Self-Organizing Maps for Exploration of Partially Observed Data and Imputation of Missing Values

Sara Rejeb\textsuperscript{a,b,*}, Catherine Duveau\textsuperscript{b}, Tabea Rebafka\textsuperscript{a}

\textsuperscript{a}LPSM, Sorbonne Université, Université de Paris \& CNRS, 4, Place Jussieu, 75252, Paris cedex 05, France
\textsuperscript{b}Safran Aircraft Engines, Réau, 77550 Moissy-Cramayel, France

Abstract

The self-organizing map is an unsupervised neural network which is widely used for data visualisation and clustering in the field of chemometrics. The classical Kohonen algorithm that computes self-organizing maps is suitable only for complete data without any missing values. However, in many applications, partially observed data are the norm. In this paper, we propose an extension of self-organizing maps to incomplete data via a new criterion that also defines estimators of the missing values. In addition, an adaptation of the Kohonen algorithm, named missSOM, is provided to compute these self-organizing maps and impute missing values. An efficient implementation is provided. Numerical experiments on simulated data and a chemical dataset illustrate the short computing time of missSOM and assess its performance regarding various criteria and in comparison to the state of the art.

Keywords: Self-organizing maps, partially observed data, missing data imputation, robustness to missingness mechanism

1. Introduction

In chemometrics, data exploration is essential and is generally the first part of the analysis of any dataset. With the explosion of data volume in many fields of application due to big data, the structure of data is often hidden behind the massive data and difficult to detect. This increases the importance of data exploration methods providing meaningful information on the internal structure and correctness of the data, as well as the relationships and redundancies among the variables. The
initial understanding of the data gained by exploratory data analysis is particularly useful for data modeling.

Common tasks of data exploration are visualization and clustering of the data. While there is plethora of methods addressing one of the tasks, self-organizing maps simultaneously provide both a low-dimensional visual data representation in form of a map and a clustering of the observations. Introduced by [1], this approach consists in mapping the data and performing vector quantization of the input space while preserving topological properties of the data even when these data are high-dimensional. These are properties that other data analysis methods do not have. For instance, $k$-means only categorises data, but the topology gets lost and the method does not allow data visualisation. Several examples of application in different areas are provided in [2] illustrating the practical interest of self-organizing maps. In [3] it is shown that self-organizing maps are a powerful tool for visualisation of high-dimensional data. A fraud detection method based on the SOM visualization and classification is proposed in [4]. In [5], self-organizing maps are used to visualize and classify complex geologic data. Moreover, self-organizing maps have proven to be of considerable value in finance, where they are able to structure, analyze, and visualize large amounts of multidimensional financial data in a significant manner [6, 7]. In short, as self-organizing maps provide easily interpretable results with a global view of the data, they have become very popular in many fields of application. Furthermore, self-organizing maps are widely used in chemometrics [8, 9, 10, 11, 12, 13], but also in biology [14], humanities [15], industry, such as health monitoring of aircraft engines [16]. Many variants of the standard self-organizing map have been developed, such as Generative Topographic Mapping [17], which is a probabilistic version of the self-organizing map, or extensions to more complex data types (mixed, textual, etc.) [18, 19] demonstrating the relevance of self-organizing maps until today.

A common issue with datasets in most fields of application are missing data. Data may be incomplete for a large variety of reasons. In surveys, for instance, they occur due to non-responses to questions that affect privacy [20, 21, 22]. In industrial applications and chemometrics, measuring instruments may have malfunctions or detection limits yielding erroneous and missing entries [23, 24]. In medical research, missing data can occur in clinical trials when patients abandon or stop taking the treatment for a certain period of time [25, 26, 27]. Missing data also frequently occur in chemical [28, 29] and environmental [30] studies. Moreover, concerning huge databases, merging several datasets from different sources can also result in missing data, as some entries may not be
recorded at all for some of the sources.

The impact of missing data on statistical results can be serious, leading to biased estimates, loss of information, decreased statistical power, increased standard errors, and weakened generalizability of findings. However, for a long time, in statistics, missing data have been treated in a very simple and inappropriate way, either by deletion of incomplete measurements or by basic data completion by mean or median values. This has changed during the last decades, by the development of many statistical methods that account for missing data in a meaningful way. There are two general approaches to deal with missing data: either a statistical method is directly adapted to the partially observed data, or first an appropriate imputation method is applied to complete the data such that the statistical method of interest can be used on the completed or augmented data.

In this paper we are interested in self-organizing maps in the presence of missing data. This problem has been considered among others by [31] by simply restricting all vector calculations to the observed entries. As such, all observed data entries are taken into account in the algorithm. However, the method performs rather poorly when the number of incomplete observations with multiple missing entries is large. Moreover, the imputation of missing data is done afterwards by replacing missing entries by the closest features on the learned map. Related approaches are presented in [32, 33, 34, 35, 36] and more recently in [37, 38].

