Getting to Know the Unknown Unknowns: Destructive-Noise Resistant Boolean Matrix Factorization

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Abstract

Finding patterns in binary data is a classical problem in data mining, dating back to at least frequent itemset mining. More recently, approaches such as tiling and Boolean matrix factorization (BMF), have been proposed to find sets of patterns that aim to explain the full data well. These methods, however, are not robust against non-trivial destructive noise, i.e. when relatively many 1s are removed from the data: tiling can only model additive noise while BMF assumes approximately equal amounts of additive and destructive noise. Most real-world binary datasets, however, exhibit mostly destructive noise. In presence/absence data, for instance, it is much more common to fail to observe something than it is to observe a spurious presence. To address this problem, we take the recent approach of employing the Minimum Description Length (MDL) principle for BMF and introduce a new algorithm, \texttt{Nassau}, that directly optimizes the description length of the factorization instead of the reconstruction error. In addition, unlike the previous algorithms, it can adjust the factors it has discovered during its search. Empirical evaluation on synthetic data shows that \texttt{Nassau} excels at datasets with high destructive noise levels and its performance on real-world datasets confirms our hypothesis of the high numbers of missing observations in the real-world data.

Keywords: Boolean matrix factorization; minimum description length; unknown unknowns

1 Introduction

As we know, there are known knowns; there are things we know we know. We also know there are known unknowns; that is to say we know there are some things we do not know. But there are also unknown unknowns – the ones we don’t know we don’t know. And ... it is the latter category that tend to be the difficult ones. [30]

While then-Secretary of Defense Donald Rumsfeld probably was not thinking about data mining when he made his famous comment about unknown unknowns, they are, arguably, the main problem data miners dealing with observation data have to face. A known unknown, in this setting, is an element for which we know we do not know its value. As an example, in a movie rating matrix we know which movies the users have not rated yet. In contrast, an unknown unknown is an element we have not observed, but we do not know if this is because it does not exist or because we have failed to observe it.

We assume that many real-world presence/absence (or binary) datasets contain unknown unknowns, and that is it is much more common to fail to observe something that is than it is to observe something that is not there. In other words, we assume there is more destructive noise than there is additive noise -- without having to pre-specify these amounts exactly. In order to find interesting patterns from the data, we consider Boolean matrix factorization (BMF) [20], a technique that has been successfully applied to a variety of data mining problems considering binary data (e.g. [14, 16, 20, 22]). In its standard form, BMF aims to find a low-rank Boolean factorization of the data that is as close to the original as possible (i.e. minimizes the reconstruction error).

But here as well the unknown unknowns are the difficult ones. When we know which observations are unknown, we can ignore them, and fit the model (i.e. the Boolean matrix factorization) only to the observed parts of the data. But we cannot ignore all unobserved values, as that leaves only the observations (1s in the data) to work with. Still, we also have to be careful that we do not consider unobserved values as important to actual observations -- as standard reconstruction error does -- as this ignores our hypothesis that real data has more unobserved than spuriously observed values.

To address this problem, we use the Minimum Description Length principle (MDL) [11, 26], which is very useful in tackling the problem of the trade-off between fitting the data well while keeping the model simple. In general, the more complexity we allow in the model, the better we can fit the data. However, having high model order comes at the cost of fitting the noise. MDL identifies the optimal balance.

In this paper we present \texttt{Nassau}, a new BMF algorithm that is designed to directly minimize description length. A key aspect is that -- unlike the majority of the previously proposed BMF algorithms -- it can correct its previous mistakes. \texttt{Nassau} is quite robust to destructive noise, which is especially beneficial for real-world data as in many domains there are zeros simply due to the lack of observation.

2 Notation

In this paper we consider matrices of Boolean values. We denote a matrix by upper-case boldface letters (A), and vectors by lower-case boldface letters (a). If A is an $n$-by-$m$ Boolean matrix, $|A|$ denotes the number of 1s in it, i.e.,
We use two-part MDL because we are specifically interested (e.g. [14]), the Boolean factors can provide better input for subsequent algorithms (e.g. [1, 31]), and for sparse input data, the factors will be naturally sparse [19]. Unfortunately, computing the least-error BMF is NP-hard, and has strong inapproximability results [18]. But there is also another, more fundamental problem: the formulation of the minimum-error Boolean rank-k problem requires a priori knowledge on k, the Boolean rank of the decomposition. And as discussed in the Introduction, using the Hamming distance essentially assumes roughly equal distribution between destructive and additive noise – something that we hypothesize is not true. In order to solve these two problems, we will use the description length instead of the reconstruction error to measure the quality of our decomposition.

