VOUW: Geometric Pattern Mining using the MDL Principle

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Abstract

- MDL is a principle that utilizes compression as a means of describing data
- There exist many MDL-based algorithms for a variety of problems
- Few (no?) solutions exist to mine grid-like data based on MDL
- VOUW is a novel approach to discover patterns and structural relations in 2D, discrete datasets
- We propose both a theoretical framework as well as a complete, optimized implementation

1 Introduction

Frequent pattern mining is the subfield of data mining that aims to find and extract recurring substructures as a means of data analytics. These patterns can be any kind of datastructure, for example item sets (transactions), graphs (networks) and sequences (in time). So far, little research has been done in applying pattern mining to raster-based, tabular data or matrices. Patterns in these types of data can be more complex as elements cannot only coincide, but their geometric relation is also important. As the first contribution of this paper, we introduce exactly this problem, that we will call **geometric pattern mining** and formally introduce notation and theoretical constructions for this problem. Potential applications include analysis of (satellite) imagery, texture recognition and clustering of matrices.



(a) 32×24 noise 'matrix' (b) Pair (146, 11) (c) Pattern 'I' occurs four times.

Fig. 1: Brief example of geometric pattern mining

Geometric Pattern Mining Geometric pattern mining is the problem of finding recurring local structure or patterns in matrices. It is different from graph mining, as a matrix is more rigid and each element has a fixed degree of connectedness/adjacency. It is also unrelated to linear algebra, other then using the term 'matrix' and a comparable style of notation. Furthermore in this context, matrix elements can only be discrete, rows and columns have a fixed ordering and the semantics of a value is position-independent. Although the concept applies to any number of dimensions, we will limit the scope to two dimensional data from here on.

The problem of geometric pattern mining can roughly be divided into three classes. The first class consists of three subclasses: finding identical patterns in (1a) an otherwise empty (sparse) matrix, (1b) differently distributed noise and (1c) similarly distributed noise. The second class contains the same subclasses but adds that the patterns can also be overlaid with noise and are therefore not identical. The third class is also a continuation of the first class and requires that the patterns are identical after some optional transformation (such as mirror, inverse, rotate, etc.). These classes also represent an increasing difficulty level and serves as a rough benchmark for the performance of an algorithm.

Let us demonstrate the problem of geometric pattern mining with a brief example. Figure 1a shows a 32×24 grayscale 'matrix' filled with noise on the interval [0; 255]. If we look at all horizontal pairs of elements, we find that the pair (146, 11) is, among others, statistically more prevalent than 'random noise' suggests. If we would continue to try all combinations of elements that 'stand out' from the background noise, Figure 1c shows that we will eventually find four copies of the letter 'I' set in 16 point Garamond Italic.

The 35 elements that make up a single 'I' in the example form a **pattern**. We can use this pattern to describe the matrix in an other way than '768 unrelated values'. For example, we could describe it as 628 unrelated values plus pattern 'I' at locations (5,4),(11,11),(20,3),(25,10), separating the structure from the accidental (noise) data. Since this requires less storage space than before, we have compressed the data. At the same time we have learned something about this data, namely that it contains four 'hidden I's.

Datamining by Compression In recent years, a class of algorithms utilizing the *Minimum Description Length (MDL) principle* [6, 3] have become more and more common in the field of explanatory data analysis. Examples of such approaches include Krimp [9] or more recently Classy [5]. The MDL principle was first described by Rissanen in 1987 [6] as a practical implementation of Kolmogorov Complexity [4]. Central to MDL is the notion that 'learning' can be thought of as 'finding regularity' and that regularity itself is a property of data that is exploited by *compressing* said data. Therefore by compressing a dataset, we actually learn its structure — how regular it is, where this regularity occurs, what it looks like — at the same time.

The problem that MDL solves first and foremost, is that of *model selection*: given a multitude of explanations (models), select the one that fits the data

best. In addition to this, MDL has also been demonstrated to be very effective in materialization of a specific model given the data. In this case, the model is predetermined and we want to find the parameters to fit the data. A similar problem class is solved in pattern mining: here the 'parameters' are the discrete building blocks that make up patterns in the data. We will specifically look at a variant of MDL called two-part MDL. As the second contribution of this paper, we present a geometric pattern mining algorithm that (1) is precise, (2) requires no parameters and (3) is tolerant to noise in the data, based on two-part MDL.