Missing data imputation can be useful to avoid incomplete data, which is crucial in data mining when methods cannot handle any missing entries. We have the ambition to combine the tasks of imputation and learning the map by a principled approach. Our motivation is the fact that any non trivial imputation method is based on some data model, and so it is natural to use the self-organizing map for imputation. Conversely, a better map may be learned when data are complete. Thus, treating both tasks simultaneously may be beneficial for the two of them.

Our approach can be viewed as an extension of the standard Kohonen algorithm for self-organizing maps and the principle of our method is given in Section 2. A mathematical presentation of the method, a new loss function that encodes our double goal of imputation and learning a self-organizing map and two algorithmic solutions are given in Section 3. Moreover, Section 4 provides an extensive numerical study assessing the robust performance of our method in various settings and in comparison to alternative methods from the literature.
2. The new method in a nutshell

This section presents the principle of our new method for self-organizing maps for partially observed data. To start with, we recall the classical self-organizing map. Let $x_1, \ldots, x_n$ be $n$ observations or measurements of dimension $p$. A self-organizing map represents a nonlinear projection of the high-dimensional data onto a low-dimensional subspace. This subspace is a two-dimensional map represented as a regular grid composed of $K$ fixed neurons. This fixed spatial arrangement of the neurons on the map is the key for the preservation of the topology of the input data when projected onto the map. Every neuron $k$ is associated with a $p$-dimensional prototype vector $w_k$, also called code vector, that is to be learned. The prototype vectors define a discretization of the data space, and each observation $x_i$ is assigned to its closest prototype. Ideally, prototype vectors of neighboring neurons on the grid are close one to another, so that data points $x_i$ that are close in the input space are also close on the map.

The Kohonen algorithm computes the self-organizing map for complete data in an iterative fashion. One randomly picked observation $x_i$ is treated at each iteration. First, the winning neuron or best matching unit is determined, which is the neuron whose prototype is the closest to measurement $x_i$. Then, all code vectors $w_k$ are updated by attracting them towards the measurement $x_i$. The attraction is the strongest for the winning neuron and very weak for the neurons that are far from the winning neuron. Those updates eventually result in an ordered map, where neighboring neurons have similar prototype vectors.

Now, when some of the measurement vectors $x_i$ contain missing entries, the Kohonen algorithm is not applicable anymore. We propose to learn the missing entries while learning the map in the following way. Our algorithm is the following: starting from some initial imputed values, like the mean values computed over the observed entries, select every measurement $x_i$ once, determine its winning neuron by considering the distance only over the observed entries of $x_i$ and then update the code vectors just as in the classical Kohonen algorithm. Then, perform an update of the imputed values using a weighted means of the closest code vectors of the partially observed measurement. Repeat this procedure until convergence. A full description is provided in Algorithm 3.

This algorithm performs both data visualization and imputation of the incomplete data. Interestingly, it is as fast as the standard Kohonen algorithm, since the update of the imputed values is immediate.
Algorithm 1: Standard Kohonen algorithm

**Input:** Data matrix $X$, size and topology of the map, neighborhood function $V_\lambda$, sequence of radii $(\lambda_t)_{0 \leq t \leq T}$ and learning steps $(\varepsilon_t)_{0 \leq t \leq T}$.

Initialize code vectors $W^{(0)}$;

Initialize the counter of iterations: $t = 0$;

**while not converged** do

Increment $t$: Set $t = t + 1$;

Choose an observation $i \in \{1, \ldots, n\}$ randomly;

Assignment: Compute winning neuron $\ell = h(x_i, W^{(t-1)})$;

Update code vectors:

for $k = 1, \ldots, K$ do

$$w_k^{(t)} = w_k^{(t-1)} + \varepsilon_t V_\lambda(k, \ell) \left( x_i - w_k^{(t-1)} \right),$$

end

end

**Output:** Code vectors $W^{(t)}$.

3. Self-organizing maps with incomplete data

In this section we first formally state the classical self-organizing map, before introducing the new loss function and two algorithms for the computation of self-organizing maps with partially observed data and missing data imputation.

