

# Visual Data Mining and Machine Learning

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**Abstract.** Information visualization and visual data mining leverage the human visual system to provide insight and understanding of unorganized data. In order to scale to massive sets of high dimensional data, simplification methods are needed, so as to select important dimensions and objects. Some machine learning algorithms try to solve those problems. We give in this paper an overview of information visualization and survey the links between this field and machine learning.

## 1 Introduction

The rationale of Information Visualization (infovis [14]) and of Visual Data Mining (VDM [44, 19]) is to leverage the very high processing capabilities of the human visual system to allow interactive exploration and analysis of massive data sets. It has been demonstrated that the low level visual system has preattentive processing capabilities (see e.g. [32]) that enables humans to detect and recognize some features without effort and extremely rapidly, generally in less than 200 ms, even in a large image. As shown in [24], relying on this type of features (e.g. the color of items, their orientation, etc.) enables to display up to one million of items without overloading the human visual system.

Information visualization faces however two major limitations of human vision. Its main limitation is its physical restriction to three dimensional (3D) displays. Moreover, while stereovision hardware (for instance based on shutter glasses) is now affordable, it is still quite uncommon: for the vast majority of users, visualization methods must rely on two dimensional displays (2D). 3D is also intrinsically limited by many problems such as occlusions, disorientation, two dimensional interaction devices, etc.

The second major limitation of human vision is pointed out in [32]: preattentive features cannot be combined freely. If more than two or three such features are used in the same image, they can interfere and reduce greatly the processing rate of the visual system. Therefore, if objects of a data set are described by more than a few attributes, their visualization becomes difficult: a trade-off between completeness of the representation and processing rate has to be made. Complete representations have therefore two drawbacks: they must rely on complex layout methods to transform high dimensional objects into 2D images and they imply a tedious browsing of the full image to obtain a complete understanding of the data set. Simplification methods are therefore needed for visual mining of massive data sets.

Machine learning (in a broad sense) and visual data mining are therefore strongly connected. Machine learning algorithms benefit from expert knowledge:

clustering is easier when the number of clusters is known *a priori*, recognition rates are higher if the training set is free of outliers and if useless variables have been removed, etc. Many of these tasks (outlier detection, number of cluster evaluation, etc.) can be performed by users via a visual inspection of the considered data set: this is exactly the purpose of visual data mining. However, as explained before, information visualization is efficient for simplified data sets: images are easier to read if they represent a small number of objects described by a small number of attributes. Machine learning algorithms can provide the simplifying methods that make visual data mining efficient: dimension reduction can be used to select important attributes, clustering allows one to replace homogeneous groups of objects by some representative examples, etc. Interactive methods can mix visualization and model construction: the user guide the modeling process via the display of results obtained so far (see e.g. [12, 35, 63, 76]).

We survey in this paper the links between visual data mining and machine learning. In section 2 we give a short introduction to information visualization and to its limits. In section 3 we survey dimension reduction techniques that can provide dimension scalability to 2D displays. In section 4 we briefly outline the links between clustering and infovis. We conclude in section 5 with a short overview of two major models coming from machine learning and extremely useful for infovis: the Self Organizing Map and the Generative Topographic Mapping.

## 2 Information visualization

According to [14] information visualization is “the use of computer-supported interactive, visual representation of abstract data to amplify cognition”. In this survey, we focus on a special type of abstract data: each object is a vector from  $\mathbb{R}^p$ , described by  $p$  real values. The descriptors are called variables, features or attributes. This model is frequently named the “table data model” in the information visualization community [19, 36].

### 2.1 Taxonomy of information visualization methods

Several attempts have been made to classify infovis methods in order to get a clear overview of the field (see e.g. [19]). Daniel Keim proposes in [43, 44] to analyze visualization methods according to three “orthogonal” axes: the **visualization technique** itself, the **interaction technique** and the **data type**. Keim’s analysis is based on the fact that interaction methods (such as zooming [7], linking and brushing [6], dynamic distortion [54], etc.) can be freely combined with visualization techniques (see section 2.3) and applied to different types of data (vectors, trees, graphs [33], text [30], etc.).

Card et al. propose in [14] a different taxonomy based on the nature of the information to be visualized. The part of this taxonomy that gathers visual data mining methods is further subdivided according to the data type, in a way quite similar to Keim’s approach.

Unfortunately, those taxonomies don't help in identifying how machine learning methods can be used to improve visualization. For instance, dimension reduction methods are buried in the class of "Geometrically-transformed displays" in [44] or considered as preprocessing steps in [19].

