the total number of trials with the characteristic less then the separating value, that are performed by participants from Group 1. For instance, **87.07%** of the trials with **Average of Y-Fixation** less then **468.4790** are performed by participants from Group 1.
 - **Group 2 in (C < V)**: fraction of the trials among the total number of trials with the characteristic less then the separating value, that are performed by participants from Group 2. For instance, **12.93%** of the trials with **Average of Y-Fixation** less then **468.4790** are performed by participants from Group 2.
- If '**with respect to Correctness**' is selected
 - **Group 1 in (C > V)**: fraction of correct trials among the total number of trials with the characteristic greater then the separating value. For instance, **1.4%** of the trials with **Trial Duration** greater then **60** seconds are correct.
 - **Group 0 in (C > V)**: fraction of incorrect trials among the total number of trials with the characteristic greater then the separating value. For instance, **98.6%** of the trials with **Trial Duration** greater then **60** seconds are incorrect.
 - **Group 1 in (C < V)**: fraction of correct trials among the total number of trials with the characteristic less then the separating value. For instance, **81.72%** of the trials with **Trial Duration** less then **60** seconds are correct.
 - **Group 0 in (C < V)**: fraction of incorrect trials among the total number of trials with the characteristic less then the separating value. For instance, **18.28%** of the trials with **Trial Duration** less then **60** seconds are incorrect.

Remark: the information gain value can be misleading. Notice that **Trial Duration** has a high information gain with separating value of **60** seconds because **60** seconds is the time limit for a trial. The high information gain on **Average of Y-Fixations** can be a result of that fixation coordinates are not scaled or not aligned properly.

