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# A Constraint on the Number of Distinct Vectors with Application to Localization 

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#### Abstract

This paper introduces a generalization of the nvalue constraint that bounds the number of distinct values taken by a set of variables. The generalized constraint (called nvector) bounds the number of distinct (multi-dimensional) vectors. The first contribution of this paper is to show that this global constraint has a significant role to play with continuous domains, by taking the example of simultaneous localization and map building (SLAM). This type of problem arises in the context of mobile robotics. The second contribution is to prove that enforcing bound consistency on this constraint is NP-complete. A simple contractor (or propagator) is proposed and applied on a real application.


## 1 Introduction

This paper can be viewed as a follow-up of [7] on the application side and [1] on the theoretical side. It proposes a generalization of the nvalue global constraint in the context of a relevant application. The nvalue constraint is satisfied for a set of variables $x^{(1)}, \ldots, x^{(k)}$ and an extra variable $n$ if the cardinality of $\left\{x^{(1)}, \ldots, x^{(k)}\right\}$ (i.e., the number of distinct values) equals to $n$. This constraint appears in problems where the number of resources have to be restricted. The generalization is called nvector and matches exactly the same definition, except that the $x^{(i)}$ are vectors of variables instead of single variables. Of course, all the $x^{(i)}$ must have the same dimension (i.e., the same number of components) and the constraint is that the cardinality of $\left\{x^{(1)}, \ldots, x^{(k)}\right\}$ must be equal to $n$.
We first show that this new global constraint allows a much better modeling of the SLAM (simultaneous localization and map building) problem in mobile robotics. In fact, it allows to make automatic and thus robust a process part of which was performed by hand. Second, we classify the underlying theoretical complexity of the constraint. Finally, a simple algorithm is given and illustrated on a real example.
Since the application context involves continuous domains, we shall soon focus on the continuous case although the definition of nvector does not depend on
the underlying type of domains.

The application is described in Section 2. We first give an informal description of what SLAM is all about. The model is then built step-by-step and its main limitation is discussed. Next, the constraint itself is studied from Section 3 to Section 5 . After providing its semantic (Section 3), the complexity issue is analyzed (Section 4), and a simple contractor is introduced (Section 5). Finally, Section 6 shows how the nvector constraint is used in the SLAM problem modeling, and the improvements obtained are illustrated graphically.
In the rest of the paper, the reader is assumed to have basic knowledge on constraint programming over real domains and interval arithmetics [9]. Domains of real variables are represented by intervals. A Cartesian product of intervals is called a box. Intervals and boxes are surrounded by brackets, e.g., $[x]$. Vectors are in boldface letters. If $\mathbf{x}$ is a set of vectors, $\mathbf{x}^{(i)}$ stands for the $i^{t h}$ vector and $x_{j}^{(i)}$ for the $j^{t h}$ component of the $i^{t h}$ vector. The same convention for indices carries over vector of boxes: $\left[\mathbf{x}^{(i)}\right]$ and $\left[x_{j}^{(i)}\right]$ are respectively the domains of $\mathbf{x}^{(i)}$ and $x_{j}^{(i)}$. Given a mapping $f$, $\operatorname{range}(f,[\mathbf{x}])$ denotes the set-theoretical image of [ $\mathbf{x}]$ by $f$ and $f([\mathbf{x}])$ denotes the image of $[\mathbf{x}]$ by an interval extension of $f$.

## 2 Application Context

The nvector constraint can bring substantial improvements to constraint-based algorithms for solving the simultaneous localization and map building (SLAM) problem. This section describes the principles of the SLAM and provides a constraint model of this application.