3 BMF with MDL

3.1 MDL: a brief introduction. The Minimum Description Length principle (MDL) [11] is a practical version of Kolmogorov Complexity. Both embrace the slogan Induction by Compression. This can be roughly described as follows: Given a set of models $\mathcal{M}$. The best model $M \in \mathcal{M}$ is the one that minimizes $L(M) + L(D \mid M)$, in which $L(M)$ is the length in bits of the description of $M$, and $L(D \mid M)$ is the length of the data when encoded with model $M$.

This is called two-part, or crude, MDL – as opposed to refined MDL, where model and data are encoded together [11]. We use two-part MDL because we are specifically interested in the model: the factors that give the best description of the data. Note that MDL requires the compression to be lossless in order to allow for fair comparison between different $M \in \mathcal{M}$, and that we are only concerned with code lengths, not actual code words.

3.2 BMF: a brief introduction. In Boolean matrix factorization, the goal is to (approximately) represent a Boolean matrix as the Boolean product of two Boolean matrices. That is, given Boolean matrix $A$, find Boolean matrices $B$ and $C$ such that $A \approx B \circ C$. In minimum-error Boolean rank-k decomposition one is given the n-by-m input matrix $A$ and rank $k$, and the goal is to find n-by-k and k-by-m factor matrices $B$ and $C$ that minimize $|A \oplus (B \circ C)|$.

Using the Boolean decomposition instead of a normal (or non-negative) can give many advantages in data mining; the binary factor matrices are often easier to interpret [18, 20], or the binary factors might be required in the application (e.g. [14]), the Boolean factors can provide better input for subsequent algorithms (e.g. [1, 31]), and for sparse input data, the factors will be naturally sparse [19]. Unfortunately, computing the least-error BMF is NP-hard, and has strong inapproximability results [18]. But there is also another, more fundamental problem: the formulation of the minimum-error Boolean rank-k problem requires a priori knowledge on k, the Boolean rank of the decomposition. And as discussed in the Introduction, using the Hamming distance essentially assumes roughly equal distribution between destructive and additive noise – something that we hypothesize is not true. In order to solve these two problems, we will use the description length instead of the reconstruction error to measure the quality of our decomposition.

3.3 MDL for BMF and problem definition. To use MDL, we have to define what our models $M$ are, how an $M \in \mathcal{M}$ describes a database, and how we encode these in bits. Miettinen and Vreeken [22] proposed a number of MDL objective functions for BMF. Here we use their so-called typed data-to-model encoding, which was shown to be both the most efficient as well as providing the best empirical performance. Below we give the main ideas of this encoding scheme. For further details we refer the reader to [22].

The description length of a Boolean $n$-by-$m$ matrix $A$ factorized into $B$ and $C$, such that $A \approx B \circ C$, is defined as

$$L(A, B, C) = L(B) + L(C) + L(A \mid B, C),$$

where $L(B)$ and $L(C)$ are the description lengths of the factor matrices and $L(A \mid B, C)$ is the description length of the matrix $A$ given this model. The columns of $B$, and analogously the rows of $C$, are encoded independently as binary vectors. One such vector can be identified by two integers: one encoding the number of nonzero elements in $B^i$ (maximum $n$), and the other encoding the index of $B^i$ among all binary strings having the same profile (maximum $\binom{n}{|B^i|}$). The number of bits for encoding the $k$ columns of $B$ is hence

$$L(B) = k \log(n) + \sum_{i=1}^{k} \log \left( \binom{n}{|B^i|} \right).$$

To reconstruct the data $A$ given the factors matrices $B$ and $C$ means we need to describe the error of this factorization. We hence have to encode the exclusive OR of the data with the model, $E = A \oplus (B \circ C)$. We can split $E$ into unmodelled 1s, $E^+$, and superfluous 1s, $E^-$, where $E^+_{ij} = 1$ if and only if $A_{ij} > (B \circ C)_{ij}$ and $E^-_{ij} = 1$ if and only if $A_{ij} < (B \circ C)_{ij}$. To avoid rewarding structure in the error matrices $E^+$ and $E^-$, they are encoded as binary strings in the same way as a column of $B$. Combined, we have $L(A \mid B, C) = L(E^+) + L(E^-)$, where