2 Related Work

Krimp [9] is probably one of the first explanatory data mining approaches using MDL and also one of the sources of inspiration for this paper. Since then, many papers on this topic have been published, such as Slim ?? (frequent item set mining, like Krimp), Classy ?? (mining classifiers), TODO: list more

The work by Campana et al. [1] also uses matrix-like input data (textures) and develops a similarity measure based on MDL. Their method, however, cannot be used for *explanatory* data analysis as they use a generic image compression algorithm that is essentially a black box.

This is still far from complete

3 Theoretical Framework

We define geometric pattern mining on bounded, discrete and two-dimensional raster-based data. We represent this data as an $M \times N$ matrix A whose rows and columns are finite and in a fixed ordering (i.e. reordering rows and columns semantically alters the matrix). For elements $a_{i,j}$, where row i is on [0; N) and column j is on [0; M), holds that $a_{i,j} \in S$, the finite set of symbols over A.

According to MDL, the shortest (optimal) description of A reveals all structure of A in the most succinct way possible and we approximate this optimal description through the compression of the original data. This optimal description A' is only optimal if we can unambiguously reconstruct A from it and nothing more — the compression is both minimal and lossless. Intuitively compression means defining A using as few building blocks as possible. We illustrate this in Figure 2. Given the matrix A we decompose it in patterns, denoted X and Y. The set of all these patterns is the **model** of a A, denoted H_A . In order to reconstruct A from this

$$A = \begin{bmatrix} 1 & \cdot & \cdot & 1 & 1 \\ \cdot & 1 & \cdot & \cdot & \cdot \\ 1 & \cdot & \cdot & 1 & \cdot \\ \cdot & 1 & \cdot & \cdot & 1 \\ 1 & 1 & 1 & 1 & \cdot \end{bmatrix}, \ I = \begin{bmatrix} X & \cdot & \cdot & Y & \cdot \\ \cdot & \cdot & \cdot & \cdot & \cdot \\ X & \cdot & \cdot & X & \cdot \\ \cdot & \cdot & \cdot & \cdot & \cdot \\ Y & \cdot & Y & \cdot & \cdot \end{bmatrix}, \ H = \{X = \begin{bmatrix} 1 & \cdot \\ \cdot & 1 \end{bmatrix}, Y = \begin{bmatrix} 1 & 1 \end{bmatrix}\}$$

Fig. 2: Example decomposition of A into instantiation I and patterns X, Y

model, we also need a mapping from the H_A back to A. This mapping represents in what MDL calls the **the data given the model** H_A . In this context we think of it as 'instructions' of how to reconstruct A. Let us call the set of all instructions required to rebuild A from H_A the **instantiation** of H_A . It is denoted by I_A in the example. The result is a notation that allows us to express matrix A as if decomposed into sets of local and global spatial information, which we will now describe in more detail.

3.1 Patterns and Instances

 \triangleright We define a **pattern** as a $M_X \times N_X$ submatrix X of the original matrix A. Elements of this submatrix may be \cdot , the empty element, which gives us the ability to cut-out any irregular-shaped part of A. We additionally require the elements of X to be adjacent (horizontal, vertical or diagonal) to at least one non-empty element and that no rows and columns are empty.

From this definition, the dimensions $M_X \times N_X$ give the smallest rectangle around X (the bounding box). We also define the cardinality |X| of X as the number of non-empty elements. We call a pattern X with |X| = 1 a **singleton pattern**, i.e. a pattern containing exactly one element of A.

Each pattern contains a special **pivot** element: pivot(X) is the first non-empty element of X. A pivot can be thought of as a fixed point in X which we can use to position its elements in relation to A. This translation, or **offset**, is a tuple q = (i, j) that is on the same domain as an index in A. We realize this translation by placing all elements of X on an empty $M \times X$ size matrix in such that the pivot element is at (i, j). We formalize this in the **instantiation operator** \otimes : \triangleright We define the **instance** $X \otimes (i, j)$ as the $M \times N$ matrix containing all elements of X such that pivot(X) is at index (i, j) and the distances between all elements are preserved. The resulting matrix contains no additional non-empty elements.

Obviously this does not yield a valid result for an arbitrary offset (i,j). We want to limit ourselves to the space of pattern instances that are actually valid in relation to matrix A. Therefore two simple constraints are needed: (1) an instance must be **well-defined**: placing pivot(X) is at index(i,j) results in an instance that contains all elements of X, and (2) elements of instances cannot overlap, meaning each element of A should be described at most once. This allows for a description that is both unambiguous and minimal.

ightharpoonup Two pattern instances $X \otimes q$ and $Y \otimes r$, with $q \neq r$ are **non-overlapping** if $|(X \otimes q) + (Y \otimes r)| = |X| + |Y|$.

