

# Infomat Manual

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**Magnus Rosell**

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# Chapter 1

## Introduction

Infomat is both a processing tool and a visualization tool. This manual will (when it is complete) deal with both. The visualization will be covered in Chapter 2 and the (non visual) processing will be covered in Chapter 3.

*Please note!* that this is work in progress: both program and manual. This manual is definitely not complete and may in part be out of date as I develop the program all the time. Still, I hope it will come to some use.

Further information can be found in the `readme.txt` file, the javadoc of the program, which can be found in the `doc` subdirectory, and on the Infomat website:

<http://www.csc.kth.se/tcs/projects/infomat/infomat/>

### 1.1 Infomat Basics

Infomat deals with objects that are called *IObjects*. Each *IObject* has a string and an id number that uniquely identifies it. It also has, when applicable, a reference to a location where the actual object is stored (like a actual text file). In this manual they often will be called *objects*, for short.

Several *IObjects* can be stored in an *IObjectGroup*, and several *IObjectGroups* constitutes an *IObjectGrouping*. Through this manual these are also called groups and groupings for short. Right now each *IObject* can belong to only one *IObjectGroup* in every *IObjectGrouping*.

The main data structure in Infomat is a matrix, called an *IMatrix*. It is an implementation of a sparse matrix<sup>1</sup>. The objects along the axes of the matrix, rows and columns, corresponds to *IObjects*. Each axes has a special *IObjectGroup* called an *IObjectSet*. An *IObjectGrouping* can only contain *IObjects* from one *IObjectSet*.

The *IMatrix* stores several *IMatrixCells* which holds information of the relation between two *IObjects*, one from each *IObjectSet*. The basic information is a count, and a derived information is called a weight.

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<sup>1</sup>I am slowly developing a dense matrix structure, which might be useful sometimes. However, for now all matrixes are handled as sparse. As the intended use of Infomat is Information Retrieval it is not a problem. For the GUI to be really useful the objects along the axes of the matrix has to be interpretable. When they are, the matrix is usually sparse and/or small.

For a typical Information Retrieval scenario the row IObjects may constitute texts, with titles and locations in the file system, and the columns words that appear in the texts. For each word that appear in a particular text an IMatrixCell with the number of appearances is stored as the count. The weight of the IMatrixCell can be calculated through a weighting scheme. An IObjectGrouping of the texts (rows) could be a clustering or a categorization of the texts.

*Any information stored in a matrix may be investigated using Infomat.*

## 1.2 Bugs

Though I have spent a considerable amount of time developing Infomat, there are probably several bugs. When using the Infomat GUI it is a good idea to keep an eye on the terminal window. Some trace text is printed there. Also, if any of the internal functions do not work properly for some reason, Infomat won't shut down. It will only be indicated by the Exceptions that are printed in the terminal. For many such Exceptions you will be able to continue working, but that particular function did not have the desired effect.

## Chapter 2

# Infomat as a Visualization and Exploration Tool

In this chapter Infomat as a visualization tool will be described. It allows you to display a matrix, and group, order and alter it. You may do this along the rows or columns. This chapter describes the GUI in an order that follow the layout. It should be considered as a reference.

The last two sections are a bit different. Sections 2.2 and 2.11 describes the most important concepts of the GUI and Section 2.12 describes the example matrix that is bundled with the program.

*There is no undo function. Save your work!*

### 2.1 Interface overview

Figure 2.1 shows the interface. The **main window** (rightmost window) is divided into five sections. At the top is the **menu**, below that the **toolbar** and under that the **grouping panel**. The **main view** is the fourth section, and the last section (the **mouse pointer information panel**) displays information of the position of the mouse pointer (the picture position and the corresponding (first) row and column object of the current matrix).

There are several other windows that appear in certain situations. From the beginning the **Overview** (top left) is shown. The **pixel window** (bottom left) is vital for obtaining textual information for parts of the matrix. Here the matrix elements for the pixel the mouse pointer is pointing to are displayed.

In the following sections the main window sections and several other windows will be described briefly. Section 2.10 describes a few GUI components that appear in several places. Finally, Section 2.12 describes a small example. First, however, a short account of the visualization.

### 2.2 Matrix Visualization

Infomat stores a matrix, which is displayed in the main view and the overview. This picture is called a *SparsePicture* and consists of *SparsePixels*.