## 2.2 A formal model

Further high level understanding of visualization methods can be obtained with the help of the formal model of Chi and Riedl [18]. In this model, raw data go through four stages via three processing steps. Each step is implemented by an operator that map the representation of the data in one stage to another representation in the next stage (the structure of the data representation is modified). A variation of this model appears in [14].

In the **data transformation** step, raw data are mapped to a mathematical representation (for instance texts are parsed into a vector model of word occurrences); this representation is called the "analytical abstraction". In the **visualization transformation** step, the analytical abstraction is transformed into another representation adapted to visualization (for instance a graph is transformed into a tree by a traversal algorithm in order to use a tree visualization method); this new representation is called the "visualization abstraction". The **visual mapping** step translates the visualization abstraction into a view/image (for instance a tree is visualized with the TreeMap method [71]). Additionally, operators can be used to modify the data representation within one stage: for instance, interaction methods can be considered as operators that modify the view or that have impact on the visual mapping step.

Keim's visualization technique axis corresponds roughly to the visualization transformation step, whereas his data axis gathers the early steps (data transformation and visualization transformation steps). The formal model gives a better understanding of the reasons why Keim's axes are more or less independent ("orthogonal"). In [17], Chi leverages his formal model to produce a taxonomy of 36 visualization methods. He shows how methods are constructed based on standard operators (data extraction, clustering, projection, etc.).

Machine learning methods fit nicely in this formal model and correspond to some operators. For instance, dimension reduction methods are generally operators from the visualization abstraction step: they produce new coordinates from the original mathematical representation.

## 2.3 Some visualization methods for high dimensional data

Keim's visualization technique axis is further subdivided into broad classes of methods. We briefly survey methods adapted to high dimensional data, using Keim's taxonomy. We refer to [44, 19, 14] for comprehensive presentation. Keim identifies four classes of methods described in the following sections.

### 2.3.1 Geometrically-transformed displays

Some methods use a layout algorithm to transform some high dimensional data into low (2 or 3) dimensional data that are displayed by standard methods.

The main standard tool is the scatter plot, i.e. the standard 2D image in which each object described by two attributes is represented by a point whose coordinates are given by the values of the attributes. For  $p$ -dimensional data, a  $p \times p$  scatter plot matrix is obtained by arranging in a matrix all the possible scatter plots. Each attribute appears both as a line and as a column: the image at position  $(i, j)$  is the scatter plot for attribute  $i$  and attribute  $j$ . The diagonal can be used to display some standard graphical representation of each attribute, for instance a histogram. Scatter plot matrices don't scale to a large number of attributes, because the number of scatter plots grows quadratically with the number of variables. Moreover, scatter plots themselves suffer from the superpositions of objects with similar (or close) attribute values. Linking and brushing techniques [6] help nevertheless the user to understand large scatter plot matrices: the user can select a region in one scatter plot and observe the results of this selection in all the plots. Dimension reduction methods (see section 3) can be used to avoid displaying the full scatter plot matrix.

Another standard 2D display is the functional plot in which  $y = f(x)$  is represented by a smooth line obtained by a high frequency sampling of the  $x$ -axis. Data in  $\mathbb{R}^p$  can be transformed into functions, for instance with the method proposed by Andrews in [2]: the coordinates of each object are used as the coefficients of a Fourier series to define a function. The display is obtained by plotting together all the functions on the  $[-\pi, \pi]$  interval (see [29] for a recent application of Andrews' curves).

Another use of functional like plot is the parallel coordinates technique [38, 39]. It consists simply in using as many vertical axes as there are attributes to represent. An object is then displayed as a polygonal line that links the values of its attributes on the corresponding axis. Parallel coordinates don't scale to a large number of objects mainly because of overlapping (see [3] for an example of a rendering method that limits this overlapping problem).

### 2.3.2 Iconic displays

Iconic methods represent each object with a complex icon or glyph [86]. Famous examples of such glyph are Chernoff's faces [16]: each object is represented by a small face where different data dimensions are mapped to different facial characteristics (face width, radius of the eyes, etc.). Other examples include the star glyph [72] and the stick-figure icon [59]. Those methods have scaling issues because representing many characteristics implies to use complex icons that use a lot of space on the screen and limit strongly the number of objects that can be visualized at once. Moreover, comparison of two objects is difficult if the corresponding glyphs are far away from each other: the problem of optimal glyph positioning is therefore quite accurate [86].