3.1. Classical Kohonen algorithm

The data matrix containing the measurements is denoted by $X = [x_1, \ldots, x_n] \in \mathbb{R}^{n \times p}$. The arrangement of the neurons on the map is given by some neighborhood function $V_\lambda : \{1, \ldots, K\}^2 \rightarrow \mathbb{R}_+$. The neighborhood radius $\lambda > 0$ describes the zone of influence around a neuron. The best prototype vectors of the self-organizing map are defined as the minimum of the loss function $F$ defined by

$$F(W) = \frac{1}{2n} \sum_{i=1}^{n} \sum_{k=1}^{K} V_\lambda(k, h(x_i, W))\|x_i - w_k\|_2^2,$$  

(1)

where $W = [w_1, \ldots, w_K] \in \mathbb{R}^{p \times K}$ is the matrix of $K$ prototype vectors and $h : \mathbb{R}^p \times \mathbb{R}^{p \times K} \rightarrow \{1, \ldots, K\}$ denotes the allocation function, attributing the closest prototype to a data point $x$. 


w.r.t. the Euclidean distance, defined as

\[ h(x, W) = \arg \min_{1 \leq k \leq K} \|x - w_k\|_2. \]  

(2)

The loss \( F \) takes into account all distances between every measurement and all code vectors, weighted by the neighborhood function evaluated on the corresponding neurons. As a result, code vectors that minimize the loss are similar if they are close on the map. In the specific case where the neighborhood function satisfies \( V_\lambda(k, \ell) = 0 \) for all \( k \neq \ell \), the loss \( F \) is the criterion minimized by the \( k \)-means algorithm. That is, clusters obtained by \( k \)-means are independent, while prototypes of a self-organizing map are organized in a topological way.

To compute the minimum of loss \( F \), [39] showed that in the given framework a gradient descent algorithm can be used, referred to as the Kohonen stochastic algorithm. For a randomly picked observation \( x_i \) with winning neuron \( \ell = h(x_i, W^{(t)}) \), the updates of the code vectors are given by

\[ w_k^{(t+1)} = w_k^{(t)} + \varepsilon_t V_\lambda(k, \ell)(x_i - w_k^{(t)}), \]  

(3)

where \( (\varepsilon_t)_{t \geq 0} \) is a sequence of decreasing learning steps. This update attracts all prototypes towards observation \( x_i \). It is also common to shrink the neighborhood by using a decreasing sequence of radii \( (\lambda_t)_{t \geq 0} \) in the neighborhood function \( V_\lambda \). The algorithm is summarized in Algorithm 1.

3.2. Notation for incomplete data

Now we consider an incomplete \( n \times p \) data matrix containing missing values. Let the matrix \( M = (m_{i,j})_{i,j} \in \{0,1\}^{n \times p} \) be the missing-data pattern which indicates where the entries are missing or masked, and that is defined by

\[ m_{i,j} = \begin{cases} 
1 & \text{if } x_{i,j} \text{ is observed} \\
0 & \text{if } x_{i,j} \text{ is missing}
\end{cases} \]

We denote \( X^{\text{obs}} \) the set of observed data values and \( X^{\text{miss}} \) the set of non-observed data entries hidden by the missing-data pattern \( M \). The complete data are denoted by \( X^{\text{compl}} = (X^{\text{obs}}, X^{\text{miss}}) \). Likewise, for the observation vector \( x_i \) we denote by \( x_i^{\text{obs}} \) and \( x_i^{\text{miss}} \) the observed and unobserved entries, respectively, and, with some abuse of notation, \( x_i^{\text{compl}} = (x_i^{\text{obs}}, x_i^{\text{miss}}) \) is the complete vector, which also corresponds to the \( i \)-th row of \( X^{\text{compl}} \).

Our goal is to adapt the model of self-organizing maps to partially observed data, and moreover, learn the values of the missing data. The motivation to treat these tasks simultaneously is that
learning missing values requires a data model, and as we are interested in self-organizing maps it is
natural to use this model for data imputation. At the same time, learning a map with completed
data may give better results compared to using only the observed part \(X^{\text{obs}}\) of the data.

3.3. New loss function

We introduce a new loss function that considers both problems: finding the best self-organizing
map and the best values for imputation of the missing data. In other words, by minimizing the new
loss function \(F_{\text{missom}}\) we search for both the best code vectors \(W \in \mathbb{R}^{p \times K}\) for the map and the best
values for the missing data denoted by \(X^*\) chosen in the set of all possible values for the missing
entries \(X^{\text{miss}}\).

To define the new criterion, an adaptation of the definition of the winning neuron is in order.
In the presence of missing values, it is natural to restrict the Euclidean distance in (2) only to the
observed entries. More precisely, for any vectors \(x^{\text{obs}} \in \mathbb{R}^{p'}\) \((p' \leq p)\), \(m \in \{0, 1\}^p\) with \(\sum_{j=1}^{p} m_j = p'\)
and code vectors \(W \in \mathbb{R}^{p \times K}\), we set

\[
h^{\text{miss}}(x^{\text{obs}}, m, W) = \arg \min_{1 \leq k \leq K} \|x^{\text{obs}} - w_k \odot m\|_2,
\]

where \(w_k \odot m\) denotes the \(p'\)-vector made of the elements \(w_{k,j}\) of \(w_k\) such that \(m_j = 1\). Now, we
define the new loss as