From here on we will use the same letter in lower case to denote an arbitrary instance of a pattern, e.g. $x = X \otimes q$ when the exact value of q is unimportant.

We briefly introduced the instantiation I as a set of 'instructions' of where instances of each pattern should be positioned in order to obtain A. As Figure 2 suggests, this mapping has the shape of an $M \times N$ matrix.

 \triangleright Given the set of patterns H, the **instantiation (matrix)** I is an incomplete $M \times N$ matrix such that $I_{i,j} \in H \cup \{\cdot\}$ for all (i,j). For all non-empty elements $I_{i,j}$ it holds that $I_{i,j} \otimes (i,j)$ is a non-overlapping instance of $I_{i,j}$ in A.

3.2 The Problem and its Solution Space

Patterns can be constructed by joining smaller patterns in a bottom-up fashion. We can define the exact way two patterns should be joined by enumerating the distance of their respective pivots. To limit the possibilities to patterns relevant to A, instances can be used as an intermediate step. Since instances are simply patterns projected on an $M \times N$ matrix, we can reverse \otimes by removing all completely empty rows and columns:

 \triangleright Let $X \otimes q$ be an instance of X, then by definition we say that $\emptyset(X \otimes q) = X$.

As Figure 3 demonstrates, we can use a simple element-wise matrix addition to sum two instances and use \oslash to obtain a joined pattern. Here we start by instantiating X and Y with offsets (1,0) and (1,1) respectively. We add the resulting x and y to obtain $\oslash z$, the union of X and Y with relative offset (1,1)-(1,0)=(0,1).

The Sets \mathcal{H}_A and \mathcal{I}_A We define the model class \mathcal{H} , the set of all possible models for all possible inputs. Without any prior knowledge, this is the search space of our algorithm. We will first look at a more bounded subset \mathcal{H}_A of all possible models for A, and \mathcal{I}_A , the set of all possible instantiations to these models. We will also take H_A^0 to be the model with only singleton patterns. As singletons are just individual elements of A, we can simply say that $H_A^0 = S$. The instantiation matrix corresponding to H_A^0 is denoted I_A^0 . Given that each element of this matrix must correspond to exactly one element of A in H_A^0 , we see that each $I_{i,j} = a_{i,j}$ and so I_A^0 is equal to A.

Using H_A^0 and I_A^0 as base cases we can now inductively define the set \mathcal{I}_A :

Base case $I_A^0 \in \mathcal{I}_A$

By induction If I is in \mathcal{I}_A then take any pair $I_{i,j}, I_{k,l} \in I$ such that $(i,j) \leq (k,l)$ in lexicographical order. Then the set I' is also in \mathcal{I}_A , providing I' equals I except: $I'_{i,j} := \emptyset \big(I_{i,j} \otimes (i,j) + I_{k,l} \otimes (k,l) \big)$

 $I'_{k,l} := \cdot$

This shows we can add any two instances together, which are by definition always non-overlapping and thus valid in A, in any order and obtain an element of \mathcal{I}_A . Eventually this results in just one big instance that is equal to A. Note that when we take two elements $I_{i,j}, I_{k,l} \in I$ we force $(i,j) \leq (k,l)$, not only to eliminate different routes to the same instance matrix, but also such that the pivot of the new pattern coincides with $I_{i,j}$. We can then leave $I_{k,l}$ empty.

$$x = X \otimes (1,0) = \begin{bmatrix} \cdot & \cdot \\ 1 & \cdot \\ \cdot & 1 \end{bmatrix}, \ y = Y \otimes (1,1) = \begin{bmatrix} \cdot & \cdot \\ \cdot & 1 \\ \cdot & \cdot \end{bmatrix}, x + y = \begin{bmatrix} \cdot & \cdot \\ 1 & 1 \\ \cdot & 1 \end{bmatrix}, \ Z = \oslash(x + y) = \begin{bmatrix} 1 & 1 \\ \cdot & 1 \end{bmatrix}$$

Fig. 3: Example of joining patterns X and Y to construct Z.

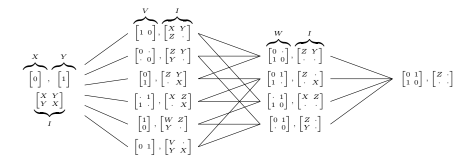


Fig. 4: Model space lattice for a 2×2 Boolean matrix. The Left hand column shows which pattern is added in each step, I is the current instantiation.