$$L(E^+) = \log(nn - |B \circ C|) + \log \left( \frac{nn - |B \circ C|}{|B \circ C|} \right),$$

and

$$L(E^-) = \log(|B \circ C|) + \log \left( \frac{|B \circ C|}{|E^-|} \right).$$
This concludes the definition of our MDL objective function. We can now define the problem we aim to solve in this paper.

**Problem 3.1. (MDLBMF)** Given a binary $n$-by-$m$ matrix $A$, the Minimum Description Length Boolean Matrix Factorization (MDLBMF) problem is to find binary factor matrices $B$ and $C$ such that we minimize the total description length

$$L(A, B, C).$$

Note that we do not require the rank of the decomposition as an input; instead we automatically find the rank (and decomposition) that minimizes the description length. Although the computational complexity of the MDLBMF problem is unknown, there are strong reasons to believe it is NP-hard [22]. Hence, we resort to heuristics.

4 Related Work

In BMF one decomposes a binary matrix into the Boolean product of two matrices while minimizing some cost function. Perhaps the intuitively most straightforward objective is to minimize the number of errors, i.e. the Frobenius norm of the residual. The $A_{ss}$ algorithm to solve BMF was proposed by [20]. Later, [13] proposed a heuristic based on a mixed-integer-programming formulation.

Error minimization is prone to overfitting, however, as more factors always allow better reconstruction. In practice, users thus have to choose the number of factors in advance. Moreover, this objective builds on the assumption that noise is equally likely to flip true 1s to 0s as it is to flip true 0s to 1s, which we argue here is not realistic.

As discussed by Faloutsos and Megalooikonomou [6], Kolmogorov Complexity, and its practical implementation, the Minimum Description Length principle [11, 26], are powerful, well-founded, and natural approaches to data mining, as they allow us to identify the most succinct and least redundant model for a dataset. MDL has been successfully employed for a wide range of data mining tasks, including discretization [7], outlier detection [28], classification [25], and clustering [17].

Miettinen and Vreeken [21, 22] recently formulated the BMF problem in terms of the Minimum Description Length (MDL) principle [11, 26] in order to solve the model order selection problem. Loosely speaking, this objective identifies the best factorization as the one that provides the best lossless compression of the data – thus automatically balancing the complexity of the factorization with the reconstruction error. The authors applied their score as post-processing for $A_{ss}$ to identify the optimal model order. In this paper we propose an algorithm that directly optimizes the description length, taking advantage of the fact that this objective can naturally handle different amounts of noise per type.

Tiling [10] is closely related to BMF, but aims at finding submatrices full of 1s. Xiang et al. [32] proposed an algorithm to mine noisy tiles – i.e. allowing 0s in the area covered by a tile. Given a collection of noisy tiles, Kontonasios and De Bie [12] iteratively discover the most interesting tile, defining interestingness through a local MDL score. Tatti and Vreeken [29] use MDL to hierarchically identify the most informative tile from data with ordered rows and columns.

Closer to our work is the $Panda$ algorithm proposed by Lucchese et al. [15]. $Panda$ performs Boolean matrix factorization, but instead of minimizing only the error, it minimizes the sum of Frobenius norms of the residual and the factor – by which the hidden assumption again is that false positives and false negatives are equally likely. Recently, the same authors proposed $Panda+$ [16] in which they optimize the TypedXOR encoding of Miettinen and Vreeken [21]. $Panda+$ will be the main competitor to $Nassau$.

5 Algorithm

In this section we present $Nassau$, a new algorithm for heuristically solving the MDLBMF problem. The existing algorithms, $Panda+$ and $A_{ss}$, never change an already-found factor. This property simplifies the search for the (locally) MDL-minimizing factorization [22], but has a drawback. The first few blocks blocks probably cover large parts of the data, but for higher-rank decompositions, these initial blocks can become too coarse-grained. $Nassau$, on the other hand, does iteratively refine previously discovered factors.