\[
F_{\text{missom}}(W, X^*) = \frac{1}{2n} \sum_{i=1}^{n} \sum_{k=1}^{K} V_{\lambda}(k, h^{\text{miss}}(x^{\text{obs}}_i, m_i, W)) \|x^{\text{obs}}_i - w_k\|_2^2,
\]

where \(m_i \in \mathbb{R}^p\) is the \(i\)-th row of the matrix \(M\) and \((x^{\text{obs}}_i, x^*_i)\) denotes the \(i\)-th measurement
vector completed with \(x^*_i\). Since

\[
\|(x^{\text{obs}}_i, x^*_i) - w_k\|_2^2 = \|x^{\text{obs}}_i - w_k \odot m_i\|_2^2 + \|x^*_i - w_k \odot (1_p - m_i)\|_2^2,
\]

where \(1_p = (1, \ldots, 1)^T \in \mathbb{R}^p\), the criterion \(F_{\text{missom}}\) can be decomposed into two parts according
to the observed and the missing entries as

\[
F_{\text{missom}}(W, X^*) = F_{\text{obs}}(W) + F_{\text{miss}}(W, X^*),
\]

where \(F_{\text{obs}}(W)\) is the part of the loss over the observed entries given by

\[
F_{\text{obs}}(W) = \frac{1}{2n} \sum_{i=1}^{n} \sum_{k=1}^{K} V_{\lambda}(k, h^{\text{miss}}(x^{\text{obs}}_i, m_i, W)) \|x^{\text{obs}}_i - w_k \odot m_i\|_2^2,
\]
Algorithm 2: missSOM algorithm

Input: Incomplete data $X^\text{obs}$, missing-data pattern $M$, size and topology of the map, neighborhood function $V$, sequence of radii $(\lambda_t)_{0 \leq t \leq T}$ and learning steps $(\varepsilon_t)_{0 \leq t \leq T}$.

Initialize imputed values $X^*(0)$ and code vectors $W^{(0)}$;
Initialize the counter of iterations: $s = 0$;

while not converged do

Increment $s$: Set $s = s + 1$;

Update code vectors by Kohonen Algorithm 1 on the augmented data with winning neurons obtained by $h_{\text{miss}}$ instead of $h$:

$W^{(s)} \leftarrow \text{Kohonen}(X^\text{aug} = (X^\text{obs}, X^{*(s-1)}))$;

Update imputed values: for $i, j$ such that $m_{i,j} = 0$,

$$x_{i,j}^{*(s)} = \frac{\sum_{k=1}^{K} V_{\lambda T}(k, h_{\text{miss}}(x^\text{obs}_i, m_i, W^{(s)})) w_{k,j}^{(s)}}{\sum_{k=1}^{K} V_{\lambda T}(k, h_{\text{miss}}(x^\text{obs}_i, m_i, W^{(s)}))}.$$ 

end

Output: Code vectors $W^{(s)}$ and imputed data $X^{*(s)}$.

and $F_{\text{miss}}(W, X^*)$ is the contribution of the imputed values $X^*$ to the loss, defined as

$$F_{\text{miss}}(W, X^*) = \frac{1}{2n} \sum_{i=1}^{n} \sum_{k=1}^{K} V_{\lambda}(k, h_{\text{miss}}(x^\text{obs}_i, m_i, W)) \|x_{i}^* - w_k \odot (1_p - m_i)\|^2.$$ 

Note that in the complete-data case, where the missing-data pattern is $M = 1_{n \times p}$, $F_{\text{miss}}(W, X^*) = 0$ for any $W$ and any $X^*$, so that the criterion $F_{\text{missom}}$ is equal to the one of the classical self-organizing map, that is, $F_{\text{missom}}(W, X^*) = F(W)$.

3.4. Minimization algorithm

For the minimization of $(W, X^*) \mapsto F_{\text{miss}}(W, X^*)$ on $\mathbb{R}^{K \times p} \times X^{\text{missing}}$ we propose to alternate the minimization in $W$ and $X^*$ while keeping the other argument fixed.

For fixed $X^*$, the function $W \mapsto F_{\text{missom}}(W, X^*)$ is similar to the objective function $F$ in (1) in the complete-data case applied to the augmented data $X^\text{aug} = (X^\text{obs}, X^*)$. The only difference lies in the definition of the winning neurons by $h_{\text{miss}}$ that appear in the neighbourhood function $V$. Thus, a Kohonen algorithm applied to $X^\text{aug}$ can be used to find the best code vectors $W$. 