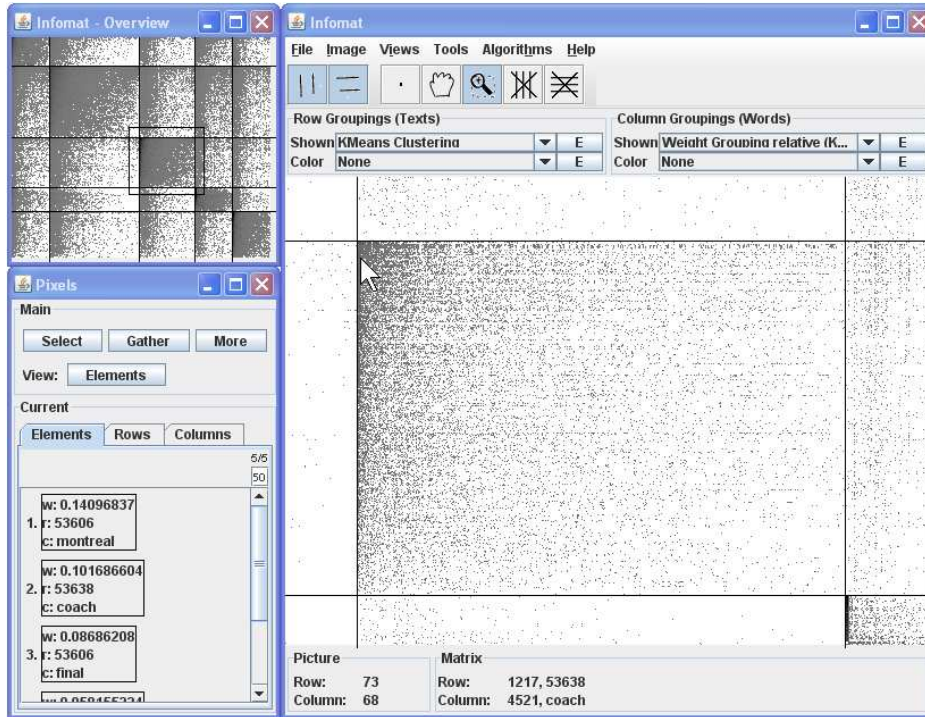


Figure 2.1: The Infomat interface. The overview window (above left) shows a set of texts (rows) clustered into five clusters. The words (columns) are clustered into five relative clusters (one word cluster per text cluster). A rectangle indicates the part which is displayed in the main window (to the right). The matrix elements represented by the pixel the mouse pointer is pointing to are listed in the pixel window (below left). The non zero elements are presented with their weight (w), and the row (r) and column (c) objects. In this example texts have numbers as names. The bottom part of the main window shows which pixel the mouse pointer is at and which is the first text and word that it represent in the current ordering (both the order number and the string).

If the matrix is bigger than the number of pixels in the main view, each SparsePixel represents several matrix elements.

The opacity of the pixels are proportional to the weight of the matrix elements they represent.

What is said in this Section is probably the most important thing to know before you start using the tool. Almost as important is what is described in Section 2.11. Before you read that you might benefit from learning a bit more about the interface, especially about how Groupings are handled, as described in Section 2.8.

## 2.3 Main View and the Overview

When you load a matrix, see Section 2.4.1, the whole matrix is displayed in the main view and the overview, but many operations result in a partial view. Which part is shown is decided thorough the *Grouping Panel*, see

Section 2.8.1. The Main view, further, may be zoomed in on any part of the partial view. The Overview always displays all of it, and indicate by a rectangle what part the main view shows.

*The main view and the overview display a part of the matrix.*

## 2.4 Menu and Toolbar

This sections contains a short account for the available menu options. As the toolbar contains convenient short cuts to some of the options it is described here as well.

The following subsections describes the content of the menus.

### 2.4.1 File

In the file menu you can save and load matrix files. It is also possible to load a “token file”, which is a single file containing several texts. Look at the example, Section 2.12, for the format.

It is also possible to save the picture in the main view as .png-file.

The Infomat Properties are some fundamental settings for the program. They are displayed and altered through the Properties-GUI which recurs for several settings through the program.

#### 2.4.1.1 File Formats

The xml-formats are quite straight forward. You should be able to figure them out by looking at the examples, see the "readme.txt".

### 2.4.2 Image-menu and Toolbar

The toolbar is divided into two sections, with two and five buttons. The five first menu options on the Image menu corresponds to the five icons in the second button section:

**Pixel selection** When the mouse is clicked on a pixel in the main view information on it is displayed in the Pixel View window, see Section 2.4.3.

**Drag** For moving the selected zoom area.

**Zoom selection** By clicking, dragging and releasing the mouse within the view that area is zoomed in.

**Delete rows** Click, drag and release to remove rows.

**Delete columns** Click, drag and release to remove columns.



The following two menu options corresponds to the leftmost two icons. They toggle the group separators on/off. The last two options in the Image menu toggles the guide lines (that helps with positions) on and off and resets the zoom entirely.

*All Image functions work in both the main view and the overview.*

### 2.4.3 Views

The different Views are the main ways to get information. The options in the Views menu are all toggle options: activating/deactivating the corresponding view.

**Pixel View** The pixel view shows information on the pixel the mouse pointer is currently pointing on and very much more. It is described in Section 2.5.

**Selection View** This part is not yet fully implemented and does not change anything.

**Similarity View** In the similarity window you can change what similarity measure is used for both rows and columns. You may also change the Properties of the chosen similarity. The chosen similarities are applied whenever appropriate: for many clustering algorithms and when sorting IObjects according to similarity in different ways.

**Overview** The overview window shows the entire matrix and indicates which part is currently visible in the main view.

**Grouping Panel** The grouping panel with all its functions is described in its own section, Section 2.8.1.

**Toolbar** The toolbar is described in Section 2.4.2 on the Image menu.

**Mouse Pointer Info** The mouse pointer information panel (at the bottom of the main window) gives direct feedback on which row and column the mouse pointer is pointing to. The leftmost values gives the screen coordinates for the pointer, while the rightmost present objects that correspond to these coordinates in the compressed matrix. There is always only one row and one column object presented. If the matrix is large it is the first object. To get more information (all row and column objects) you should use the pixel view, see Section 2.5.

### 2.4.4 Tools

There are several tools:

**Evaluation** Choose grouping to evaluate (and if you want to make an external evaluation a reference grouping) and press “Evaluate”. The measures can be saved and loaded in an xml-format.

**Matrix Summary** gives some basic matrix information.

**Export to Text** exports the currently selected grouping(s).

**Search** is a search tool similar to search engines. It is described in Section 2.6.

**Stoplist** is a rather complex tool that is described in Section 2.7.

**Purge matrix** removes all objects that are not displayed in the overview. If you for instance have deleted certain uninteresting objects from a grouping this function removes them from the matrix and from all other groupings. Purge matrix is applied the moment you chose it from the menu, without any options window.