Group Path Comparison

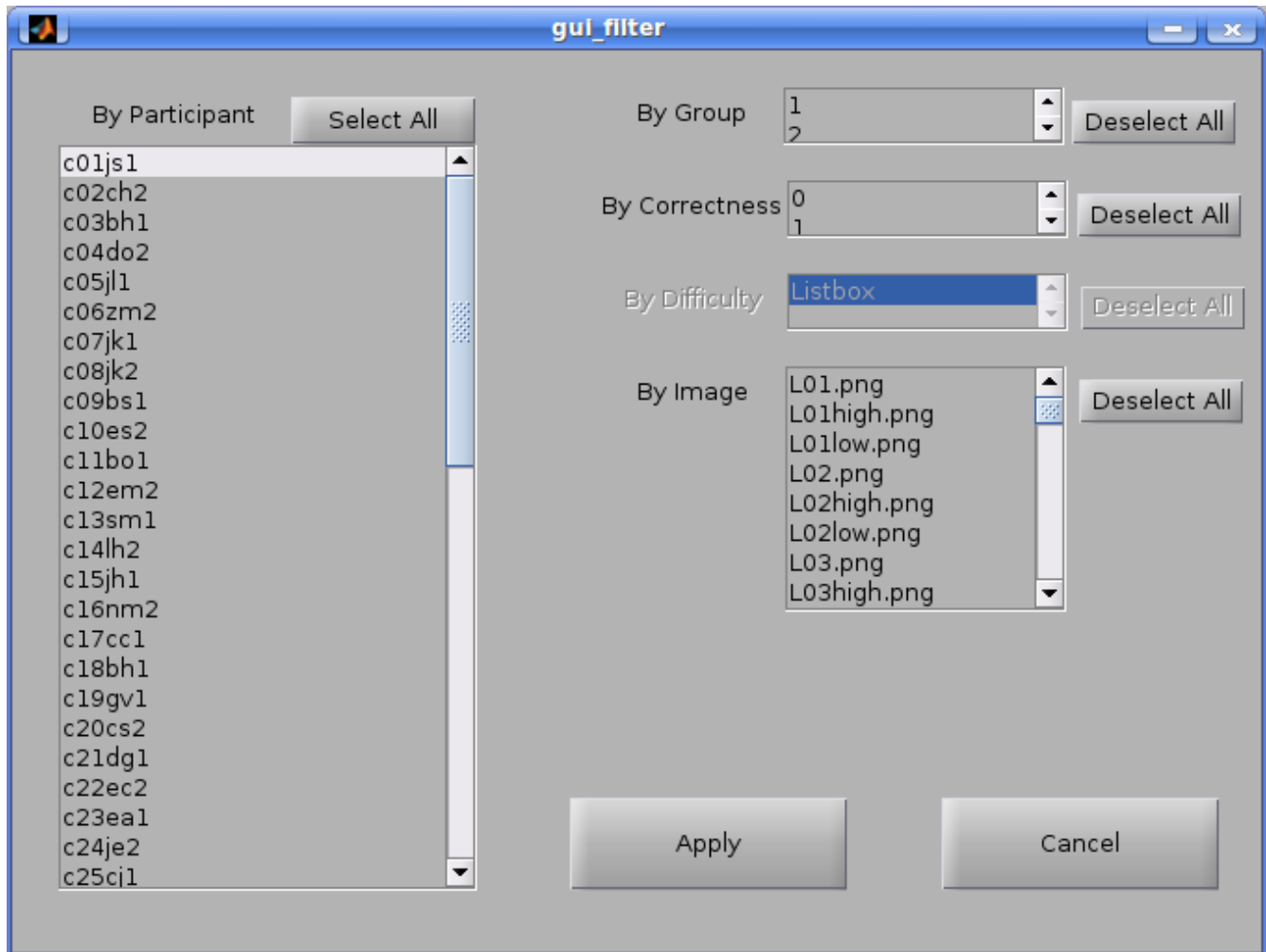


The '**Group Path Comparison**' tool allows one to draw a path on the provided grid and compare it to the eye-tracking fixation data.

To begin, click on the given grid. This will create an initial point to begin the drawn path. Subsequent clicks will draw a line between the point being clicked on and the previous point. The world is your oyster; no matter how simple or complex your path, our software will work. For the rest of this demonstration, our feline friend will be used as an example path.

At any time, one can hit the '**Start a new path**' button, which will clear the grid and allow you to start anew.

After drawing a path, select the **Filter** button. From here, one is able to determine the subset of data to be compared to the drawn path.



Remember, the **Ctrl+Click** function allows the selection of multiple elements.

As you can see in the above image, data can be filtered by individual participants, group number, correctness of responses, and images.

When using the filter, ALWAYS select which participants are being filtered, otherwise, the system defaults to a single participant.

When you are happy with your filter, hit the **Apply** button to save your settings.

Now you are ready to start the comparison algorithms! Hit the **Compare** button button. This can also take quite a bit of time, depending on the size of the subset of data being considered.

Once this process is done, one can select the type of comparison method being used, and **T.E.A.R.S.** Will display the top 5 minimum and maximum distances (trial number and participant), as well as the average distance and standard deviation. On the next page, we will describe each of the comparison methods in detail.

Levenshtien distance : A metric for determining the 'distance' between two strings. In the case of eye-tracking data, this string is a sequence of fixations. The distance is given by the number of single character insertions, deletions, and substitutions to change one string into another. Of course, for computational feasibility in this instance, the space on which fixation information is plotted **MUST** be discretized by a large-enough mesh.

ScanMatch : This is an improvement on using Levenshtien distance for fixation data, implementing the Needleman-Wunsch algorithm to compare eye-movement sequences. In general, ScanMatch is robust in measuring sequence deviation, where it becomes more inaccurate with respect to wild geometric deviation. (Scaling is very important here.)

MultiMatch : Compared to ScanMatch, most implementations of MultiMatch are more geometrically robust, whereas it is not as good at measuring sequence deviations. There are four implementations of MultiMatch used in **T.E.A.R.S.**, each measuring different aspects of the data. Unlike in ScanMatch, areas of interest are not used.

Shape : Vector distance between aligned saccade pairs, measure of scanpath shape similarity.

Length : Measures similarity in saccadic amplitude by finding the length between vector endpoints.

Direction : A measure of angular distance between saccade vectors, useful for determining shape similarity when amplitudes are wildly different.

Position : Measures scan path similarity based on Euclidean distance.

Cross-recurrence : In general, Cross-recurrence is a good tool for the analysis of temporal dynamics of fixation data. **T.E.A.R.S.** makes use of four different CR methods, described below.