8
In turn, when $W$ is fixed, the minimization of $X^* \mapsto F_{\text{miss}}(W, X^*)$ boils down to minimize $X^* \mapsto F_{\text{miss}}(W, X^*)$. This problem has a unique explicit solution given for all $1 \leq i \leq n$ and $j$ such that $m_{i,j} = 0$ by

$$x_{i,j}^* = \frac{\sum_{k=1}^{K} V_\lambda(k, h_{\text{miss}}(x_{i}^{\text{obs}}, m_{i}, W))w_{k,j}}{\sum_{k=1}^{K} V_\lambda(k, h_{\text{miss}}(x_{i}^{\text{obs}}, m_{i}, W))}. \quad (4)$$

That is, the imputed values are a weighted mean of the prototype vectors weighted according to the neighborhood function.

To summarize, the algorithm updates imputed values for the missing data and applies the classical Kohonen algorithm with adjusted winning neuron function $h_{\text{miss}}$ to learn the map. This is repeated until convergence or until a maximum number of iterations chosen by the user is attained. The algorithm is described in Algorithm 2.

As initial values for the imputed values $X^*$, one can simply impute the sample mean or median of the variables obtained over the observed entries.

3.5. Accelerated version

Algorithm 2 happens to be expensive in terms of computing time, in particular when the number of iterations is large, since the entire standard Kohonen algorithm is carried out during each iteration. A speed up is obtained by interwining updates of the missing data and the iterations of the Kohonen Algorithm 1. More precisely, we propose to update the missing data at every epoch, that is, after every pass through the data. This procedure gives rise to Algorithm 3. As such, the Kohonen algorithm is carried out only once, while in the initial Algorithm 2, the entire Kohonen algorithm is applied repeatedly. Thus the computing time of the accelerated version of missSOM is comparable to the computing time of the standard Kohonen Algorithm 1, since the update of the imputed values is fast.

While the first version of the missSOM algorithm has some theoretical justification, the accelerated version lacks this foundation. A numerical study given in Appendix 6.1 shows that the Algorithms 2 and 3 provide very similar maps and hence justifies the utilization of the accelerated version, which achieves a significant gain in computing time.

Note that while the selection of the observations $x_i$ in Algorithm 3 is deterministic, it is possible to use a random selection scheme.
4. Numerical experiments

In this section, the performance of the proposed method missSOM is evaluated and compared to alternative methods. A simulation study is conducted to assess the quality of the representation of the data via the map and the accuracy of imputed values under various conditions.

4.1. Performance criteria

Let $W^*$ be the code vectors of the final self-organizing map, $X^*$ the imputed values and $\hat{x}_i = (x_i^{\text{obs}}, x_i^*)$ the completed observation vectors.

The quality of the map as a representation of the data can be evaluated by two criteria. First, the quantization error defined as the average of the squared distances between the observations and their nearest prototype vector given by

$$E = \frac{1}{n_{\text{obs}}} \sum_{i=1}^{n} \| x_i^{\text{obs}} - w_{h^{\text{miss}}(x_i^{\text{obs}}, m_i, W^*)} \|_2^2,$$

where $n_{\text{obs}} = \sum_{i,j} m_{i,j}$ is the number of observed entries in the data, informs on whether the prototype vectors are good representations of the data. Second, the topographic error evaluates the preservation of the topology of the data in the map by the proportion of observations for which the winning neuron and the second closest neuron are not neighbors, i.e. not connected on the grid. It is defined as

$$T = \frac{1}{n} \sum_{i=1}^{n} e(\hat{x}_i),$$

where

$$e(\hat{x}_i) = \begin{cases} 0 & \text{if the neurons } h^{\text{miss}}(x_i^{\text{obs}}, m_i, W^*) \text{ and } v^{\text{miss}}(x_i^{\text{obs}}, m_i, W^*) \text{ are adjacent} \\ 1 & \text{otherwise} \end{cases}$$

with $v^{\text{miss}}(x_i^{\text{obs}}, m_i, W^*) = \arg \min_{j \neq h^{\text{miss}}(x_i^{\text{obs}}, m_i, W^*)} \| x_i^{\text{obs}} - w_j^* \odot m_i \|_2^2$ the second closest neuron to measurement $x_i^{\text{obs}}$. The topographic error is expected to be small when the map is well organized and ordered. As both errors are computed only on the observed entries, the focus is put on the quality of the map with respect to the observed part of the data. The accuracy of the imputed values is assessed separately by the imputation error, which is defined as the root mean square error and quantifies the quality of the imputed values compared to the true missing values.
4.2. Datasets

For the numerical experiments, two settings are considered. First, the dataset *wines* from the UCI machine learning repository [40] is used, which contains the results of a chemical analysis of 178 wines on 13 quantitative variables. We generate 100 perturbed datasets by adding gaussian noise with mean 0 and standard deviation equal to one tenth of the mean value in each variable.