The construction of \mathcal{I}_A also implicitly defines \mathcal{H}_A . While this may seem odd — defining models for instantiations instead of the other way around — note that there is no unambiguous way to find one instantiation for a given model. Instead we find the following definition by applying the inductive construction:

$$\mathcal{H}_A = \{ \{ \emptyset(I) \mid I \in I \} \mid I \in \mathcal{I}_A \}. \tag{1}$$

So for any instantiation $I \in \mathcal{I}_A$ there is a corresponding set in \mathcal{H}_A of all patterns that occur in I. This results in an interesting symbiosis between model and instantiation: increasing the complexity of one decreases that of the other. This construction gives a tightly connected lattice as shown in Figure 4.

3.3 Encoding Models and Instances

From all parametrized models in \mathcal{H}_A we want to select (approximate) the model that describes A best. We use two-part MDL to quantify how well a given model and instantiation matrix fit A. Two-part MDL tells us to minimize the sum of $L_1(H_A) + L_2(A|H_A)$, two functions that give the length of the model and the length of 'the data given the model', respectively. In this context, the model is the set of patterns H_A and the data given the model is I_A , the accidental information needed to reconstruct the data from H_A .

In order to compute their lengths, we need to decide on a way to encode H and I first. This encoding is of great influence on the length functions and therefore on the ability to accurately quantify the fit of a given H and I to A. Although the decision for this encoding is obviously prone to bias, practice shows that good results can be had once certain conditions are met: (1) all data is encoded (lossless) and (2) the encoding is as concise as possible (nothing but the data is encoded). Based on these conditions we give length functions for pattern sets and instantiations, but we do not actually need to encode them.

The fictional encoder sequentially sends each symbol in the datastream to the 'decoder' using a code word. Information theory tells us that we optimal length of a code word is given by $-\log(p)$, where p is the exact probability that the code

Table 1: Length computation for the different classes of matrices. The total length is the sum of the listed terms.

| Matrix | Bounds | # Elements Positions | Symbols |
|---------------------|--------------|----------------------|----------------|
| $L_p(X)$ Pattern | $\log(MN)$ | (A) | $\log(S)$ |
| $L_1(H)$ Model | N/A | $L_N(H)$ N/A | $L_p(X \in H)$ |
| $L_2(I)$ Inst. mat. | . $constant$ | $\log(MN)$ implicit | $L_{pp}(I)$ |

word occurs in the output. We therefore need not compute the actual code words, just their probabilities. For this to work, both the encoder and hypothetical decoder must know either the exact probability distribution or agree upon an approximation beforehand. Such approximation is often called a **prior** and is used to fix 'prior knowledge' that does not have to be encoded explicitly.

The length function for incomplete matrices. To losslessly encode A' we have to encode both H and I individually. As both instances and patterns are both matrices it is tempting to utilize the same length function for both. Empirical evidence has shown that this is not a good idea though. For example, we do not consider certain values such as the size of the instance matrix because it is constant, however, the size of each individual pattern is not. We therefore have to construct a different length function for each type of matrix. These are listed in Table 1 When encoding I, we observe that it contains each pattern $X \in H$ multiple times, given by the **usage** of X. Using **prequential plug-in code** [3] to encode I enables us to omit encoding these usages separately, which would create unwanted bias. Prequential plug-in code gives us the following length function for I. We elaborate on the derivation of this equation in Appendix A.

$$L_{pp}(I \mid P_{plugin}) = -\sum_{X_i \in h}^{|H|} \left[\log \frac{\Gamma(\text{usage}(X_i) + \epsilon)}{\Gamma(\epsilon)} \right] + \log \frac{\Gamma(|I| + \epsilon|H|)}{\Gamma(\epsilon|H|)}$$
 (2)

Each length function has four terms. First we encode the total size of the matrix. Since we assume MN to be known/constant, we can use this constant to define the uniform distribution $\frac{1}{MN}$, such that $\log MN$ encodes an arbitrary index of A. Next we encode the number of elements that are non-empty. For patterns this value is encoded together with the third term, namely the positions of the non-empty elements. We use the previously encoded M_XN_X in the binominal function to enumerate the ways we can place the (|X|) elements onto a grid of M_XN_X . This gives us both how many non-empties there are as well as where they are. Finally the fourth term is the length of the actual symbols that encode the elements of matrix. In case we encode single elements of A, we simply assume that each unique value in A has an equal possibility of occurring. For the instance matrix, which encodes symbols to patterns, the prequential code is used as demonstrated before. Notice that L_N is the universal prior for integers[7] that can be used to encode integers on an arbitrary range.