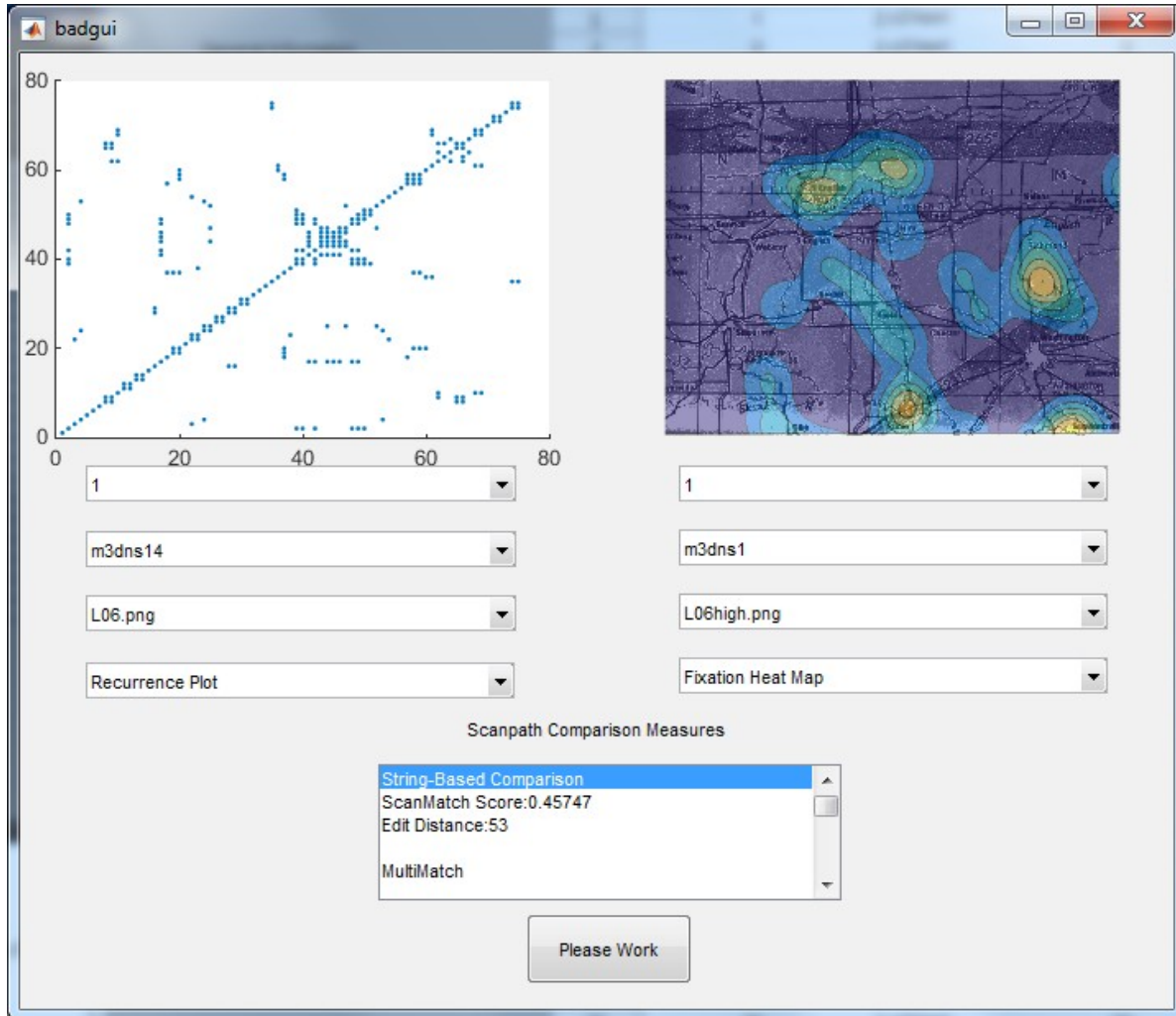
Recurrence : Good at measuring how often observers return to previously fixated image positions

Determinism : Measures the proportion of recurrent points forming diagonal lines and represents repeating gaze patterns in the recurrence diagram.

Laminarity : Generally indicates that specific areas of a scene are repeatedly fixated.

Center of recurrence mass : Measure that indicates approximately where in time most of the recurrent points are situated.

Pairwise Path Comparison



The '**Pairwise Path Comparison**' window calculates various scanpath comparison measures, given two trials selected by the user.