**Transpose matrix** speaks for itself. This is applied directly.

## 2.4.5 Algorithms

The options in the algorithm menu are:

**Clustering Algorithms** Here you can choose between several clustering algorithms. See Section 2.9.

**Filter Matrix** The Filter Matrix algorithm is straight forward: alter the Properties and hit the apply-button.

**Weight Matrix** In the Weighting window you can choose between different weighting schemes and alter their properties. When you are satisfied, hit the apply-button. The weightings considered the rows to be the objects, and the columns the representation.

Some things in the properties need explanations:

**tf according to Croft 1983**

$$\text{tf}_{i,j} = c_1 + (1 - c_1) \frac{n_{i,j}}{\max_i n_{i,j}} \quad (2.1)$$

**idf according to Croft and Harper 1979**

$$\text{idf}_i = c_2 + \log \frac{n - n_{\text{word}(i)}}{n_{\text{word}(i)}} \quad (2.2)$$

where  $n_{i,j}$  is the number of times word  $i$  appears in document  $j$  ( $\max_i n_{i,j}$  is the number of times the most frequent word in text  $j$  appears). In the properties  $c_1$  is called (*Local row/global column*) *weight importance factor* and  $c_2$  *Global column weight belief factor*.

**Cluster Sorter** The cluster sorter is work in progress. It allows you to sort the clusters of the current clustering of the rows in order of their value for a evaluation measure. Choose the reference grouping, choose “ascending” or “descending” and hit “Sort by value”.

**Description** This is used to construct a description of a clustering of one dimension in the other dimension. for text clustering groups of words that constitute descriptions for the text clusters. Choose description method (currently only the centroid description), choose whether to use reduction and relative reduction. The two later are described below. Here they are used to decide how many objects should be in the description groups, and (if chosen) to reduce the described clustering accordingly.

**Reduction** This algorithm allows for reduction of all groups in a grouping. There are two types: reduction and relative reduction (choose by clicking the button at the top). The reduction removes objects from each group that are last in the list and/or not similar enough to the centroid for the entire group. The relative reduction considers the groups of a grouping of the other dimension as the centroids for the groups in the grouping about to be reduced. For it to work the grouping along the other dimension has to have the same number of groups.

## 2.4.6 Help

Try them! :)

## 2.5 Pixel View

Through the Pixel View you get textual information about the matrix as ordered by the current groupings. This is a rather complex tool and is described in some detail here.

When you open it the first time it has just two panels. From the top the *main* and *current* panels. When the mouse pointer points to a particular pixel the current panel displays all the matrix elements that the pixel represents.

### 2.5.1 The View button and the Current Lists

The *View*-button in the main panel lets you choose between elements, rows and columns. For elements the *Elements*-tab in the current panel displays all the matrix elements that are represented by the pixel the mouse pointer points to. They are presented as pairs, like: (row-object, column-object) followed by a value, the weight of the matrix element. If you choose the *Rows(Columns)*-tab the row(column)-objects for the matrix elements are presented with the weight of the corresponding elements.

When the *View*-button is set to *Rows(Columns)* the *Elements*-tab does not show anything. The *Rows(Columns)*-tab shows the row(column)-objects associated with the picture row(column).

*The selection (next section) is affected in the same way by the View-button.*

### 2.5.2 Selection

The *Select*-button in the main panel opens (and closes) two panels: the *select* and *selected* panels. These panels allow you to study some objects more thoroughly.

*To select anything pixel selection has to be on. See Section 2.4.2.*

The *Select*-button in the *select* panel lets you choose between *Single* and *All in area*. If you click the mouse on a pixel when set to *Single*, the objects in the current lists are stored in the selected lists. When set to *All in area* you can select several objects by a click-drag-release procedure.

The objects stored in the selected lists are the recently selected. They stay there when you move the mouse.

### 2.5.3 Gathering

The *Gather*-button in the main panel opens (and closes) two panels: the *Copy selected* and *Gathered* panels. The gathered panel allow you to store the addition of several sets of selected objects (as described in the previous Section, 2.5.2).

The *All*, *Elements*, *Rows* and *Columns* buttons in the *Copy selected* panel adds the corresponding list of the selected objects.

The *How* button lets you choose between three things. When it is set to *Accumulate* if an object is already in the gathered list its value is increased with the value it has in the selected list. When it is set to *Add* the latest value for the object is stored, and when it is set to *Set* all previous objects are removed and the selected ones are added.

### 2.5.4 More

The *More*-button in the main panel opens (and closes) three panels: the *Remove Gathered*, *Sort Gathered to Selected*, and *Select Gathered* panels.

The *Remove Gathered* panels allow you to remove the gathered objects from the matrix. The elements are removed from the matrix (it thus affects all groupings), while the rows and columns are only removed from the current shown groupings. To remove these from the matrix and all groupings you need to use the *purge matrix* option in the *Tools* menu, see Section 2.4.4.

The *Sort Gathered to Selected* panel allows you to sort the gathered row(column)-objects in order of similarity to the all row(column)-objects of the selected lists:

**RowRow** The similarity of each gathered row to all the selected row. The row similarity measure (see Section 2.4.3) is used to extract the representation and calculate the similarity.

**RowCol** The similarity of each gathered row to all the selected columns considered as a representation, using the row similarity measure.