### 2.3.3 Dense pixel displays

In pixel oriented methods [45, 43], each attribute of each object is represented by an unique pixel, via its color. This family of methods scales to large number of objects and/or attributes. It has however ordering problems, as the insights on the data it provides depend strongly on the quality of pixel arrangements: pixels corresponding to related objects and/or related attributes should be close in the image. As pointed out in [43], dimension/attribute ordering is in fact an NP-complete problem and only sub-optimal solutions can be obtained in reasonable time.

### 2.3.4 Stacked displays

Stacked displays correspond to methods in which the image is partitioned recursively in such a way that each level of the hierarchy represent one or several attributes of the data (see [51] and [23] for instance). As for many other layout methods, the quality of the visualization strongly depends on the ordering of the dimensions.

## 2.4 Visual data mining tasks

The main goal of VDM is to enable users to explore massive data sets and to search for interesting information. As pointed out in [34], visualization allows to find patterns in the data by proximity and similarity reasoning: plots of the characteristics of objects might reveal dependency between variables as well as clusters of objects. More generally, as shown by P. Hoffman in [36], visual data mining is efficient for many classical tasks, such as: cluster detection, outlier detection, feature importance assessment, feature correlation, prior classification analysis, etc.

Another goal of VDM is to display the results of mining algorithms [47], for instance association rules or frequent patterns extracted from a database, clusters (either extracted by the algorithm or pre-specified in case of supervised learning), etc.

While the bulk of VDM methods is dedicated to unsupervised problems, there is also an important need of visual methods for supervised problems such as classification. Due to size constraints, we won't however cover this important field in this paper (an example of the visualization of classification is given in this volume by [40]).

## 2.5 Links with machine learning

While integrating machine learning and information visualization appears clearly as potentially rewarding (see e.g. the paper [15] by Chen, editor-in-chief of *Information Visualization*), such integration is still rare, with some remarkable exceptions such as multidimensional scaling (MDS, see section 3.1) which is a standard method in VDM and the Self-Organizing Map (SOM, see section

5.1) whose visualization capabilities have been widely recognized by the infovis community.

It is obvious that most of the methods designed to display high dimensional data have scaling problems. In order to display voluminous data sets in which objects are described by numerous variables, they must rely on simplification methods that reduce either the number of variables or the number of objects (or both). Numerous machine learning methods have been designed to tackle those problems. However, infovis methods tend to favor user intervention over automatic methods. For instance, in order to avoid saturation of the human vision system, information visualization uses frequently the concept of “focus+context” (see chapter 4 of [14]): the general idea is to provide a detailed view of a part of the data while retaining as much context information as possible. Distortion techniques have been used to implement this idea (see [54] for a survey). User intervention consists in choosing the interesting part of the data by browsing the summarized version (see [62, 50] for well known examples).

A possible way for building more links between machine learning methods and visualization methods would be to favor user intervention and control. Ward describes for instance in [87] how visual clues allow user to monitor dimension reduction and clustering techniques in order to check whether important information might have been removed. Visual representations of the quality of dimension reduction methods have been produced (see e.g. [4, 5, 9, 76]). Their generalization to e.g. manifold learning methods (see section 3.2), might be a first step toward the integration of those methods in infovis algorithms.

### 3 Dimensionality reduction

While dimension reduction [25, 13, 68] is generally considered as an important pre-processing task in infovis (see e.g. [19]), it seems that only standard methods, such as principal component analysis (PCA) and multidimensional scaling (MDS), are commonly used in visual data mining.

Among classical methods, and apart from PCA, projection pursuit [26] is quite popular in infovis. As Independent Component Analysis (ICA) [41, 37] (but for different reasons), projection pursuit is looking for linear combination of the original features that are non Gaussian. Generalization of PCA such as Hastie’s principal curves and surfaces [31], neural network based non linear PCA [21] or Kernel Principal Component Analysis [69], are not widely used in VDM.

#### 3.1 Multidimensional scaling

In fact, the most popular non linear projection method in infovis is multidimensional scaling (MDS) framework. The main idea of MDS is to compare distances between the objects in the low dimensional space to the corresponding dissimilarities in the original space. Different *stress functions* (i.e. measure of the distortion of the distances) lead to different algorithms. Torgerson’s original MDS (classical metric MDS [79]) works for original data in an Euclidean space and tries to preserve inner products. It also corresponds to finding the linear

projection of the data that preserves best the square euclidean distances between the original observations. It can be shown to be equivalent to PCA.