5.1 The main algorithm. $Nassau$ (Algorithm 1) interleaves the addition of new blocks and updating of already-found ones. As new blocks add more fine-grained structure, the coarser older factors become obsolete and can be replaced.

The algorithm starts by finding a set of seeds that provide the starting point for finding the factorization (Line 3; see Section 5.2). After initialization, $Nassau$ starts its first phase (Lines 4–10), where it repeatedly adds a new block

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**Algorithm 1 Nassau**

<p>| Input: $A \in {0,1}^{n \times m}, t, \tau, \theta \in (0,1), M &gt; 0$ |</p>
<table>
<thead>
<tr>
<th>Output: $B \in {0,1}^{n \times k}$ and $C \in {0,1}^{k \times m}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1: function Nassau($A, t, \tau, \theta, M$)</td>
</tr>
<tr>
<td>2: $B' \leftarrow \emptyset^{n \times 0}, C' \leftarrow \emptyset^{0 \times m}$</td>
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<tr>
<td>3: Seeds $\leftarrow GetSeeds(A)$</td>
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<tr>
<td>4: repeat</td>
</tr>
<tr>
<td>5: $B \leftarrow B', C \leftarrow C'$</td>
</tr>
<tr>
<td>6: $[b, c] \leftarrow GetBlock($A, $B, $C, $Seeds$)</td>
</tr>
<tr>
<td>7: $B' \leftarrow [B, b], C' \leftarrow [C, c]$</td>
</tr>
<tr>
<td>8: if $M$ rounds since last update then</td>
</tr>
<tr>
<td>9: $[B', C'] \leftarrow Cyc1Upd($A, $B', C', $Seeds, $0, \theta$)</td>
</tr>
<tr>
<td>10: until $L(A, B', C') \geq L(A, B, C)$</td>
</tr>
<tr>
<td>11: while not converged do</td>
</tr>
<tr>
<td>12: $[B, C] \leftarrow Cyc1Upd($A, $B, C, $Seeds, t, \theta$)</td>
</tr>
<tr>
<td>13: $t \leftarrow t \cdot \tau$</td>
</tr>
<tr>
<td>14: return $B, C$</td>
</tr>
</tbody>
</table>
to the existing factorization until the description length does not improve any more. The blocks are found using the FindBlock routine (Algorithm 2, see Section 5.3). To adjust the already-found factors, Nassau regularly calls CyclUpd (Algorithm 3, Section 5.4).

When additional factors do not decrease the objective further, we enter the last phase of the algorithm (Lines 11–13), which entails refinement of the discovered blocks. This step is explained in Section 5.4.

Nassau accepts several parameters that control its execution. Parameters \( t \) and \( \tau \) control the simulated annealing by giving the initial temperature and update ratio, respectively, while \( M \) controls the frequency at which we update the found factors. Hence, at the expense of run time, higher \( t \) and lower \( \tau \) and \( M \) potentially provide better results. Parameter \( \theta \) controls the mining process and is explained in Section 5.3.

5.2 Finding seed columns. Finding a good block to add is a hard problem. Hence, we take an approach similar to [16, 20], and start by finding a good collection of seed columns. The seed column vectors are collected in the \( n \)-by-\( s \) matrix \( \text{Seeds} \). Later, the FindBlock algorithm will use these seeds to build the final blocks. In principle, we can use any method to create these seeds, including those used in [16, 20], but here we present our approach, which is based on restarted random walks.

A good seed captures the correlation between data rows. Consider an \( n \)-by-\( m \) binary matrix \( A \) and the corresponding bipartite graph \( G = (V_{\text{rows}} \cup V_{\text{cols}}, E) \). The correlated rows of \( A \) correspond to the highly interconnected nodes in \( V_{\text{rows}} \), and if two nodes are correlated, then a short random walk starting from one of them is likely to frequently visit the other [27]. If we restart each walk from the same node frequently, the fraction of time we spend on each node indicates how related it is to the origin of the walk.