**ColCol** The similarity of each gathered column to all the selected columns, using the column similarity measure.

**ColRow** The similarity of each gathered column to all the selected rows considered as a representation, using the column similarity measure.

The *Select Gathered* panel lets you move the gathered objects to the selected panel. There are four straight forward buttons: *All*, *Elements*, *Rows* and *Columns* sets the selected objects, overwriting the previous selected objects. The *El for RC* extracts the matrix elements that intersect with the rows and columns of the gathered objects and sets them as selected objects. The *RC for El* does the opposite.

The last row of the *Select Gathered* panel lets you extract the representation for the objects. With the *C for R* button you set the columns of the selected list to the objects that represents the rows of the gathered list according to the row similarity. The *C for R* button uses the column similarity analogously.

## 2.6 Search

The search tool works similarly to a search engine. You compose a query and then get the resulting list of objects that correspond to that query. It is possible to search for both rows and columns.

The tool consists of three panels. The left one lets you formulate a query; the list of rows and/or columns at the bottom. Enter space separated strings in the text field (you can use javas syntax for regular expressions if you start with a left bracket ([)) and press one of the buttons below. You could also import any list using the “Load” button.

The middle panel is divided into two parts. Press the “Search” button to get the result for the query in the result list below.

The right panel lets you display the search result graphically as a grouping in the main and overview. Choose wether to consider rows or columns, to display the result or the query, if the grouping should consist of one group or one group with the rest in a rest group, and whether the result should be displayed directly or as a coloring.

## 2.7 Stoplist

The stoplist tool is an implementation of the common notion of a stoplist in information retrieval. It can do a little bit more though.

The stoplist window has four panels. The leftmost shows several Properties, that might be altered. The middle panel allows you to load and save a list of ordinary strings from/to a simple text file.

The rightmost third of the stoplist window consists of two panels. The top panel is a list of IObject:s that can be removed (stopped). These might be loaded from an xml file (and saved as well).

The button *From Strings to IO* allows you to convert the strings into IObject:s that can be removed from the matrix. Only IObject:s that exist in the matrix are generated. IObjects may be converted into a list of strings using the *From IO to Strings* button.

In rightmost bottom panel *Main* you choose which matrix dimension that is considered. The *Apply* button removes the IObject:s currently in the IObject list from the matrix and all groupings.

*Using the “From IO to Strings” button you can save any list of objects in a simple text format.*

## 2.8 Groupings and Groups

Infomat stores a matrix. It is displayed in the main view and the overview in order of a row and a column grouping. A grouping consists of one or several groups, which together contain all or some of the IObjects in the matrix. This sections describes how the groupings and groups are managed.

### 2.8.1 Grouping Panel

Through the grouping panel all handling of the groupings is devised. It is divided into two sections, one for rows and one for columns. They work similarly.

The topmost drop down menu displays the currently selected grouping. When you choose the grouping here the order of the objects along the dimension (rows or columns) changes.

The bottom drop down menu selects the coloring grouping. For the rows this leads to a coloring of the pixels, and for the columns a coloring of the background columns. The pixels are averaged over the matrix elements they represents, while the column coloring is averaged over the entire columns.

When the *E*-button beside each drop down menu is pressed a grouping edit window is displayed. It is described in the next Section, 2.8.2.

### 2.8.2 Grouping Edit Window

The grouping edit window looks a little different depending on which of the four groupings it concerns. They all have the following sections:

**Name panel** Here the name of the grouping is displayed. You can alter it.

**Groups panel** Here all the groups are displayed. For each group you can alter the name and press the *E*-button, which opens up a group edit window. It is described in Section 2.8.3.

**Reordering panel** By changing the order of the numbers in the text filed and pressing the *Apply* button you can change the order of the groups in the grouping. If you leave a group out it is deleted - a very convenient way to remove one or more groups.

**File panel** Here you can load and save groupings. For either to work there has to be a matrix loaded.

For coloring groupings you can change the color of each group in the groups panel. The change does not take effect until you press the *Apply* button in the coloring panel which (for coloring groupings) is located between the reordering panel and the file panel. There you can also reset the coloring to the default colors.

The opacity of the pixels can be altered in the “row show” grouping edit window and the opacity of the column coloring in the “column coloring” grouping settings window. By default the column coloring opacity has a lower range than the pixel opacity.

### 2.8.3 Group Edit Window

The group edit window have the following sections:

**Info panel** Here the name of the grouping is displayed. You can alter it.

**Main panel** Here, you can apply any changes you make in the list panel to the actual group, using the *Apply List Order* button.

**List** When you open a group edit window the list panel contains all the IObjects in the group. You can alter it in many ways, using the list manipulations. The similarity that is used is the row or column similarity from the Similarity View, see Section 2.4.3. For the manipulations to affect the group you have to press the *Apply List Order* button.

## 2.9 Clustering Algorithms

There is a Clustering Algorithm Window. In it you can decide if you want to cluster rows or columns. You choose algorithm in a combo box. The algorithms all have some properties that can be altered, like for instance the number of clusters. The algorithm window explains these properties rather well.

**K-Means** K-Means clustering

**Bisecting K-Means** Bisecting K-Means clustering

**Relative Clusterer** An algorithm that cluster the columns (or rows) relative the rows (columns). The column objects that have the highest weight in the first row cluster is assembled into a first column cluster, and so on.

**Appearance Relative Clusterer** An algorithm that cluster the columns (rows) relative to the rows (columns). A column group is created for each row group. Each column object is put in the column group corresponding to the row group in which it first appears (the first non-zero matrix element in the column). The objects are ordered in the column group according to the order in which they appear. Ties are broken by their id number.