Commonly used MDS are based on Kruskal's version [48, 49]. Let us denote  $\delta_{i,j}$  the dissimilarity between objects  $i$  and  $j$  in the original space and  $d_{i,j}$  the Euclidean distance between the low dimensional representations of  $i$  and  $j$ . A generic stress function is given by  $\frac{\sum_{i,j} w_{i,j} (f(\delta_{i,j}) - d_{i,j})^2}{\sum_{i,j} (d_{i,j})^2}$ , where  $f$  is a transformation of the original dissimilarities and the  $w_{i,j}$  are weighting coefficients. Variations on the normalization method, the coefficients and on the transformation lead to Sammon's non linear mapping (NLM [66], see also [55] for a smooth version based on a MLP), to Curvilinear Component Analysis (CCA [20]) and to other variants such as the non metric scaling (in which  $f$  is monotone) that tries to preserve ranking between dissimilarities rather than their actual values.

It should be noted that MDS methods are quite computationally intensive. As a consequence a lot of work has been done in order to reduce their actual cost (see [73] in this volume). Fast Map for instance [22], a well known dimension reduction algorithm in the visualization community, is in fact an approximate realization of MDS, as shown in [60].

### 3.2 Manifold learning

While MDS methods overcome some limitations of PCA and related methods, they still have an important shortcoming: if the data happen to belong to a subspace of the original space, MDS methods might fail to discover this fact when the shape of the subspace is complex. The class of manifold learning techniques try to overcome this limitation for instance by feeding a MDS like method with "smart" dissimilarities. One popular idea of those methods is to work at the local level: the structure of the manifold around a data point is described by the  $k$  nearest neighbors ( $k$ -nn) of this point in the original space.

The field of manifold learning is evolving very quickly and numerous methods have been proposed to address the problem. Two recent surveys [13, 68] present current tendencies of the field. Among popular methods, we can mention Isomap [75], Curvilinear Distances Analysis (CDA [52]), Locally Linear Embedding (LLE [65, 67]) and Laplacian Eigenmaps [8].

### 3.3 Latent variable models

In latent variable models, the high dimensional observed data  $t_1, \dots, t_n$  in  $\mathbb{R}^p$  are supposed to be generated from corresponding low dimensional unobserved (or latent) data  $x_1, \dots, x_n$  in  $\mathbb{R}^q$  with  $q < p$  via the general formulation  $t = y(x; W) + \epsilon$ , where  $y$  is a function of the latent variables  $x$  and of some parameters  $W$ , and where  $\epsilon$  represents some noise.

The simplest and oldest latent variable model is the one of factor analysis (see e.g. [13]). In this model,  $y$  is a linear function of both  $x$  and  $W$ :  $t = Wx + \mu + \epsilon$  (where  $\mu$  is the expectation of  $t$ ). Moreover, we assume that  $x$  has Gaussian distribution (with identity covariance matrix) and  $\epsilon$  is also Gaussian with covariance matrix  $\Psi$ . The parameters of the model can be estimated via an

EM algorithm. In practice however, additional constraints are used to simplify the estimation, for instance by assuming that  $\Psi$  is known. Additional (strong) assumptions allows one to show that PCA is a special case of factor analysis.

Factor analysis is revisited in [78] to produce a probabilistic PCA (PPCA) model for which a closed-form solution is given. The main advantage of this model over PCA is its tolerance to missing data, but it can also be extended to a mixture of PPCA models [77]: data are assumed to be generated by a mixture of local factor models. This provides multiple local linear projections of the same data set. A related hierarchical latent model has also been proposed [12]. Further development of similar models can be found in [85] (see also section 5.1.1).

### 3.4 Limitations of dimension reduction for infovis

An important limitation of advanced dimension reduction methods, apart from their computational cost, is that many of them don't always produce interesting visualization. As pointed out in e.g. [13], Kernel PCA for instance, is more adapted to feature extraction than to dimension reduction as PCA is conducted in the high dimensional kernel induced feature space. Manifold learning is also intrinsically limited by the fact that low dimensional manifolds cannot in general be projected to two dimensions without introducing distortions. Earth maps are well known examples of this problem. A possible solution can be found in a recent tearing method proposed in [53].

Finally, the main problem is to evaluate the impact of the dimension reduction on the ability to conduct data mining tasks: if the neighborhood relationship between objects are not correctly preserved by the reduction methods, for instance, the corresponding visualization can lead to false conclusion because close objects can be mapped to distant points and *vice versa* and can therefore produce visualizations that lead to false conclusion. Linear projection methods always reduce distances between points and can therefore project outliers close to the bulk of the other data, for instance. Non linear methods introduce more complex distortions which mix compression and stretching. Moreover, as pointed out in [70], some manifold learning methods, such as LLE, don't even explicitly preserve distances.