Let \( P(i \mid j) \) be the probability of reaching node \( i \) from node \( j \). We do one random walk for each node on the left part of the graph, and on every step we have a fixed probability \( \varepsilon \) to return back to the starting node. Otherwise, we go to one of the neighboring nodes, selecting them uniformly at random. Now, the fraction of time we spend in node \( i \) in a random walk starting from node \( v \) is

\[
\Pi_v(i) = \varepsilon \cdot 1(v = i) + (1 - \varepsilon) \sum_{(i,j) \in E} \Pi_v(j)P(i \mid j).
\]

The above equation can be solved using the dominant eigenvector, similar to [27]. We generate one seed for each data row \( v \), and the seed has 1 on every row where the random walk starting from \( v \) spends sufficient time.

5.3 Finding the blocks. Given the input matrix and the current factorization, FindBlock (Algorithm 2) finds a new block to be added to the factorization.

**Algorithm 2 FindBlock**

| Input: matrices \( A \in \{0,1\}^{n \times m} \), \( B \in \{0,1\}^{m \times k} \), \( C \in \{0,1\}^{k \times m} \), Seeds \( \in \{0,1\}^{m \times s} \), \( \theta \in (0,1) \) |
| Output: block \( (b^*, c^*) \) |

1. \( \text{function FindBlock}(A, B, C, \text{Seeds}, \theta) \)
2. \( b^* \leftarrow 0^m \), \( c^* \leftarrow 0^m \)
3. for \( i = 1 \) to \( s \) do
4. \( b \leftarrow \text{Seeds}^i \) \( \triangleright \) The \( i \)th seed column.
5. repeat
6. \( c \leftarrow \arg \max_{c \in \{0,1\}^m} \text{cover}(A, [B, b], [C], \theta) \)
7. \( b \leftarrow \arg \max_{b \in \{0,1\}^m} \text{cover}(A, [B, b], [C], \theta) \)
8. until stopping criteria are satisfied
9. if \( L(A, [B, b^*], [C]) < L(A, [B, b], [C]) \) then
10. \( b^* \leftarrow b \), \( c^* \leftarrow c \)
11. return \( (b^*, c^*) \)

The algorithm tries every seed one by one, and uses alternating updates. To compute the updates, we ignore all the locations where the existing factorization has a non-zero, as we cannot change that value. Then, given a fixed column factor \( b \), we build the corresponding row factor \( c \) by testing, for every data column, if using \( b \) to cover that column would improve the cover function:

\[
\text{cover}(A, B, C, \theta) = \theta \left| \{(i,j) : a_{ij} = 1 \land (B \circ C)_{ij} = 1\} \right| - \left| \{(i,j) : a_{ij} = 0 \land (B \circ C)_{ij} = 1\} \right|.
\]

We use the cover function as a surrogate to the full description length, partially to make this update step faster, and partially to avoid some local optima. Further local optima can be avoided by adjusting parameter \( \theta \); setting \( \theta = 1 \) returns locally optimal blocks, but we can introduce some locally nonoptimal decisions (that can yield to better global behaviour) by setting \( \theta \) to a slightly larger (or smaller) value.

5.4 Updating the factors. As was mentioned above, blocks found earlier might become outdated – that is, they become less useful in covering data, or they can even become redundant. This happens if we have added new blocks that overlap with previous ones and cover the same data better. In these cases we can improve the factorization by updating the old blocks. CyclUpd (Algorithm 3) iteratively replaces blocks with better alternatives. This is performed by first removing a block, and then finding a replacement by calling FindBlock with the current factors (Line 4).

Note that the objective is not strictly decreasing: FindBlock is heuristic and is not guaranteed to improve the result. However, we might still want to keep the update to avoid getting stuck in a local minimum. In order to accomplish this, we apply a technique similar to simulated annealing. We use a temperature parameter \( t \), which controls the probability of accepting a new block that does not improve the score. If a new block decreases the objective, we always accept it; otherwise we will only do so with a probability...
Algorithm 3 CyclUpd

Input: matrices $A \in \{0, 1\}^{n \times m}$, $B \in \{0, 1\}^{n \times k}$, $C \in \{0, 1\}^{k \times m}$, Seeds $\in \{0, 1\}^{n \times k}$, $t > 0$, $0 < \theta < 1$.

Output: Factors $B^* \in \{0, 1\}^{n \times k}$ and $C^* \in \{0, 1\}^{k \times m}$.