4 A Search Algorithm

Pattern mining problems often yield vast search spaces and geometric pattern mining is no exception. Since we are already looking for an approximate result (by definition of two-part MDL) it makes sense to use a greedy strategy, a heuristic widely used in many MDL-based approaches [9, 8, 5].

Given these heuristics, an inductive algorithm is devised similar to the lattice in Figure 4: we start with a completely underfit model (left of the lattice), where there is one instance for each matrix element. On each iteration we combine two patterns, resulting in one or more pairs of instances to be merged (one step right in the lattice). We pick the pair of patterns that improve the compression ratio the most and we repeat these steps until no improvements are possible.

4.1 Finding candidates

The first step of the algorithm is to find the 'best' **candidate** pair of patterns to merge. Candidates are denoted as a tuple (X, Y, δ) , where X and Y are patterns and δ the relative offset of X and Y as they occur in the data. All possibilities form an enormous vector space. Fortunately, we need only pairs of patterns and offsets that actually occur in the instance matrix. This means at each step we can directly enumerate candidates from the instantiation matrix and never even look at the original data.

The **support** of a candidate, written $\sup(X,Y,\delta)$, tells how often it is found in the instance matrix. Computing support is not completely trivial, as one candidate occurs multiple times in 'mirrored' configurations, such as (X,Y,δ) and $(Y,X,-\delta)$, which are equivalent but can still be found separately. Furthermore, due to the definition of a pattern, many potential candidates cannot be considered by the simple fact that their elements are not adjacent.

Peripheries. For each instance x we define its periphery: the set of instances which are positioned such that their union with x produces a valid pattern. This set is split into the anterior- ANT(X) and posterior POST(X) peripheries, containing instances that come before and after x in lexicographical order, respectively. This enables us to scan the instance matrix once, in lexicographical order. For each instance x, we only consider the instances POST(x) as candidates, thereby eliminating any (mirrored) duplicates.

Self-overlap. Self-overlap happens for candidates of the form (X, X, δ) . In this case, too many or too few copies may be counted. Take for example a straight line of five instances of X. There are four unique pairs of two X's, but only two can be merged at a time, in three different ways. Therefore, when considering candidates of the form (X, X, δ) , we also compute an *overlap coefficient*. This coefficient e is given by the equation $e = (2N_X + 1)\delta_i + \delta_j + N_X$. This equation essentially transforms δ into a one-dimensional coordinate space of all possible ways that X could be arranged *after* and *adjacent* to itself. For each instance x_1 a vector of bits V(x) is used to remember if we have already encountered a combination x_1, x_2 with coefficient e, such that we do not count a combination

| Algorithm 1 Find Candidates | Algorithm 2 Merge Candidate |
|---|---|
| | |
| Input: I | Input: H, I |
| Output: C | 1: $C \leftarrow \text{Find Candidates}$ |
| 1: for all $x \in I$ do | 2: $(X, Y, \delta) \in C : \forall_{c \in C} \Delta L((X, Y, \delta)) \leq \Delta L(c)$ |
| 2: for all $y \in POST(x)$ do | 3: $\Delta L_{best} = \Delta L((X, Y, \delta))$ |
| 3: $X \leftarrow \emptyset(x), Y \leftarrow \emptyset(y)$ | 4: if $\Delta L_{best} > 0$ then |
| 4: $\delta \leftarrow \operatorname{dist}(X, Y)$ | 5: $Z \leftarrow \emptyset(X \otimes (0,0) + (Y \otimes \delta))$ |
| 5: if $X = Y$ then | 6: $H \leftarrow H \cup \{Z\}$ |
| 6: if $V(x)[e] = 1$ continu | ue 7: for all $x_i \in I \mid \emptyset(x_i) = X$ do |
| 7: $V(y)[e] \leftarrow 1$ | 8: for all $y \in POST(x_i) \mid \emptyset(y) = Y$ do |
| 8: end if | 9: $x_i \leftarrow Z, y \leftarrow \cdot$ |
| 9: $C \leftarrow C \cup (X, Y, \delta)$ | 10: end for |
| 10: $\sup(X, Y, \delta) += 1$ | 11: end for |
| 11: end for | 12: end if |
| 12: end for | 13: repeat until $\Delta L_{best} < 0$ |

 x_2, x_3 with an equal e. This eliminates the problem of incorrect counting due to self-overlap.