The global distortion of a projection method can be assessed with neighborhood preservation measures [82, 42]. These measures allow the user to decide whether she should trust the projection or not. Detailed and local analysis of the distortion can be done with the visualization methods proposed in [4, 5].

## 4 Reducing the number of objects

Limiting the number of objects to display is extremely important for most of the visualization methods. Clustering algorithms have been used for this task in order to provide scalability to some visualization methods, especially those, such as parallel coordinates, that are impaired by superposition. Objects are



then replaced by a representative object (a prototype) chosen in the cluster to which they belong (see [87]).

It should be noted that this type of automatic data reduction techniques have been used in infovis only quite recently, e.g. in [63, 27]. However, rather than simply relying on prototypes produced by clustering algorithms, summarized displays include visual representation of the clusters themselves. The method proposed in [27] (and latter generalized in [28]) displays both prototypes and the variability in clusters, using some color coding. Moreover, as the simplification is based on hierarchical clustering, the user can interactively choose the amount of simplification.

## 5 Specialized models

### 5.1 Self-Organizing Map

The great success of the Self-Organizing Map (SOM [46]) as a visualization tool might be a consequence of the simultaneous simplifications of a data set that it operates: it acts both as a clustering algorithm and as a non linear projection method. Moreover, the grid structure solves nicely the superposition problems associated to algorithms surveyed in section 3: complex representation of the prototypes can be used without superposition. The basic SOM has been completed by many visualization enhancements ranging from component planes to the U-matrix [81]. Surveys of visualization methods based on SOM can be found in [83, 84, 34].

Extensions to the SOM have been designed based on some important discoveries of the infovis community (see in particular [34]). The “focus+context” concept for instance has motivated the introduction of hyperbolic SOM [64] and more recently of hierarchically growing hyperbolic SOM [57] (see also [1] for a combination of neural gas [56] with the “focus+context” principle).

Other recent works on visualization methods for the SOM include graph based approaches [61], P-Matrix [80], connectivity matrix [74], etc.

#### 5.1.1 Generative Topographic Mapping

In addition to the latent models surveyed in 3.3, which operate only a (local) dimension reduction, a very interesting non linear model, the Generative Topographic Mapping (GTM) has been proposed [11]. As the SOM, this model can be seen as doing both a dimension reduction and some form of clustering. The model is based as the SOM on a grid of points  $(x_i)_{1 \leq i \leq k}$  chosen in the low dimensional latent space. A set of  $m$  non linear functions  $(\phi_j)_{1 \leq j \leq m}$  is used to map the latent space to  $\mathbb{R}^m$  (the mapping is denoted  $\Phi$ ). The observed data  $t$  are assumed to be distributed as a mixture of  $k$  Gaussians with a common covariance matrix  $\beta^{-1}\mathbf{1}$  and with centers given by  $(W\Phi(x_i))_{1 \leq i \leq k}$ , where  $W$  is a  $p \times m$  parameter matrix. An EM algorithm is used to fit  $W$  and  $\beta$  to the data.

The application of GTM to visualization is based on the fact that each observation  $t_i$  induces a posterior distribution in the latent space. This distribution

can be summarized by its mean or by its mode, and provides this way a non linear projection of each observation to the latent space. An important difference with the SOM is that when the mean is used to represent an observation, its position is not constrained in the grid of points and the projection is therefore smooth. If the mode is considered, then the visualization is quite similar to what can be obtained with the SOM.

The initial GTM model has been modified and adapted in many ways (see [10]). For the visualization aspect, interesting developments include the visualization of the distortion (magnification factors) [9] and a hierarchical GTM model proposed in [76], which is an extension of the hierarchical local linear projection developed in [12]. Variations of the GTM can be used to identify outliers [58].

## 6 Conclusion

Integrating machine learning and information visualization is potentially rewarding, as demonstrated by successful visual data mining tools such as the Self-Organizing Map and the Generative Topographic Mapping. A lot of integration work remains however to be done in order to benefit from advanced results of both domains. User control and interaction, aesthetic layout, linking-and-brushing remain for instance quite rare in machine learning oriented programs, whereas advanced dimension reduction and clustering methods are seldom use in infovis. As pointed out in [15], tightening the bounds between machine learning and information visualization is one of the challenge in which both community might find very rewarding results.

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