**Random Clustering** Just what it sounds like.

**Location Grouper** constructs a grouping based on the location of the objects in the file system, if this information is available.

*The clustering algorithms are applied to the whole matrix, not just the part that is displayed at the moment!*

## 2.10 Standard Components

The GUI makes use of some standard components that appear in several places. This section describes some of their functions in more detail.

### 2.10.1 Buttons

Most buttons have direct effect. There however, are several alternating buttons that only sets the contexts for actions. The typical example the *Choose rows or columns*: button in the Clustering Algorithms window. It alternates between the words *Rows* and *Columns* when you press it, the visible being what you have chosen. Most alternating buttons has a leading text ending with a colon and it should be rather obvious from the context.

### 2.10.2 Properties

A lot of functions could be applied in several different versions. Instead of presenting all of them separately they have properties that you can alter. These properties can be saved in an xml format and recalled. There is also a default setting.

The properties gui is easy to understand. Each property has a value that is displayed. You can alter it by typing in a new value and hitting enter. If your new value is allowed the value of the property changes, if it is not it does not.

Some property values are completely open, you can type in anything you want. If you type something inappropriate here (like for instance -5 clusters in a clustering algorithm) and apply the function Infomat throws java Exceptions.

### 2.10.3 Lists

Lists appear in several functions. They can display IObjects, and IMatrixCells and simple java Strings. Lists have many functions. Depending on the context not all of them are available. It is for instance not possible to load and save IMatrixCells.

The list gui consists of two parts: the list of objects and above that a few functions. The objects are presented in a textual way, often accompanied by a value indicating their order. If the name of the object appears like a button you may open the object in a simple viewer by clicking it. The objects also have a checkbox that you may tick.

The function part of the gui has at most three rows, from the top:



**File** The file row lets you load and save the list of objects. The loading is usually restricted by a `IObjectGroup`, meaning that you will only actually load those that is in that particular group. For the Group Edit window it is the corresponding `IObjectGroup`. In most other cases it is the `IObjectSet` of the `IMatrix` corresponding to the rows or columns depending on the context.

**Sel** The selection row allows you to handle selected objects. The *Sel* button selects all objects, the *Desel* button deselects all objects, the *Rm* button removes all currently selected objects, and the *Inf* button inverses the selection.

**Order** Here you can reorder the objects in the list. There are several possible orderings. You choose between them in the combo box and applied them on the objects by pressing the *Apply* button.

This has no other effect than the ordering in the list. For an ordering to have effect on anything else you have to do more. On a group for instance you have to hit the *Apply List Order* button, see Section 2.8.3.

The two first uses the similarity measure along the rows or columns depending on the context. These are only available in some of the lists.

**Sim. to Sel.** Sorts all objects in order of similarity to the selected objects.

**Similarity** Sorts the objects in order of similarity to all the objects in the list.

**Literal** Sorts the object in literal order.

**Random** Makes a random permutation of the objects

**Invert** Inverts the order of the objects.

**Original** When the list is displayed for the first time it has a particular order. Through this you can revert to it. There is one exception. When hit the *Apply List Order* button in the *Group Edit Window* the new order is set as the original.

To the right in the function part of the gui there is an indicator that shows how many of the objects in the list that are displayed (“50/172”). There is also a small text field in which you may alter the maximal number of objects that should be displayed. Type a new number and press enter.

The gui only present a small part of the entire list for efficiency reasons.

## 2.11 The Matrix – Grouping Concept

Infomat is quite a complex tool. The single most important thing to keep in mind when using it is that the view presents the matrix through a row and a column grouping. The matrix may contain several objects that are not visible. Some functions work on the visual groupings and some work directly on the matrix behind them. This section describes some implications of this fact.

Each grouping is a view of the matrix. Use the purge matrix option in the tools menu to force the matrix to contain only the objects in the current row and column groupings. The objects are also removed from all other groupings simultaneously.

Some functions work on the groupings and some work on the matrix directly. When you remove matrix elements you always remove them directly from the matrix. Row or column objects, on the other hand, are removed either from the current grouping or the matrix directly, depending on the tool you use.

A list of functions that remove row and column objects from the groupings and not from the matrix:

- Through the toolbar and image menu.
- Through the Pixel View window.
- Through the Group View window (after you have pressed the *Apply List Order* button).

When you have removed objects and/or matrix elements (and purged the matrix), remember to weight the matrix again, using the Weight Matrix function in the algorithm menu.

## 2.12 Example

In the directory `/Infomat/example/` you find a few files to start with. Read more in the `readme.txt` file.

# Chapter 3

## Infomat as a Processing Tool

This chapter explains some of the possibilities with Infomat when it is not used with the graphical user interface as a visualization tool. It is divided into two sections, that discuss the command prompt possibilities and the program structure. The later to help programmers to use the Infomat classes for other tasks.

### 3.1 Command Prompt Usage

The simplest class to use is the `ExampleClusterer`. It generates a clustering of the example set that is distributed with Infomat. How to run it is described in the `readme.txt` file and Section 3.1.1.

There are so many parameters to set that I have decided to not let the user set all of them in the command prompt. Instead, for the other classes described here you have to save `Properties` files for the different functions of the program. Properties files are xml files with values. There is a lot of functionality in Infomat that use them.

#### 3.1.1 ExampleClusterer

How to run the `ExampleClusterer` is described in the `readme.txt` file. It runs the K-Means algorithm on the English example. You can not alter anything from the command prompt. This class does the same thing every time, but the result may differ due to K-Means random initialization.