1: function CyclUpd($A, B, C, \text{Seeds}, t, \theta$)
2: $B^* \leftarrow B, C^* \leftarrow C$
3: for $t = 1$ to $k$ do
4:   $[b, c] \leftarrow \text{FindBlock}(A, B^{-l}, C^{-l}, \text{Seeds}, \theta)$
5:   $B^* \leftarrow [B^{-l}, b], C^* \leftarrow [C^{-l}, c]$  \>	ext{Replace block}
6:   if $L(A, B^*, C^*) < L(A, B, C)$ then
7:     $B \leftarrow B^*, C \leftarrow C^*$
8:   else
9:     $B \leftarrow B^*, C \leftarrow C^*$ with probability $t$
10: if $L(A, B^*, C^*) < L(A, B^*, C^*)$ then
11:   $B^* \leftarrow B, C^* \leftarrow C$
12: return $B^*, C^*$

6 Experiments

In this section we empirically evaluate Nassau and compare its performance to that of two competing algorithms.

6.1 Algorithms and parameters. We used three algorithms, Nassau,\footnote{The Nassau code is available for research purposes at http://people.mpi-inf.mpg.de/~pmiettin/nassau/; for the other two algorithms, we used code provided by the authors.} Panda+ [16], and Asso [20]. For Asso, we followed the approach of [22] to compute the MDL-minimizing factorization. These algorithms take various parameters. For Nassau we used the same parameters in all experiments, and we set the initial annealing temperature $t = 0.8$, temperature scaling $\tau = 0.6$, weight $\theta = 1.1$, and launched the cyclic updates in every $M = 5$ rounds. Asso takes one parameter, the rounding threshold $\tau$. For synthetic experiments, we tried values from 0 to 1 at increments of 0.1; for real-world experiments, we followed the setup of [22], and tried the values from 0.1 to 0.95 with increments of 0.025. In all cases, we selected the value of $\tau$ that yields the minimal MDL cost factorization. For Panda+, we minimize the description length (i.e., the $J_2$ error function) and use randomization with 10 re-starts. Other parameters were left at their default values.

Panda+ optimizes an encoding that differs slightly to the one we focus on. For fair comparison we thus ran a set of baseline experiments with both encodings. This, however, did not give results that were different in any significant way, and hence we do report these results separately – the fact that the encoding we use gives very similar results to the encoding used by Panda+ was also noted in [22].

### 6.2 Synthetic data.

We first use synthetic data to test the algorithms on data of known ground truth and characteristics. To that end, we generated 1000-by-800 matrices by first creating two binary factor matrices $B^*$ and $C^*$, computing their product $A = B^* \circ C^*$, and applying noise to $A$ in order to obtain the final input matrix $A$. In addition to the results from the algorithms, we also report the results obtained with the original factor matrices $B^*$ and $C^*$. We call this the True Model. The amount of noise is measured in percentages of non-zeros in $A$. That is, $p\%$ of destructive noise flips an expected $p|A|/100$ of the $1$s of $A$, while $p\%$ of additive noise flips an expected, $p|A|/100$ of the $0$s.

When generating the matrices, we varied one parameter and kept the rest fixed. In particular, we evaluated different levels of additive and destructive noise, density of the matrix, and rank (model order). All results are averaged over 10 instances per configuration. When not varied, the rank was kept at 15, the density at 8%, and additive noise at 3%. To evaluate how well the algorithms perform with increasing amounts of destructive noise, we ran two sets of experiments, one with 15% destructive noise and one with 50%.

**Destructive noise.** We start by testing the effects of the destructive noise, shown in Figure 1a. Initially Nassau and Asso are indistinguishable from the true model, while Panda+ has much higher description length. At around 40% destructive noise, however, Nassau starts to perform better than Asso, and continues in that way until the end. Meanwhile, we observe that while Panda+ under-performs overall, it does relatively well when the data becomes very sparse (high levels of destructive noise). This may mean its search procedure is well-equipped to find blocks in sparse data.

**Additive noise.** The purpose of this test is to find how robust the algorithms are to additive noise, that is when $0$s are turned to $1$s. Keeping the destructive noise constant at 15%, we find that up to 60% additive noise both Nassau and Asso still find models virtually indistinguishable from the
true model, while Panda+ is consistently worse except at 70% noise (Figure 1b).