### 2.1 Outline of the SLAM problem

The SLAM problem can be described by an autonomous robot moving for a period of time in an unknown environment where the self-location cannot be performed accurately with the help of external equipments (typically, the GPS), and the observation of the environment can only be performed by the robot itself. Examples of such environments include under the sea and the surface of other planets (where the GPS is unavailable). Both limitations are also present indoor where the GPS is often considered unreliable.
In the SLAM problem, we have to compute as precisely as possible a map of the environment (i.e., the position of the detected objects), as well as the trajectory of the robot. The input data is the set of all measures recorded by the robot during the mission from its embedded sensors. These sensors can be divided into two categories: proprioceptive, those that allow to estimate the position of the robot itself (e.g.: a gyroscope) and exteroceptive, those that allows to detect objects of the environment (e.g.: a camera). Note that the positions of the robot at the beginning and the end of the mission are usually known with a good precision. Objects are called landmarks (or seamarks under the sea).

Since nothing can be observed from outside (should it be a landmark or the robot itself), uncertainties of measures get accumulated every time step. The more the mission lasts, the more the robot gets lost. Likewise, detection of landmarks is achieved with less and less accuracy. However, when the robot detects again a landmark that was already placed with a good accuracy on the map, its position can be adjusted from that of the landmark and the whole process of estimation (trajectory and map building) can be refined (see Figure 1). Hence, localization


Fig. 1: A simple SLAM example. The robot is a mouse and landmarks are trees. Uncertainties on the robot position (resp. trees) are represented by blank (resp. hatched) ellipsis. (a) At initial time, the robot knows where it is. (b) The uncertainty on the robot position increases while it moves. When a second tree is detected, there is a significant uncertainty on its real position. (c) The first tree is met again and used to adjust the position of the robot. (d) With a backward computation of the trajectory, uncertainties on the previous positions and observations are reduced.
and map building are two connected goals. This interdependence is one of the reason that makes traditional probabilistic approaches inadequate. In contrast, it makes no difficulty in the constraint programming framework, as shown below.

### 2.2 Basic Constraint Model

Let us now focus on a basic modeling of the SLAM. Many details on a real experiment (description of the robot, experimental setup, full constraint model, etc.) can be found in [7] and [8] that deal with SLAM in a submarine context.

The SLAM problem is cast into a CSP as follows. First, the motion of the autonomous robot obeys a differential equation: $\mathbf{p}^{\prime}(t)=\mathbf{f}(\mathbf{u}(t))$, where $\mathbf{p}(t)$ is the position of the robot in space, $\mathbf{u}(t)$ a vector of $m$ inputs (speed, rotation angles, etc.) and $\mathbf{f}$ a mapping from $\mathbb{R}^{m}$ to $\mathbb{R}^{3}$. This equation can be cast into a CSP using a classical interval variant of the Euler method. Details can be found in [7] and [8]. The discretization introduces a set of $(N+1)$ variables $\mathbf{p}^{(0)}, \ldots$, $\mathbf{p}^{(N)}$ where $\delta_{t}$ is the time lapse between to measures and $N \delta_{t}$ the total duration of the mission. These variables represent a discretization of the trajectory $\mathbf{p}$, i.e.,

$$
\begin{equation*}
\forall i, 0 \leq i \leq N, \quad \mathbf{p}^{(i)}=\mathbf{p}\left(t_{0}+i \delta_{t}\right) \tag{1}
\end{equation*}
$$

has to be fulfilled. The discretization also introduces $N$ constraints.
Thus, the CSP provides a rigorous enclosure of the trajectory, i.e., for every possible input $\mathbf{u}(t)$ there exists a feasible tuple $\left(\mathbf{p}^{(0)}, \ldots, \mathbf{p}^{(N)}\right)$ such that the trajectory $\mathbf{p}$ corresponding to the inputs satisfies (1).