The result is essentially two xml-files, with names like:

- `clustering.1222852580758.xml`
- `clusteringEvaluation.1222852580758.xml`
- `clusteringResult.1222852580758.xml`

where the number is the system clock time. The first file contains a full clustering result.

The second file is the index file for a textual presentation of the clustering result. You may look at it in a browser and follow the links.

The third file contains an evaluation: a lot of measures. It can also be viewed with a browser. It represents a `Measures` object corresponding to

such a class. Section 3.2.2 describes it in more detail, but you will be able to use such files without reading that.

### 3.1.2 What Properties are Available?

The class `WriteProperties` helps with setting up default values for many Properties.

```
..Infomat/>java infomat.WriteProperties <resultPath>
```

writes one Properties file each for several classes to the specified directory.

#### 3.1.2.1 Altering Properties

You can open the xml files in an editor and change the values for the different properties. The properties are rather self explaining. Each of them has a name, a value and a description, that explains its purpose. A Property:

```
<p><n>Name</n><v>Value</v><d>Description</d></p>
```

Some of the Properties that are more difficult to understand have `PropertyValueDescriptions` (`<pvd>` in the xml), that may help you as well. If a Property has one or several `PropertyValueDescriptions` you have to choose one of those values!

*Some Properties are Strings.* In order for that to work they have to have a `<str>`-tag within the value field, for example: `<v><str>a string</str></v>`. All files, and paths are treated as strings.

When you want to set a value to nothing it is accomplished like this: `<v>null</v>`. This goes for Strings as well.

#### 3.1.2.2 Some of the Properties

Here is a short list of some of the Properties files that are generated by the `WriteProperties` class:

**Dot\_Product\_Similarity\_Properties.xml** Properties for the dot product similarity.

**Evaluation\_Properties.xml** Properties for the evaluation.

**IMatrixCell\_Filter\_Properties.xml** Properties for removal of rows, columns and matrix elements.

**IObjectGrouping\_Text\_Result\_Properties.xml** Properties for exportation of results as browsable pages.

**IObjectGroupingIO\_Properties.xml** Properties for exportation of full clustering results. (Not browsable. The files you want if you want to use the result somewhere else.)

**KMeans\_Properties.xml** Properties for the K-Means algorithm.

**Stoplist\_Properties.xml** Properties for the use of a stoplist. These properties are in addition to a file of stop objects (words most of the time), when such a file is specified. If there is no file specified these properties are applied alone. The stoplist file is not applied if this properties file is not in the directory.

The stop file is usually specified in the `Clusterer_Properties.xml` or `Experimentator_Properties.xml` file described below.

**TFIDF\_Weighter\_Properties.xml** Properties for the tf\*idf-weighting scheme.

### 3.1.3 Clusterer

The `Clusterer` runs one clustering, evaluates it and writes the clustering result and the evaluation result to files:

```
..Infomat/>java infomat.Clusterer <properties path>
```

A toy example can be found in `...Infomat/examples/clusterer/`. The subdirectory `properties/` contains properties that sets up a clustering.

```
..Infomat/>java infomat.Clusterer examples/clusterer/properties/
```

generates results in `...Infomat/examples/clusterer/result/`. They consist of two files like for the `ExampleClusterer`.

You may alter the Properties in `properties/`, remove some, or substitute them for others<sup>1</sup>. The `Clusterer` will inform you if something is missing, or if there are too many properties, making them ambiguous.

The properties path has to include several `Properties` files, the most important being the `Clusterer_Properties.xml` file, which sets up the basics for the clustering. Here is a description of the Property:s in it:

**Result Path** A string specifying where the result should be written.

**Stoplist** A string specifying a file with stop-objects (words for instance). Opposite dimension to "Rows as Matrix Dimension". Combined with stoplist Properties from file.

**Rows as Matrix Dimension** Whether rows or columns are regarded. Takes one of two values: `true` and `false`. (Overrides the similarity measures dimension.)

**Matrix Source** A group of Property:s that indicate how the `IMatrix` should be loaded. One of the three has to have a value. The other two has to have value `null`. The three possibilities:

**Matrix File** A string specifying an `IMatrix` file.

**Token File** A string specifying a Token file.

**Inpath** A string specifying a path from where files are read recursively.

---

<sup>1</sup>Like for instance replacing the K-Means algorithm with the Bisecting K-Means algorithm.

**Comparison Grouping** A group of Property:s that indicate another grouping. If there is any the results will be compared to it in the Evaluation. Anyone of the two could have a value, but only one. The two possibilities:

**IObjectGrouping File** A string specifying a file with a grouping that works with the IMatrix.

**LocationGrouper** Construct a grouping by looking at the location of the objects (true or false).

### 3.1.4 Experimentator

With the **Experimentator** class you can perform rather complicated experiments. The principles for running the experiments are the same as for the **Clusterer**, see Section 3.1.4.1. The results may, however, become very hard to overview. Section 3.1.4.2 explains how to generate convenient tables based on the results.

#### 3.1.4.1 Running an Experiment

The **Experimentator** class is very similar to the **Clusterer**:

```
..Infomat/>java infomat.Experimentator <properties path>
```

The difference is that the properties path now may contain a hierarchy of directories that defines several clusterings. At the leaf directories all properties along the path from the root directory must define a clustering. If it does not the **Experimentator** will abort and report this.

The results for each of the clusterings are written in a similar hierarchy with the result path as the root. For each of these directories a subdirectory **properties/** is constructed. In it all properties for the corresponding properties path directory is saved, so it is easy to match results with properties.