Varying density. We varied the density of the noise-free matrices from 2% to 20%. At 15% destructive noise (Fig. 1c), Nassau and Asso discover models on par with the true model, while Panda+ is clearly much worse. With 50% destructive noise (Fig. 1d), Nassau and Panda+ are virtually indistinguishable until the density reaches 16%, after which the performance of Panda+ starts to degrade.

Model order. Last, we varied the model order (rank) between 2 and 20. With 15% destructive noise, Nassau and Asso obtained exact results, while Panda+ was mostly underestimating (Fig. 1e). With 50% destructive noise (Fig. 1f), Nassau over-estimates slightly more than Panda+, although for both methods we see occasional high variance.

6.3 Real-world datasets. Next, we consider 10 real-world datasets, most of which are publicly available. Below we give a short description for each, and an overview in Table 1.

4News is a subset of the 20Newsgroups data, containing the usage of 800 words over 400 posts for 4 newsgroups.

Abstracts represents the words in all abstracts of the papers accepted at the ICDM conference up to 2007, where the words have been stemmed and stop words removed.

DBLP conf. contains records of which of the 19 conferences the 6980 authors had published in. The dataset is collected from the DBLP database and it is pre-processed as in [18].

DBLP co-auth. is a (symmetric) co-authorship matrix of a subset of the authors in the DBLP conf. data.

Dialect contains presence data of dialectal linguistic properties in 506 Finnish municipalities.

DNA Amp. contains information on DNA copy number amplifications. Such copies activate oncogenes and are hallmarks of nearly all advanced tumours.

Firewall 2 describes the reachability between two IP addresses.[5] It has an exact Boolean decomposition of 10 factors [5], and hence should be highly compressible.

Mammals consists of presence records of European mammals within areas of 50 by 50 kilometers.

Mushroom contains edibility records of mushrooms.

Lastly, Paleo consists of fossil records per location.

6.4 Quantitative evaluation of real-world results. We start by studying the compression ratios of the real-world data sets. By compression ratio, we mean the description

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2The authors are grateful to Ata Kabán for pre-processing the data [18].
3Available upon request from the author [2]
length $L$ obtained by the algorithm divided by the description length of the data using an empty model, i.e. when no factors are used to represent it, $L_0$. The smaller this ratio is, the better the discovered factorization compresses the data. The results are presented in Table 2.

Out of the ten datasets in Table 2, Nassau obtains the best compression ratio in five – and in each of these cases it outperforms the second-best result by at least one percentage point. Panda+ is the best for the two sparse text datasets, 4News and Abstracts, although for 4News Nassau is only 0.4 and Asso only 0.9 percentage points worse. For all practical purposes, we would consider 4News a tie. Similarly, Asso is better in DBLP co-auth., but only by 0.3 percentage points, more clearly in Firewall 2, and by a wide margin in Mushroom.

If we consider the two cases where Nassau is significantly worse than the best method, Abstracts (against Panda+) and Mushroom (against Asso), we notice that in both cases Nassau reports significantly smaller model order than the best method (3 vs. 128 and 4 vs. 59, respectively). It seems that in these cases the random walks were unable to find good seed vectors. Indeed, when we re-ran Nassau on Mushroom but generating the seeds as Asso does, we obtained a compression ratio of 56.4 with 44 factors – a significant improvement confirming that the random walks do not necessarily produce good seeds for every data set.

**Table 2**: Compression ratio $L\% = 100 \times L/L_0$ (smaller is better) and model order $k$ for the real-world datasets.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Nassau</th>
<th>Panda+</th>
<th>Asso</th>
</tr>
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<td>93.6</td>
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<td>86.7</td>
<td>97.2</td>
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<tr>
<td>DBLP co-auth.</td>
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<td>92.4</td>
<td>90.0</td>
</tr>
<tr>
<td>DBLP co-auth.</td>
<td>94.1</td>
<td>95.9</td>
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<tr>
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<td>48.8</td>
</tr>
<tr>
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<td>1.7</td>
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<td>Mammals</td>
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<td>64.6</td>
</tr>
<tr>
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<td>Paleo</td>
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**Scalability.** The implementations of these methods are not fully comparable. Panda+ is a full-C implementation, Asso is a mixture of C and Matlab, while Nassau is written purely in Matlab. Also, the different methods use different levels of parallelization, and the running time for all these algorithms is heavily influenced by the eventual model order. For Asso, the number of different rounding thresholds $\tau$ to try, and for Panda+, the number of random re-starts, also have obvious effects. That said, in our experiments, Panda was generally the fastest, followed by Nassau and Asso, but the order could vary from dataset to dataset. All methods were able to finish all datasets within 24 hours, except for Asso with DBLP co-auth., which took more than four days.