### 2.3 Introducing Detections

Now that the motion of the vehicle has been cast into a CSP, let us take into account detections. In the mission, $n$ landmarks have to be localized. Their coordinates will be denoted by $\mathbf{o}^{(1)}, \ldots, \mathbf{o}^{(n)}$. Once the mission is over, a human operator scans the waterfall of images provided by exteroceptive sensors. When a group of pixels are suspected to correspond to a landmark, a box encompassing the corresponding area is entered as a potential detection. The position of a pixel on the image can be directly translated into a distance between the landmark and the robot. First, the detection time $\tau(i)$ (i.e., the number of time steps since $t_{0}$ ) and the distance $r_{i}$ are determined. Then, the landmark number $\sigma(i)$ is identified which amount to match detections with each others. Finally, a distance constraint $\operatorname{dist}\left(\mathbf{p}^{(\tau(i))}, \mathbf{o}^{(\sigma(i))}\right)=r_{i}$ between the landmark and the robot is added into the model. Therefore, in [7], the model was augmented as follows:

$$
\left(\mathcal{P}^{\prime}\right)\left\{\begin{array}{l}
\text { additional variables: } \\
\mathbf{o}^{(1)} \in\left[\mathbf{o}^{(1)}\right], \ldots, \mathbf{o}^{(n)} \in\left[\mathbf{o}^{(n)}\right] \\
\text { domains: } \\
{\left[\mathbf{o}^{(1)}\right]:=(-\infty,+\infty), \ldots,\left[\mathbf{o}^{(n)}\right]:=(-\infty,+\infty)} \\
\operatorname{additional} \text { constraints: } \\
\operatorname{dist}\left(\mathbf{p}^{(\tau(i))}, \mathbf{o}^{(\sigma(i))}\right)=r_{i} \quad(i=1 . . k)
\end{array}\right.
$$

Humans are subject to three types of mistakes: (1) Omission, a landmark on the waterfall is missed by the operator; (2) Illusion, the operator adds a detection where there is no landmark; (3) Mismatching, the operator makes a wrong identification (do not match detections properly). On the one hand, the two first types of mistakes have been experimentally proven as irrelevant ${ }^{1}$. On the other

[^0]hand, if identifying the type of a landmark is fairly easy, recognizing one particular landmark is very hard. In other words, the main difficulty in the operator's task is matching landmarks with each others. The bad news is that mismatching makes the model inconsistent. Hence, the third type of mistakes is critical and requires a lot of energy to be avoided. Up to now, in the experiments made in $[7,8]$, matching was simply performed using a priori knowledge of seamark positions.

### 2.4 Our contribution

The ambition of our work is simply to skip the operator's matching phase. The idea is to use the knowledge on the number of landmarks to make this matching automatically by propagation. This is a realistic approach. In all the missions performed with submarine robots, a set of beacons is dropped into the sea to make the SLAM possible. The positions of the beacons at the bottom is not known because of currents inside water (consider also that in hostile areas they may be dropped by plane) but their cardinality is.

## 3 The Number of Distinct Vectors Constraint

Consider $k$ vectors of variables $\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}$ of equal dimension (which is 2 or 3 in practice) and an integer-valued variable $n$. The constraint nvector ( $n$, $\left.\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}\right)$ holds if there is $n$ distinct vectors between $\mathbf{x}^{(1)}$ and $\mathbf{x}^{(k)}$ which can be written:

$$
\text { nvector }\left(n,\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}\right) \Longleftrightarrow\left|\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}\right|=n
$$

where $\left|\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}\right|$ stands for the cardinality of the set of "values" taken by $\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}$. An equivalent definition that better conform to the intuition behind is that the number of distinct "values" in $\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}$ equals to $n$.
A family of constraints can be derived from nvector in the same way as for nvalue [2]. The constraint more specifically considered in this paper is atmost_ nvector that bounds the number of distinct vectors:

$$
\operatorname{atmost\_ nvector}\left(n,\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}\right) \Longleftrightarrow\left|\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}\right| \leq n .
$$

It must be pointed out that, with continuous domains, the constraints nvector and atmost_nvector are operationally equivalent. Indeed, excepted in very particular situations where some domains are degenerated intervals (reduced to single points), variables can always take an infinity of possible values. This means that any set of $k$ non-degenerated boxes always share at least $n$ values. In the nvector constraint, only the upper bound ("at most $n$ ") can actually filters. This remark generalizes the fact that all_diff is always satisfied with nondegenerated intervals. Figure 2a shows an example with $n=2$ (the variable is ground) and $k=5$ in two dimensions. Figure 2b shows the corresponding result of a bound consistency filtering.