4.2 Gain computation

After candidate search we have a set of candidates C and their respective supports. The next step is to select the candidate that gives the best gain: the improvement in compression by merging the pattens in the candidate. For each candidate $c = (X, Y, \delta)$ the gain $\Delta L(A', c)$ is comprised of two parts: (1) the negative gain of adding the union pattern Z to the model H, resulting in H' and (2) the gain of replacing all instances x, y with relative offset δ by Z in I, resulting in I'. We use the length functions L_1, L_2 to derive an equation for gain:

$$\Delta L(A',c) = \left(L_1(H') + L_2(I')\right) - \left(L_1(H) + L_2(I)\right)$$

$$= L_0(|H|) - L_0(|H| + 1) - L_p(Z) + \left(L_2(I') - L_2(I)\right)$$
(3)

As we can see, the terms with L_1 are simplified to $-L_p(Z)$ and the model's length because L_1 is simply a summation of individual pattern lengths. The equation of L_2 requires the recomputation of the entire instance matrix' length, which is expensive considering we need to perform it for every candidate, every iteration. However, we can rework the function L_{pp} in Equation (2) by observing that we can isolate the logarithms and generalize them into:

$$\log_G(a,b) = \log \frac{\Gamma(a+b\epsilon)}{\Gamma(b\epsilon)} = \log \Gamma(a+b\epsilon) - \log \Gamma(b\epsilon)$$
 (4)

Which can be used to rework the second part of Equation (3) in such way that the gain equation can be computed in constant time complexity.

$$L_2(I') - L_2(I) = \log_G(U(X), 1) + \log_G(U(Y), 1)$$

$$- \log_G(U(X) - U(Z), 1) - \log_G(U(Y) - U(Z), 1)$$

$$- \log_G(U(Z), 1) + \log_G(|I|, |H|) - \log_G(|I'|, |H'|)$$
(5)

Notice that in some cases the usages of X and Y equal that of Z, which means additional gain is created by removing X and Y from the model.

4.3 Mining patterns

In the second part of the algorithm we select the candidate (X,Y,δ) with the best gain and merge X and Y to form Z, as explained in Section 3.2. We linearly traverse I to replace all instances x and y with relative offset δ by instances of Z. (X,Y,δ) was constructed by looking in the posterior periphery of all x to find Y and δ , which means that Y always comes after X in lexicographical order. The pivot of a pattern is the first element in lexicographical order, therefore pivot(Z) = pivot(X). This means that we can replace all matching x with an instance of Z and all matching y with \cdot . This concludes the baseline version of the algorithm, which is listed in Algorithm 2.

4.4 Improvements

Local search. Given a matrix containing some pattern X, the algorithm can arrive to X in different ways. Exploring these combinatorics can tell us how efficiently the algorithm arrives at X. By definition we know that the fundamental operation is to combine exactly two patterns on each step. Given this, the number of steps in which X can be constructed lies between $\log_2 |X|$ and |X| - 1.

To improve efficiency on large patterns without sacrificing the objectivity of the original heuristics, we add an optional local search. It is a result of the observation that the algorithm generates a large pattern X by adding small elements to a incrementally growing pattern, resulting in a behaviour that approaches |X|-1 steps. Instead of taking all |X|-1 steps to arrive at X, we can try to predict which elements will be added to X and merge them directly. Given that we selected candidate (X,Y,δ) and merged X and Y into Z, now for all m resulting instances $z_i \in z_0, \ldots, z_{m-1}$ we try to find pattern W and relative offset δ such that holds:

$$\forall_{i \in 0...m} \exists_{w} \in ANT(z_i) \cup POST(z_i) \cdot \oslash(w) = W \wedge dist(z_i, w) = \delta$$
 (6)

This yields zero or more candidates (Z, W, δ) , which are then treated as any candidate: candidates with the highest gain are merged first until none exists with positive gain. This essentially means that we run the baseline algorithm only on the peripheries of all z_i , with the condition that the support of the candidates is equal to that of Z. Therefore we only expand Z during local search and we do not create new patterns.