In the second setting, 100 datasets are simulated from a multivariate gaussian mixture with dimension $p = 5$, four equal-sized groups and $n = 2000$ observations. The correlation among all pairs of components is equal to 0.5 and for every dataset, the 4 gaussian means are drawn independently from a centered normal distribution with standard deviation equal to 5.

In both settings, missing values are generated using the *ampute* function from the R package *mice* [41]. Different proportions of missing values, namely 5%, 20% and 40%, and different mechanisms of missingness are considered. The literature on missing data traditionally distinguishes three mechanisms that lead to missingness [42] and it is well known that the performance of imputation methods may be sensitive to the mechanism at work. The mechanism is said to be *missing completely at random* (MCAR), when the causes of missing values are independent from the data and the probability of being absent is the same for all items. That is, a subset of observations is chosen at random using independent Bernoulli variables with fixed success probability for all entries. In contrast, when the probability that a value is missing depends on the values of the observed variables, the mechanism is called *missing at random* (MAR). In our simulations, to obtain MAR, for each variable, missing values are obtained using a logistic regression model depending only on the other variables. Finally, when the probability of being absent also depends on the unobserved value, the mechanism is called *missing not at random* (MNAR) and we can consider missing data simulated by using a logistic regression model depending on all variables.

4.3. Alternative methods

Our study is twofold. On the one hand, we compare *missSOM* to the state of the art on self-organizing maps. The simplest method (referred to as *deletion* in the figures) consists in deleting the observations containing missing values and applying the standard Kohonen algorithm to the remaining data. Incomplete observations are classified once the map is built by assigning them to their closest prototypes and missing entries are imputed by the corresponding values of the winning prototypes. Cottrell’s approach is a variant of the classical self-organizing maps appropriate to deal
with incomplete observations [31]. Cottrell adapts the Kohonen algorithm by simply restricting all vector calculations to the observed entries. The main difference with missSOM is that data imputation is performed only after learning the map by imputation with the values of the closest prototypes.

On the other hand, in our experiments, we compare missSOM to the state of the art on missing data imputation. The literature provides numerous general imputation methods. A basic approach (here referred to as mean) is the imputation by the mean value of the observed variables. A non-parametric approach called missForest predicts missing values using a random forest trained on the observed part of the dataset. Moreover, the method referred to as knn is a $k$-nearest neighbors approach, which is implemented in the VIM package [43]. Missing values are imputed iteratively by a weighted average of the $k$ closest observations. Finally, assuming a gaussian mixture model for the data, the Amelia package [44] performs imputation using the expectation-maximization algorithm and a bootstrap approach to iteratively estimate missing values. In our numerical study, the imputation error of missSOM is compared to the one obtained by all these methods. In addition, we also compare the map obtained by missSOM with the maps obtained by the classical Kohonen algorithm applied to the complete dataset, where missing values are imputed by the aforementioned imputation methods.

As a benchmark we consider a self-organizing map trained on the complete data set (corresponding to a missing rate of 0%) and compute the error rates only on the observed values that are provided to the other methods. This method is referred to as complete-data SOM.

4.4. Parameters of the methods

The missSOM algorithm is applied with the default parameters in the package missSOM. That is, the neighborhood function is a gaussian and the sequence $(\lambda_t)_{t \leq T}$ decreases from $\lambda_0 = 4.58$ to $\lambda_T = 0.5$ for the wine dataset and from $\lambda_0 = 8.89$ to $\lambda_T = 0.5$ for the gaussian mixture data. The maximum number of epochs is $T = 100$. Concerning the map, it has a hexagonal topology and the grid is composed of $K = 9 \times 7$ neurons for the wines data and $K = 16 \times 14$ neurons for the gaussian mixture data. The choice of the size of the map depends on the sample size and on the objectif of the analysis. In general, missSOM has the same parameters as classical SOM and they can be chosen in the same way as for SOM.

For the alternative methods, the maximum number of iterations is set to 100 and for the other
parameters the default values are used.

4.5. Comparison to the state of the art on self-organizing maps

Figure 1 and 2 present the boxplots of the topographic and the quantization error for all SOM-type methods in both settings. They serve to evaluate the quality of the representation of the data using a self-organizing map computed with the different approaches. The corresponding imputation errors in Figure 3 allow to judge their performances as imputation methods.

First, we observe that when the percentage of missing data is low, most methods have very similar performance. With increasing percentage of missingness, the problem becomes harder and differences among the methods appear. But as errors are evaluated only on the observed part of the data, errors are not necessarily increasing. Namely quantization errors for the wines data decrease with increasing missingness. This may be a consequence of the higher variance and less structure of the wines data compared to the Gaussian mixture data. So when deleting entries from the wines data, the data variance decreases and the resulting map is a better representation of the observed part of the data. Concerning the mechanisms of missingness, it appears that the quality of the map does not depend on it for any SOM-type method. However, imputation is impacted by the mechanism. Imputation is the easiest under MCAR and the hardest under MNAR.