To choose the trials you wish to compare, you must select the group, subject and mapid (trial), in the first, second and third drop down menus respectively. The names listed in each drop-down menu are taken from the raw data (Excel files). In the fourth drop-down menu you can choose which visual representation of the scanpath you wish to view. To compute the various scanpath comparison measures, click the '**Please Work**' button at the bottom of the GUI.

In the example above, the data was grouped in “pilots” and “students” labeled ‘1’ and ‘2’ respectively. Each member (subject) of a group was given a unique name, in this case we have selected “m3dns14” and “m3dns1”. The trials themselves are labeled by the image files that the subject viewed, in this case “L06.png” and “L06high.png”. Finally, we have chosen to display the recurrence plot and fixation map.

Scanpath Visualization

Fixation Maps

To generate the fixation maps, we place a 2-D Gaussian (bump) centered on each fixation in the scanpath. By doing this we create a surface on our 780x540 plot, where the height at a given point represents the attention given to that point on the image. We then can generate a contour plot of this surface and impose on it the original map (if one exists).

For the scaled fixation maps, the height of each Gaussian bump was scaled by the duration of the corresponding fixation. The unscaled maps treats each fixation equally.

Scan Path

For the scan path plot, we simply place a line plot of the scan path over the original image (or a blank plot window if no image exists)

Recurrence Plot

The recurrence plot is an n-by-n scatter plot, where n is the length of the fixation sequence (scanpath) in question. A filled-in circle is placed at the point (i,j) if fixation i and fixation j are “recurrent” in the sense described in the “Cross-Recurrence analysis” section below. Essentially, two fixations are recurrent if they are close based on Euclidean distance. These are the diagrams from which the recurrence measures are derived.

Scanpath Comparison

String-Based Comparisons

For string-based comparisons, each scanpath is first encoded as a string. Usually this is done by superimposing a grid onto the image and assigning each fixation a character based on which grid element it is contained in. One can also use areas of interest (AOIs) for creating strings.

Edit Distance: The distance is given by the number of single character insertions, deletions, and substitutions to change one string into another. This distance is given by a non-negative integer, a distance of 0 indicates two identical strings and a distance of n indicates the minimal number of “edits” required to transform one string into another.

Edit distance can also be computed using AOIs for trials which use the same image. See **Areas of Interest**.

ScanMatch : This is an improvement on using Levenshtien (edit) distance for fixation data, implementing the Needleman-Wunsch algorithm (used for comparing DNA sequences) to compare eye-movement sequences. ScanMatch scores returns a number in the interval (0,1], where 1 indicates the same string and increasing string dissimilarity as the score gets closer to 0.

MultiMatch : MultiMatch returns several scores based on various geometric and temporal aspects of the given scanpaths. MultiMatch first simplifies both scanpaths (by combining some pairs of successive saccades/fixations), then uses a graph-theoretic path minimization algorithm to align the two paths. Given the simplification/alignment, MultiMatch then computes five different measures by comparing paired saccades/fixations (the pairing given by the alignment). Scores are given by a real number between 1 and 0, here 1 indicates maximal similarity and 0 indicates maximal dissimilarity. If any fixations are outside of the range (assuming 780x540 image), MultiMatch returns nothing.

Shape: Measure of overall scanpath shape similarity, based on aligned saccades

Length: Measures similarity in saccadic amplitude of two scanpath.

Direction: A measure of angular distance between saccade vectors, useful for determining shape similarity when amplitudes are wildly different.

Position: Measures scan path similarity based on Euclidean distance.

Duration: Compares aligned fixations based on duration

Cross-Recurrence:

For cross-recurrence analysis (CRQA), two scanpaths are reduced to a sequence of fixations and truncated so that both sequences have the same length. Given two fixations from two sequences we wish to compare, we can say these fixations are recurrent if their Euclidean distance is less than 64 pixels. Based on this definition of recurrence, we can define several measures. **T.E.A.R.S.** makes use of four different CR methods, described below.

Recurrence: Measures total percentage of recurrence over all pairs of fixations. A value of 100 indicates maximal possible recurrence. In this case, both fixation sequences would lie with the *same* 64 pixel radius ball. A value of 0 indicates no recurrence, meaning for both sequences, no given fixation from sequence 1 is within 64 pixels of a fixation in sequence 2 and vice versa.