Reusing candidates. We can improve performance by reusing the candidate set and slightly changing the search heuristic of the algorithm. The **Best-N** heuristic selects multiple candidates on each iteration as opposed to the baseline **Best-1** heuristic that only selects a single candidate with the highest gain. Best-N selects candidates in descending order of gain until no candidates with positive gain are left. Furthermore we only consider candidates that are all *disjunct*, because when we merge candidate (X, Y, δ) , remaining candidates with X and/or Y have unknown support and therefore unknown gain.

5 Experiments and Results

To asses the practical performance of the VOUW algorithm, we will primarily use the synthetic dataset generator RIL that was developed specifically for this purpose. RIL utilizes random walks to populate a matrix with patterns of a given size and prevalence, up to a specified density. We fill the remainder of the matrix with uniform noise which allows us to think of the density of patterns as the signal-to-noise ratio (SNR). The objective of the resulting experiment is that we try to find all of the signal (the patterns) and none of the noise. Figure 6 gives an overview of what the generated data looks like, how it is mined and evaluated.

Evaluation Completely random data (noise) is unlikely to be compressed. The SNR tells us how much noise is present in the data and thus conveniently gives us an upper bound of how much compression could be achieved. We use the ground truth SNR versus the resulting compression ratio as a benchmark to tell us how close we are in finding all the structure in the ground truth.

Because the compression ratio alone does not tell us the quality of the results, we also compare the ground truth matrix with the compressed result. In order to do this, we use the notion that elements that have been encoded with singleton patterns, could evidently not be compressed. These elements must therefore be noise. We reconstruct the original matrix from the compressed result, while we omit any singleton patterns. This essentially gives us a matrix of 'positives' (signal) and 'negatives' (noise). By comparing each element with the corresponding element in the ground truth matrix, the *true positives* can be calculated. This subsequently gives us traditional figures for *precision* and *recall*.

Figure 5a plots input SNR versus compression as well as precision/recall for different matrix sizes. We expect the resulting compression ratio to be close to $1-\mathrm{snr}$. In Figure 5b shows that patterns with a low prevalence have a lower probability of being 'detected' by the algorithm as they are more likely to by accidental/noise. We see that increasing the matrix size also increases this threshold. Finally we look at the influence of the two improvements upon the baseline algorithm as described in Section 4.4. Table 2 shows both precision/recall figures as well as wall clock time. In terms of quality, local search improves the results quite substantial in some cases (other experiments have shown that it can dramatically lower the detection threshold as well) while Best-N notably

lowers precision. Both improve speed by an order of magnitude, although the improvements given by Best-N are clearly superior.

6 Conclusion

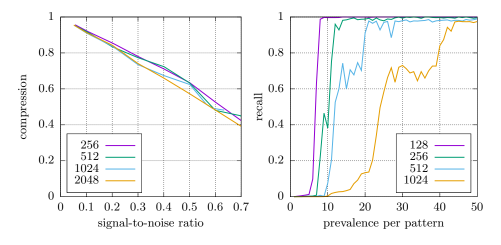
We have introduced geometric pattern mining, the problem of finding recurring structures in matrices or raster-based data, that we had not previously seen in literature. Compared to most pattern mining problems, it adds a layer of encoding geometric relations of data elements. Furthermore the problem can be split into three classes, of which the first class has been the focus of this paper.

We have also presented Vouw, an algorithm based on the MDL principle. The baseline algorithm is capable of generating high-quality results from synthetic datasets produced with our own dataset generated Ril. We have shown that the compression ratio achieved by the algorithm is on-par with what we would expect given the density (SNR) of the input data. By default, the algorithm has a bottleneck in the candidate search, which can be alleviated by the two demonstrated optimizations of the heuristics. It should also be mentioned that more optimizations on part of the implementation are possible (most notably parallelization), but these are outside the scope of this paper.

In future work we would like to generalize the formal definition and notation to n-dimensional data. Moreover, the framework can be expanded to cover all three problem classes. For the algorithm, we believe that it can benefit from more refinement on part of the encoding scheme in an effort to lower the threshold of pattern detection in larger matrices. Lastly we believe that expanding Vouw into similarity measure and clustering such demonstrated by [1], holds a very interesting premise.

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- (a) The influence of SNR in the ground truth
- (b) Prevalence versus recall

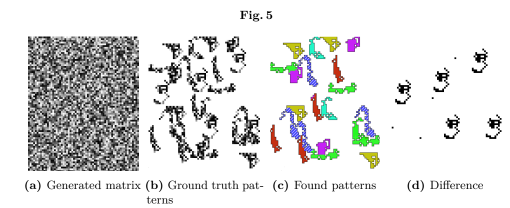


Fig. 6: Example of how synthetic input is generated and evaluated.