In the root directory for the properties there has to be a **Experimentator\_Properties.xml** file, which sets up the basics for all the clusterings. It has the same Property:s as the **Clusterer\_Properties.xml** file (see Section 3.1.3), plus one:

**Number of Repetitions** Number of times the clustering is done for each set of properties.

So in each leaf directory of the result path hierarchy there will be **Number of Repetitions** number of results, for clusterings as specified in the corresponding leaf directories of the properties path.

The **Experimentator** works most efficiently if the preprocessing is defined (through Properties files) in the topmost directories, as it only keeps one matrix in memory at the time<sup>2</sup>.

There are two toy examples<sup>3</sup>:

---

<sup>2</sup>Actually two: the original from the **Experimentator\_Properties.xml** file, and the current preprocessed one.

<sup>3</sup>These and the following examples have line breaks for typographical reasons. They should, however, of course, be written as a single line in the command prompt.

```
..Infomat/>java infomat.Experimentator
                    examples/experimentator/1/properties/
```

which generates results in `examples/experimentator/1/results/`, and

```
..Infomat/>java infomat.Experimentator
                    examples/experimentator/2/properties/
```

which generates results in `examples/experimentator/2/results/`. Looking at these should give you an idea of how to use the `Experimentator`.

Notice that the second example do not have any `IObjectGrouping_Text_Result_Properties.xml` files, which leads to that no textual result files are generated. This is very convenient when you run large experiments with many repetitions, as the results otherwise tend to get huge.

### 3.1.4.2 Extracting Results from an Experiment

The package `mro.util.experimentation` contains classes that helps you extract parts of the results generated by the `Experimentator`. This is actually a very nice functionality for any kind of experiment, but it will be described here in the context of the `Experimentator`. The main objective is to calculate average values and standard deviations for the results in the result directories.

You run the `Experimentator` like this:

```
..Infomat/>java
    mro.util.experimentation.ExperimentResultGenerator <params>
```

If you give no parameters it displays some information. The possible parameter combinations are:

1. `-resultPath <res-path> -averagePath <avg-path>`
2. `-structure <table-file.xml> -latex <res-file.tex>`
3. `-structure <table-file.xml> -matlab <res-file.m>`
4. `-structure <table-file.xml> -csv <res-file.csv>`

We will deal with these in order here. The first generates a similar hierarchical directory structure as the one in the result path (`<res-path>`) in the average path<sup>4</sup> (`<avg-path>`). It will produce an average result file, called `average.xml`, for every directory that has result files. It will also copy any properties files. You can browse the average path hierarchy to compare average results.

Try this for the results for example one:

```
..Infomat/>java -cp classes/
    mro.util.experimentation.ExperimentResultGenerator
        -resultPath examples/experimentator/1/results/
        -averagePath examples/experimentator/1/avg/
```

---

<sup>4</sup>Do not choose the average path to be the same as the result path. If you do that twice the averages from the first run will be included in the new averages.

The three other possible parameter combinations have a lot in common. They all read a structure file in xml-format, that holds information on which directories to process. There are examples on such structure files in both `tables/` directories in the Experimentator examples. There are lots of possibilities in the structure files. By looking at the DTD in the beginning of the file you will be able to figure them out. Section 3.1.4.3 describes the structure file in more detail.

The second possible parameter combination generates a latex table. It reads a structure file (`<table-file.xml>`) in xml-format and writes a latex file (`<res-file.tex>`) corresponding to this structure. Try this<sup>5</sup>:

```
..Infomat/>java -cp classes/
    mro.util.experimentation.ExperimentResultGenerator
        -structure examples/experimentator/1/tables/structure.xml
        -latex examples/experimentator/1/tables/table.tex
```

Compile the resulting latex file and you have a document you can look at. It is a table with average results and (standard deviations). You may want to alter some of the typography in the table, but is a good start when you want to include results in a text of some sort.

The last two parameter combinations also reads the structure file, but they disregard all row and column titles and generate two matrixes corresponding to the structure: one with the average values and one with the corresponding standard deviations. Parameter combination three generates two matlab matrixes in a file, that you can call from matlab to start working with the values. The last combination generates a similar *csv*-file (a semi-colon-separated file) that can be read by for instance MS Excel.

There is one more example to try:

```
..Infomat/>java -cp classes/
    mro.util.experimentation.ExperimentResultGenerator
        -structure examples/experimentator/2/tables/structure.xml
        -matlab examples/experimentator/2/tables/mat.m
```

This structure file use different settings for the extraction of values. Look at the `<pathtable>`. For two values it rounds them to two decimals, for one value to two significant figures, and for one value it keeps the whole calculated value. The last is especially interesting if you will go on working with the values in matlab or some other application!<sup>6</sup>

### 3.1.4.3 The Structure File

The previous section showed how to use the Experimentator. This section describes the structure xml file that is used by the Experimentator.

The most complicated part of the structure file is:

`<measureabbreviationtable>`

---

<sup>5</sup>For this and the following example to work you have to have run the Experimentator examples in Section 3.1.4.1.

<sup>6</sup>You would not want to work with rounded values!



```

<mae><abb>ASS</abb><n>Intrinsic Measures</n>
<n>Evaluated Grouping</n>
<n>Weighted Average Self Similarity</n></mae>
<mae><abb>NMI</abb><n>Extrinsic Measures</n>
<n>Global Extrinsic Measures</n><n>NMI</n></mae>
</measureabbreviationtable>

```

This table defines abbreviations (ASS and NMI in this example) for measures that are used in the actual table. Each `<mae>` (measure abbreviation) consists of an abbreviation and a number of names. The names specify a Measure in a Measures xml file, by giving a series of names of MeasureGroup:s and a name for a leaf Measure. The abbreviations are used further down in the structure file, in the `<pathtable>`.