Next, we see that the deletion method is very unstable. In some scenarios its error rates are among the worst, and, more importantly, when too many values are missing, the method breaks down and does not produce any result. Indeed, on small datasets as wines with 20% of missingness, the number of complete observations is smaller than the size of the map and thus classical SOM is just not applicable. On the large gaussian mixture dataset, the deletion method produces the best topographic error, but the associated quantization error is disastrous, disqualifying the approach.

The quantization error of the complete-data SOM method is constant when varying the amount of missingness, because the underlying map remains the same. For the topographic error, an increase is observed which is due to the definition of the error. The error determines the closest and second closest prototypes only with respect to the observed entries, while the map was optimized by taking into account the complete data. This explains why missSOM and Cottrell have lower topographic errors, as their maps are learned with a notion of closeness restricted on the observed entries.

Finally, we observe that Cottrell's method always achieves the best quantization errors, directly followed by missSOM. Concerning the topographic error, missSOM is consistently doing better.
Figure 1: Topographic error of SOM-type methods for various amounts and mechanisms of missingness on the wines (a) and the gaussian mixture data (b).

than Cottrell. Thus, in terms of quality of the map and representation of the data, none of the methods outperforms all others, and Cottrell and missSOM have both a good global performance. Now, considering the imputation error, there is a clear winner. When the proportion of missingness is important, missSOM outperforms all other methods in every scenario. This confirms us in the use of missSOM with respect to Cottrell, especially when accurate imputation is desired.

4.6. Comparison to the state of the art on missing-data imputation

In the second part of the simulation study, missSOM is compared to general imputation methods. Figure 4 shows that as an imputation method missForest is unbeaten regardless of the missingness mechanism and the percentage of missing data. On gaussian mixture data, knn also provides accurate imputations. The imputation accuracy achieved by missSOM is high and missSOM is mostly among the best imputation methods.

Finally, Figure 5 and Figure 6 provide insights on the quality of the data representation, when classical SOM is applied to the data with missing values imputed by the different imputation methods. In this respect, missSOM clearly outperforms all the other methods. In particular, missForest provides self-organizing maps of poorer quality, especially in terms of preservation of the topology.
Figure 2: Quantization error of SOM-type methods for various amounts and mechanisms of missingness on the wines (a) and the gaussian mixture data (b).

Figure 3: Imputation error of SOM-type methods for various amounts and mechanisms of missingness on the wines (a) and the gaussian mixture data (b).
Figure 4: Imputation error of missSOM and classical imputation methods for various amounts and mechanisms of missingness on the wines (a) and the gaussian mixture data (b).

Figure 5: Topographic error of missSOM and classical imputation methods for various amounts and mechanisms of missingness on the wines (a) and the gaussian mixture data (b).
of the input data. This confirms the initial motivation of missSOM: in the presence of missing values and when a self-organizing map is required, it is better to learn the map simultaneously with the imputed values than to treat the two tasks separately.

A further advantage of missSOM is its computing time, which is about 10 times shorter than the one of missForest.

To summarize, we have seen that missSOM is the method of choice when both tasks data imputation and data representation are desired. However, when the focus is only on data imputation, then better alternatives as missForest exist.

5. Conclusion

In this paper, we have proposed an extension of the self-organizing map for partially observed data, referred to as missSOM. The proposed method addresses simultaneously the two problems of computing a self-organizing map and imputing missing values. A numerical study assesses the good performance of missSOM regarding various criteria and in comparison to the state of the art. While this paper focuses on the standard Kohonen algorithm, in future work we may address the
transfer of our approach to other existing variants of self-organizing maps on more complex data types such as mixed data to enable them to deal with missing data.

6. Appendix

6.1. Validation of accelerated missSOM algorithm

Table 1 compares the results of the basic missSOM Algorithm 2 and its accelerated version Algorithm 3 on the simulated gaussian mixture data in various conditions. On the one hand, for all settings the errors are totally equivalent. This indicates that the accelerated version provides the same self-organizing maps and very similar imputations as the basic missSOM algorithm. On the other hand, we see that in terms of computing time we gain two orders of magnitude. Hence, the use of the accelerated algorithm instead of the basic version is completely justified.

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Algorithm 3: Accelerated missSOM algorithm

**Input:** Incomplete data matrix $X^{\text{obs}}$, missing-data pattern $M$, size and topology of the map, neighborhood function $V_{\lambda}$, sequence of radii $(\lambda_t)_{0 \leq t \leq T}$ and learning steps $(\varepsilon_t)_{0 \leq t \leq T}$.

Initialize imputed values $X^{*}(0)$ and code vectors $W^{(0)}$.