Table 2: Influence of optimizations with respect to the baseline algorithm

| | Precision/Recall | | | | Average time | | | |
|------------|------------------|--------------|---------|---------|------------------|--------------|-----------------------------|--------|
| Size | None | Local search | Best-N | Both | None | Local search | Best-N | Both |
| 256^{1} | .98/.98 | .99/.99 | .93/.98 | .95/.99 | 29s | 1s | 2s | 1s |
| 256^{2} | .99/.8 | .99/.88 | .96/.82 | .99/.89 | $2m\ 32s$ | 9s | 5s | 5s |
| 512^{1} | .98/.97 | .99/.99 | .87/.97 | .93/.98 | $5m\ 26s$ | 8s | 20s | 6s |
| 512^{2} | .97/.93 | .99/.99 | .94/.91 | .97/.90 | $26m\ 52s$ | 2m 32s | 24s | 65s |
| 1024^{1} | .97/.98 | .99/.99 | .84/.98 | .92/.96 | 21m 34s | 44s | 37s | 34s |
| 1024^{2} | .98/.98 | .99/.99 | .93/.96 | .98/.97 | $116 \mathrm{m}$ | 7m 31s | $1\mathrm{m}\ 49\mathrm{s}$ | 3m 31s |

 $^{^{1}}$ signal-to-noise ratio of .05, 2 signal-to-noise ratio of .3

A Appendix

A.1 Prequential Plugin-Code

To encode the instance matrix we use the **prequential plug-in code** [?]. The prequential plug-in code is defined for sequences of one item at a time and updates the probability of each item as it is encoded, such that the probability need not be known in advance. It has the favorable property of being asymptotically equal to the optimal code for large sequences. Say we want to encode all elements $I_i \in I$, we define:

$$P_{plugin}(y_i = I_i \mid y^{i-1}) = \frac{|\{y \in y^{i-1} \mid y = I_i\}| + \epsilon}{\sum_{X \in H} |\{y \in y^{i-1} \mid y = X\}| + \epsilon}$$
(7)

Here y_i is the i-th element to be encoded and y^{i-1} is the sequence of elements encoded so far. We initialize the base case (no element has been sent yet) with a pseudocount ϵ , which gives $P_{plugin}(y_1 = I \mid y^0) = \frac{\epsilon}{\epsilon \mid H \mid}$. We pick $\epsilon = 0.5$ as it is used generally with good results.

Let us adapt this principle to the problem of encoding patterns. The first step here is to determine the probability that each unique element (instance of a pattern) in I occurs.

 \triangleright Given a set of instances I, we define usage $(X) = |\{I_i \in I \mid I_i = X\}|$.

From this definition we see that the **usage** of a pattern is a sum of how often it occurs as an instance. We can use this function to simplify things a little by realizing that we actually know the precise number of instances per pattern on the side of the decoder, but not as the decoder. This information can be used to slightly rephrase Equation 2 to be able to encode items in arbitrary order. This produces the length function of the instance matrix I as follows: 1 :

$$L_{pp}(I \mid P_{plugin}) = \sum_{i=1}^{|I|} -\log \frac{|\{y \in y^{i-1} \mid y = I_i\}| + \epsilon}{\sum_{X \in H} |\{y \in y^{i-1} \mid y = X\}| + \epsilon}$$

$$= \sum_{X_i \in h}^{|H|} -\log \prod_{j=0}^{\text{usage}(X_i) - 1} \frac{j + \epsilon}{\sum_{k=1}^{i-1} U(X_k) + j + \epsilon |H|}$$

$$= -\log \frac{\prod_{j=0}^{X_i \in H} \prod_{j=0}^{\text{usage}(X_i)} j + \epsilon}{\prod_{j=0}^{|I| - 1} j + \epsilon |H|}$$

$$= -\sum_{X_i \in h}^{|H|} \left[\log \frac{\Gamma(\text{usage}(X_i) + \epsilon)}{\Gamma(\epsilon)}\right] + \log \frac{\Gamma(|I| + \epsilon |H|)}{\Gamma(\epsilon |H|)}$$
(8)

¹ Here we use the fact that we can interchange sums of logarithms with logarithms of products and that those terms can be moved around freely. Moreover we convert the real-valued product sequences to the Gamma function Γ , which is the factorial function extended to real and complex numbers such that $\Gamma(n) = (n-1)!$.