Determinism: Measures the percentage of recurrent points forming diagonal lines and represents repeating gaze patterns in the recurrence diagram. High determinism indicates recurrences between two scanpath occur in sequential order in both scan paths.

Laminarity: Measures the percentage of recurrent points forming horizontal or vertical lines in the recurrence diagram. High laminarity indicates that for some fixations there are recurrent fixations which appear in sequential order.

Center of Recurrence Mass: Measure that indicates approximately where in time most of the recurrent points are situated.

Fixation Position Based Measure: These scanpath comparison methods essentially use only the coordinates of the fixations comprising each scanpath.

Mannan Distance:

“Analyzes the overall similarity between two scanpaths by computing the linear distances between two scanpaths by computing the linear distance between the fixations in the first scanpath and the nearest neighbor in the second scanpath [and vice versa]”

The following two scanpath comparison measures are based on the data used to generate the fixation maps. To generate the fixation maps, we place a 2-d unit Gaussian (bump) centered on each fixation in the scanpath. By doing this we create a surface on our 780x540 plot, where the height at a given point represents the attention given to that point on the image.

Several methods of comparison were developed for these surfaces. For the **KullBeck Liebler Distance** and the **Correlation Coefficient**, we treat these surfaces as probability density functions (PDFs) on the 780x540 space and use methods from probability theory to compare various PDFs.

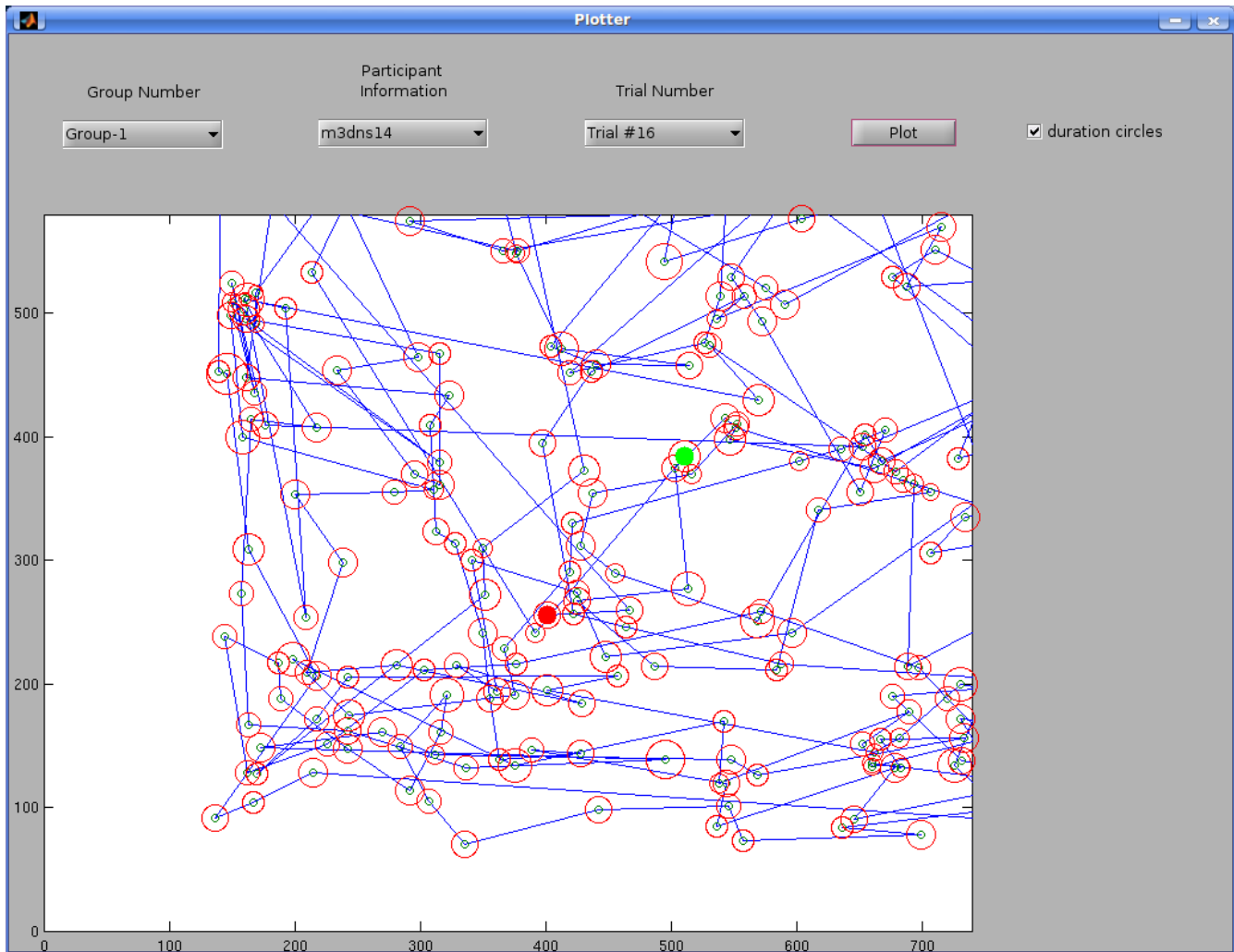
KullBeck Liebler Distance:

Returns a non-negative real value. A value of 0 indicates maximal similarity (both PDFs are the same) a value greater than 0 indicates increasing dissimilarity

Correlation Coefficient:

Returns a real number in the interval $[-1,1]$. A value approaching 0 indicates increasing dissimilarity and a value with absolute value approaching 1 indicates increasing similarity.

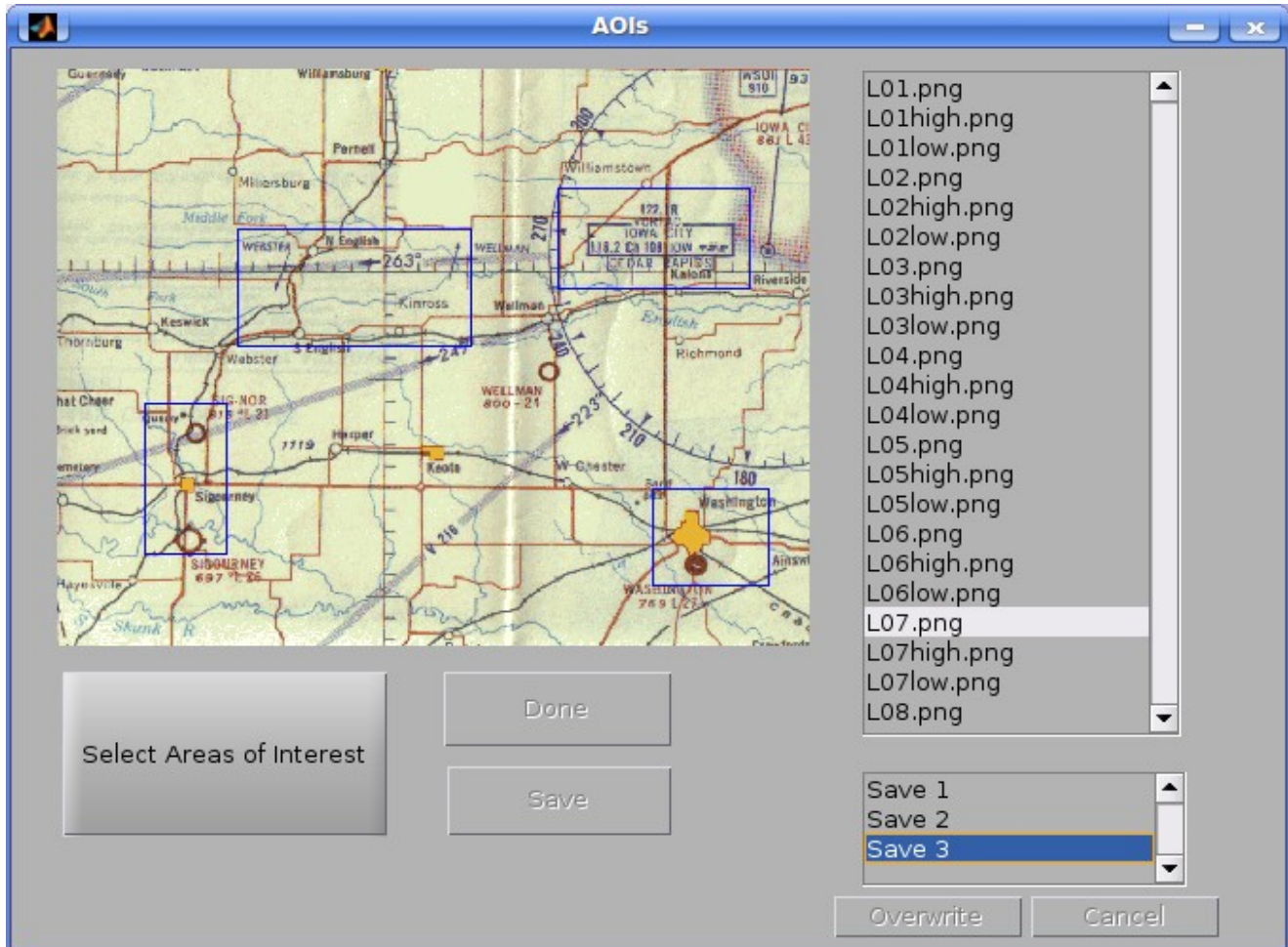
Fixation Plotter



The '**Plotter**' is a very simple tool to use! First, select the Group Number from the leftmost drop-down menu. This will populate the '**Participant**' drop-down menu directly to its right. Select one of these, and the '**Trial**' drop-down menu will be populated. Select one of these, and hit the '**Plot**' button. This will plot the fixation data for that trial, with the filled-in green circle representing the first fixation, and the filled-in red circle representing the final fixation.

The check-box '**duration circles**' will create a representation of how long each spot is being fixated on; the larger the radius of the open circle, the longer that fixation-duration.

Areas of Interest



T.E.A.R.S provides analyses based on user-input '**Areas of Interest (AOIs)**' for images used in a Project (see **Pairwise Path Comparison**). The Areas of Interest menu is where the user can draw and save these AOIs.

The names of the images saved in the '**images**' folder of the current Project will be listed on the left side of the Areas of Interest window. Select an image's name to display it. Any existing AOIs saved for the selected image will appear in the smaller listbox; clicking on these will show the saved AOIs on the current image.

To draw AOIs for an image, select that image's name and then click '**Select Areas of Interest**'. This will allow you to draw AOIs on the image until pressing '**Done**'. (**MATLAB** may report an "Interruption during mouse selection" error when '**Done**' is pressed, which can be safely ignored.) To erase the AOIs you have just drawn and select new AOIs, click '**Select Areas of Interest**' to draw new AOIs for the current image, or select a new image. To save the AOIs you have just drawn, click '**Save**'. You will see a new Save added to the save list for the current image.