Initialize the number of epochs: $t = 0$;

**while** not converged **do**

Increment $t$: Set $t = t + 1$;

Set $\tilde{W}^{(0)} = W^{(t-1)}$;

**for** $i = 1, \ldots, n$ **do**

Assignment: Compute winning neuron $\ell = \text{h}^{\text{miss}}(x_i^{\text{obs}}, m_i, \tilde{W}^{(i-1)})$;

Update code vectors:

**for** $k = 1, \ldots, K$ **do**

$$w_k^{(i)} = \tilde{w}_k^{(i-1)} + \varepsilon_t V_{\lambda_i}(k, \ell) \left( (x_i^{\text{obs}}, x_i^{* (t-1)}) - \tilde{w}_k^{(i-1)} \right).$$

**end**

**end**

Set $W^{(t)} = \tilde{W}^{(n)}$;

Update imputed values: for $i, j$ such that $m_{i,j} = 0$,

$$x_{i,j}^{* (t)} = \frac{\sum_{k=1}^{K} V_{\lambda_i}(k, \text{h}^{\text{miss}}(x_i^{\text{obs}}, m_i, W^{(t)})) w_{k,j}^{(t)}}{\sum_{k=1}^{K} V_{\lambda_i}(k, \text{h}^{\text{miss}}(x_i^{\text{obs}}, m_i, W^{(t)}))}. $$

**end**

**Output:** Code vectors $W^{(t)}$ and imputed data $X^{* (t)}$. 
| Error Type       | Technique        | 5% MCAR | MAR | MNAR | 20% MCAR | MAR | MNAR | 40% MCAR | MAR | MNAR |
|------------------|------------------|---------|-----|------|----------|-----|------|----------|-----|------|
| Topographic error| basic missSOM    | 0.337 (0.047) | 0.345 (0.035) | 0.336 (0.037) | 0.313 (0.032) | 0.355 (0.063) | 0.339 (0.046) | 0.294 (0.056) | 0.303 (0.057) | 0.306 (0.083) |
|                  | accelerated missSOM | 0.352 (0.039) | 0.345 (0.049) | 0.338 (0.040) | 0.301 (0.035) | 0.299 (0.039) | 0.322 (0.042) | 0.282 (0.035) | 0.290 (0.059) | 0.285 (0.040) |
| Quantization error| basic missSOM    | 0.405 (0.115) | 0.394 (0.042) | 0.407 (0.143) | 0.327 (0.115) | 0.326 (0.116) | 0.330 (0.112) | 0.225 (0.074) | 0.239 (0.060) | 0.233 (0.077) |
|                  | accelerated missSOM | 0.413 (0.149) | 0.409 (0.139) | 0.407 (0.130) | 0.351 (0.177) | 0.340 (0.116) | 0.341 (0.115) | 0.361 (0.081) | 0.245 (0.071) | 0.260 (0.074) |
| Imputation error  | basic missSOM    | 0.601 (0.140) | 0.598 (0.154) | 0.613 (0.171) | 0.659 (0.125) | 0.686 (0.141) | 0.759 (0.108) | 0.758 (0.090) | 0.795 (0.095) | 0.871 (0.093) |
|                  | accelerated missSOM | 0.615 (0.097) | 0.624 (0.096) | 0.654 (0.103) | 0.685 (0.092) | 0.694 (0.087) | 0.770 (0.075) | 0.734 (0.060) | 0.731 (0.069) | 0.848 (0.072) |
| ARI              | basic missSOM    | 0.922 (0.092) | 0.929 (0.089) | 0.937 (0.071) | 0.849 (0.088) | 0.846 (0.108) | 0.823 (0.089) | 0.486 (0.108) | 0.640 (0.101) | 0.623 (0.086) |
|                  | accelerated missSOM | 0.923 (0.090) | 0.923 (0.086) | 0.922 (0.082) | 0.841 (0.092) | 0.845 (0.081) | 0.805 (0.080) | 0.647 (0.094) | 0.629 (0.109) | 0.598 (0.089) |
| Computing time   | basic missSOM    | 33.169 (0.546) | 32.940 (0.737) | 31.860 (0.396) | 31.076 (0.517) | 32.988 (0.776) | 32.948 (0.530) | 33.031 (0.583) | 31.920 (0.061) | 32.846 (0.0644) |
|                  | accelerated missSOM | 0.394 (0.018) | 0.387 (0.006) | 0.376 (0.009) | 0.557 (0.008) | 0.561 (0.017) | 0.556 (0.006) | 0.790 (0.016) | 0.751 (0.011) | 0.784 (0.022) |

Table 1: Comparison of the basic missSOM Algorithm 2 and its accelerated version Algorithm 3 in terms of different errors and computing time (in seconds) on the gaussian mixture data.