The `<table>`:s of the `<columntitles>` and the `<rowtitles>` allow you to set up how the headers for both columns and rows should look like in the latex file. `<caption>` defines a caption that will be used in the latex file.

Finally, the `<pathtable>` is where everything happens. here the values for the measures specified in the `<measureabbreviationtable>` are extracted from particular directories. The `<commonpath>` allows you to specify a start for the path to the directories.

The `<pathtable>` consists of rows `<ptr>` and columns `<ptc>`. Each column has the following structure:

```

<ptc><m>ASS</m><rd>3d</rd><p>KM/</p></ptc>

```

where `<m>` should contain a measure abbreviation, and `<p>` a path to a directory (which will be concatenated to the `<commonpath>`).

`<rd>` is optional. If it is not present the full value of the measure will be extracted. This is useful when you like to continue working with the values. If it is present, it should be a figure followed by a `d` or be an `i`. The first leads to values rounded to the specified number of decimals, the second leads to values rounded to integers.

## 3.2 Program Structure

This section deals with the programming issues of Infomat and aims at helping the developer use the different functions. More detailed information about all classes can be found in the javadoc in the directory `/Infomat/doc/`.

The simplest example is the `ExampleClusterer`. How to run it is described in the `readme.txt` file. By looking at the code the programmer will also get the first idea of how to use Infomat when writing own programs.

For now this section is very incomplete. As a first help I give a small uml-diagram in Figure 3.1 for the most central data-structure classes used by Infomat. These classes can all be found in the `infomat.vectorspace` package and its subpackages.

### 3.2.1 Properties

This Section gives a short introduction to the Properties class.

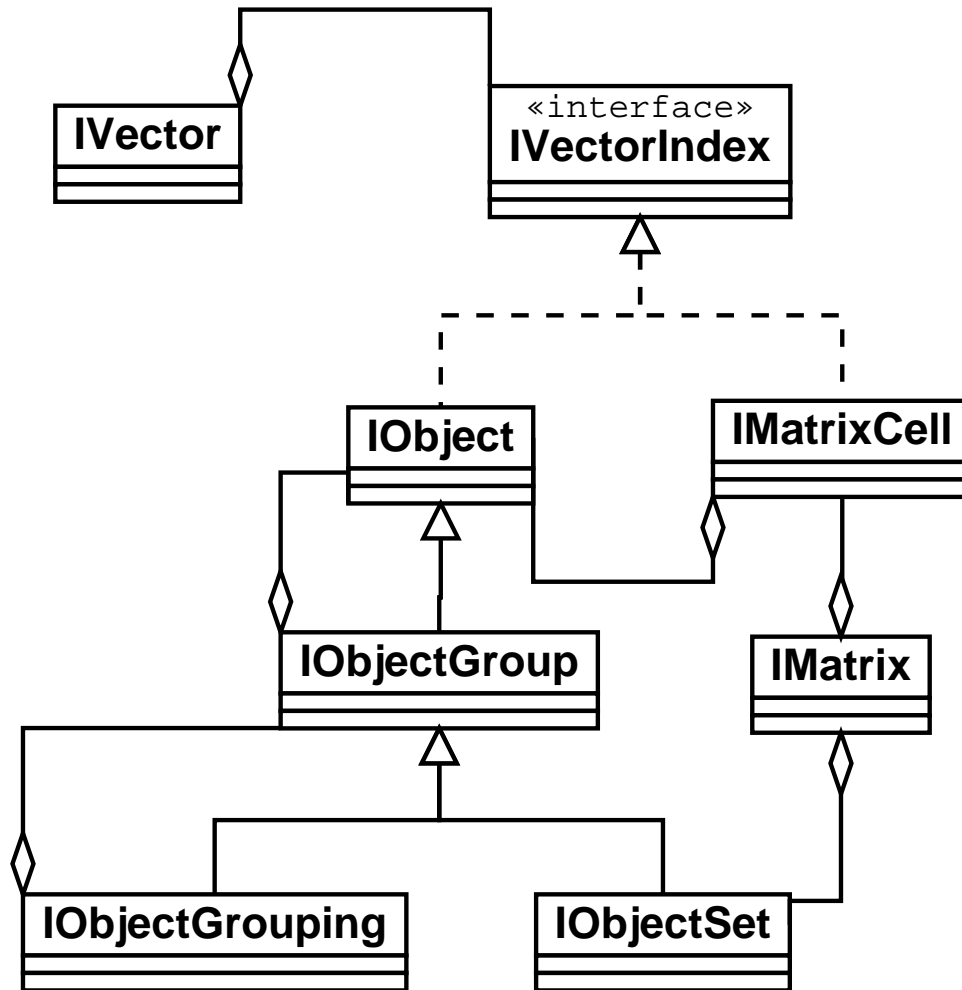


Figure 3.1: UML. Part of the Infomat data-structure

Objects of the **Properties** class contain a lot of **Property**:s that can be grouped into **PropertyGroup**:s to provide more order. Groups can contain groups in a hierarchy.

A short discussion on how to handle xml files representing **Properties** objects can be found in Section 3.1.2.1.

### 3.2.2 Measures

This Section gives a short introduction to the **Measures** class.

Objects of the **Measures** class contains a lot of **Measure**:s that can be grouped into **MeasureGroup**:s to provide more order. Groups can contain groups in a hierarchy.