6.5 Qualitative evaluation of real-world results. We studied the results on two datasets, DBLP co-auth. and Mammals, more carefully in order to understand the types of results we can obtain from Nassau (and its competitors). As Table 2 shows, DBLP co-auth. is not very compressible, while Mammals is reduced to almost half of its original size.

**DBLP co-auth.** The DBLP co-auth. data is a graph, i.e. a symmetric matrix, and while none of the methods tested here assume symmetry, we would expect that a good factorization would be (approximately) symmetric. This, however, is not the case for Panda+ or Asso: in symmetric decompositions, the two factor matrices would be the same, but for Asso, 1.2% of the values differ, and for Panda+, 1.6% of the values in the factor matrices differ. For Nassau, only 0.3% of the values are different.

Looking at the factors, we observed that in particular Asso finds very skewed blocks that have many rows but few columns, or vice versa, being clearly unable to capitalize on the symmetry of the matrix. Panda+ returns more square-like blocks, but they still have almost four times as many as columns (or vice versa) on average. The blocks Nassau returns have an average rows/columns ratio of 1.7, giving the most square-like blocks.

We show two exemplar factors found by Nassau in the DBLP co-auth. data. It shows the known collaborations between the authors in a factor. Here, Nassau has identified two (quasi-) cliques of famous data mining and machine learning researchers with strong collaboration patterns.

**Mammals.** Both Panda+ and Asso perform so-called hierarchical factorization (rank-$(k-1)$ factorization) is part of the rank-$k$ factorization), while Nassau is designed to avoid that. We studied these different behaviours with the Mammals data and using the first four left-hand-side factors (i.e. columns of $B$) for each method. As the rows of the data correspond to locations in Europe, we visualize the factors...
using maps in Figure 3. For Nassau the first four factors are very compact and correspond very closely to countries in Europe. For Panda+, the first factor covers almost the whole area of Europe that was part of the data, and the other three factors also cover large, not very well-defined areas. The same holds true for Aso, although to a slightly lesser extent. In other words, the results by Nassau in Figure 3 correspond to species that are very specific to certain areas, while both Panda+ and Aso selected common species.

7 Discussion

The experiments show that Nassau performs well in practice. The evaluation on synthetic data shows it is resistant to high amounts of destructive noise, reliably discovering factorizations of the correct model order and of complexities close to the ground truth. For real-world data Nassau discovered the most succinct descriptions for five out of ten datasets, while being a close second for three more. Manual inspection showed that the factors Nassau finds are concise, clean, and more interpretable than those of Aso and Panda+. Overall, the experiments permit two conclusions: 1) Nassau performs very well when the data exhibits large amounts of destructive noise, and 2) our hypothesis that real-world data contains large amounts of destructive noise is warranted.

We see two main avenues for further research. First of all, the seed generation process of Nassau can be improved. Preliminary experiments suggest that by adding the seeds that Aso considers to Nassau, even better factorizations can be discovered – alternatively, it will be interesting to see if in addition the seed generation process of Panda+ can be incorporated. The second line of research is optimization, of both the process and the implementation.

8 Conclusions

We studied the problem of Boolean matrix factorization under the assumption that in general it is more likely not to have observed the value of a cell than to have done so – and that the reliability of both observations differs. We formalized this problem by the Minimum Description Length principle, and proposed the new algorithm that directly optimizes the description length of the factorization. Nassau is non-hierarchical and is capable of fixing errors it has previously made. Empirical evaluation on synthetic data shows that Nassau is very robust to high levels of destructive noise. Experiments on real-world data show it finds more compact and intuitive factors than the state of the art, which are more inclined to ‘simply’ cover large parts of the data.

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References

Figure 3: **We discover compact descriptions** The first four factors discovered on the *Mammals* data using, from top to bottom, Nassau, Panda+, and Asso.