Fig. 2: (a) The atmost_nvector constraint with $n=2$ and $k=5$. The domains of $\mathbf{x}^{(i)}$ is the cross product of two intervals $\left[\mathbf{x}^{(i)}\right]=\left[x_{1}^{(i)}\right] \times\left[x_{2}^{(i)}\right]$. The constraint is satisfiable because there exists two vectors $\mathbf{v}^{(1)}$ and $\mathbf{v}^{(2)}$ such that $\forall i, 1 \leq i \leq n,\left[\mathbf{x}^{(i)}\right] \cap\left\{\mathbf{v}^{(1)}, \mathbf{v}^{(2)}\right\} \neq$ $\emptyset$. The set of all such pairs $\left(\mathbf{v}^{(1)}, \mathbf{v}^{(2)}\right)$ is represented by the two little dashed boxes. (b) Result of the bound consistency with respect to the atmost_nvector constraint. The domain of $\left[\mathbf{x}^{(4)}\right]$ encloses the two little rectangles.

## 4 Operational Complexity

Enforcing generalized arc consistency (GAC) for the atmost-nvalue constraint is NP-hard [3]. Worse, simply computing the minimum number of distinct values in a set of domains is NP-hard [2]. However, when domains are intervals, these problems are polynomial $[1,6]$. The question under interest is to know if atmost-nvector is still a tractable problem when domains are boxes (vector of intervals), i.e., when the dimension is greater than 1 . We will show that it is not. Focusing on this type of domains is justified because continuous constraints are always handled with intervals. This also implies that bound consistency is the only acceptable form of filtering that can be applied with continuous domains. ${ }^{2}$ We shall restrict ourselves to the calculation of the minimum number of distinct values (also called the minimum cardinality) and to boxes of dimension 2, i.e., rectangles. As noticed in [2], the minimum number of distinct values shared by a set of rectangles is also the cardinality of the minimum hitting set. A hitting set is a collection of points that intersects each rectangle. In the two following subsections, we consider the equivalent problem of finding the cardinality of the minimum clique partition of a rectangle graph.

### 4.1 Rectangle Graphs and Clique Partitions

A rectangle graph is a $n$-vertices graph $G_{r}=\left(V_{r}, E_{r}\right)$ that can be extracted from $n$ axis-aligned rectangles in the plane by (1) creating a vertex for each

[^1]a rectangle, and (2) adding an edge when there is a non-empty intersection between two rectangles. Figure 3 depicts the two views of a rectangle graph.
A clique partition of a graph is a collection of complete subgraphs that partition all the vertices. A minimal clique partition is a clique partition with the smallest cardinality (i.e., with the smallest number of subgraphs). Since any set of pairwise intersecting rectangles intersect all mutually (by Helly's theorem), a $k$-clique in a rectangle graph represents the intersection of $k$ rectangles in the geometrical view. For instance, the 3-clique $S 1$ of $G_{r}$ in Figure 3a represents the intersection of three rectangles $(R 4, R 5, R 6)$ in Figure 3b, depicted by the black rectangle $S 1$. As a consequence, looking for the minimum hitting set or the


Fig. 3: A rectangle graph and its geometrical representation (axis-aligned rectangles).
minimum clique partition are indeed equivalent problems for rectangle graphs. The final problem under consideration can then be formulated as follows:

## Rectangle Clique Partition (RCP)

- Instance: A rectangle graph $G_{r}=\left(V_{r}, E_{r}\right)$ given in the form of $\left|V_{r}\right|$ axisaligned rectangles in the plane and $k \leq\left|V_{r}\right|$.
- Question: Can $V_{r}$ be partition into $k$ disjoint sets $V_{1}, \ldots, V_{k}$ such that $\forall i, 1 \leq i \leq k$ the subgraph induced by $V_{i}$ is a complete graph?