Each image holds a maximum of three Saves. If you click '**Save**' for an image with three existing Saves, you will be prompted to overwrite an existing Save, or '**Cancel**'. If you click '**Yes**' at the overwrite prompt, you will be able to select the existing Save you wish to overwrite. Clicking '**Overwrite**' will permanently overwrite the selected Save with the current AOI data. Clicking '**Cancel**' here or at the overwrite prompt will return you to the option of saving current AOIs or selecting new ones.

AOIs can be used in **Pairwise Path Comparison** to compute **edit distance** for associated trials. In contrast to the edit distance which is reported automatically, where characters are assigned to each fixation based on their location in a 5x5 grid, fixations under the AOI edit distance are assigned to characters based which AOI they fall in. For instance, a trial with three AOIs and fixations falling in the following order:

1 – 1 – No AOI – No AOI – No AOI – 2 – 3 – 3 – 3

would be given the string 'aaddbccc'. Note that the smaller alphabet will tend to make AOI edit distances much smaller than standard edit distance.

To compute AOI edit distance, go to the **Pairwise Path Comparison** menu and select the same image for both trials. If there is saved AOI data for this image, AOI edit distance for each Save will be reported at the end of the **Scanpath Comparison Measures**.

References

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4. Dewhurst, R., Nystrom, M., Jarodzka, H., Foulsham, T., Johansson, R. & Holmqvist, K. (in press). It depends on how you look at it: scanpath comparison in multiple dimensions with MultiMatch, a vector based approach. *Behaviour Research Methods*
5. Mitchell, Tom M. *Machine Learning*. New York: McGraw-Hill, 1997. Print.
6. Scanmatch: <http://seis.bris.ac.uk/~psidg/ScanMatch/>
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