Proposition 1. $R C P$ is NP-complete.
The fact that RCP belongs to NP is easy to prove: the given of a $k$-partition is a certificate that can be checked in polynomial time. The rest of the proof, i.e., the reduction of a NP-complete problem to RCP is given below. Note that the transformation below is inspired from that in [10] but contains fundamental differences.

### 4.2 Building a Rectangle Graph from a Cubic Planar Graph

The problem we will reduce to RCP involves planar graphs which require the introduction of extra vocabulary. An embedding of a graph $G$ on the plane is a
representation of $G$ (see Figure 4) in which points are associated to vertices and arcs are associated to edges in such a way:

- the endpoints of the arc associated to an edge $e$ are the points associated to the end vertices of $e$,
- no arcs include points associated to other vertices,
- two arcs never intersect at a point which is interior to either of the arcs.

A planar graph (Figure 4a) is a graph which admits an embedding on the plane (Figure 4b), and a cubic graph is a 3 -regular graph, i.e., a graph in which every vertex has 3 incident edges. A rectilinear embedding is an embedding where every arc is a broken line formed by horizontal and vertical segments (Figure 4c).


Fig. 4: A (cubic) planar graph, one of its embedding on the plane, and one of its rectilinear embedding.

We are now in position to discuss about the transformation itself. Consider a planar cubic graph $G_{P}$. First, Tammassia and al. gave a polytime algorithm in [11] for computing a 5 -rectilinear embedding of a planar graph, i.e., a rectilinear embedding where each broken line is formed by 5 segments. This is illustrated by Figure 5 . Second, this embedding can be transformed into a rectangle graph


Fig. 5: A 5-rectilinear embedding of the cubic planar graph given in Figure 4 (segments are delineated by arrows).
using the patterns depicted on Figures 6a and 6b:

- Every segment of the 5-rectilinear embedding of $G_{p}$ is replaced by a rectangle such that if two segments do not intersect in the 5 -rectilinear embedding, the corresponding rectangles do not intersect (should they be flat).
- For segments having a vertex at a common endpoint, the corresponding rectangles intersect all mutually in the neighborhood of this vertex (Figure 6a). Remember that the case of a leaf node is irrelevant because $G_{p}$ is cubic.
- For segments having a bend in common, the rectangles are disjoint.
- An extra rectangle is added in the neighborhood of each bend. This rectangle intersects the two rectangles associated to the segments (Figure 6b). The three rectangles cannot intersect all mutually due to the previous point.

(a) Transformation of segments at a common endpoint.

(b) Transformation of segments having a bend in common.

Fig. 6: Atomic operations to transform the 5-rectilinear embedding of a cubic planar graph (left) into the geometrical view (middle) of a rectangle graph (right).

### 4.3 Reduction From Cubic Planar Vertex Cover

Consider the well-known vertex cover problem. This problem remains NP-Complete even for cubic planar graphs [12]. It is formally stated as follows:

## Cubic Planar Vertex Cover (CPVC)

- Instance: A cubic planar graph $G_{p}=\left(V_{p}, E_{p}\right)$ and $k \leq\left|V_{p}\right|$.
- Question: Is there a subset $V^{\prime} \subseteq V_{p}$ with $\left|V^{\prime}\right| \leq k$ such each edge of $E_{p}$ has at least one of its extremity in $V^{\prime}$ ?

Lemma 1. Let $G_{p}$ be a n-vertex m-edges cubic planar graph and an integer value $k \leq n$. Let $G_{r}$ be the rectangle graph obtained by the transformation of §4.2. The answer to CPVC with $\left(G_{p}, k\right)$ is yes iff the answer to $R C P$ with $\left(G_{r}, k+\right.$ $4 \times m$ ) is yes.

Proof. This proof is illustrated in Figure 7.
Forward implication: Assume a $n$-vertex $m$-edges cubic planar graph $G_{p}$ has a vertex cover $V^{\prime}$ of cardinality $k$. We shall build a partition $\mathcal{P}$ of $G_{r}$, initially
empty. To each edge $e=\left(v_{i}, v_{j}\right)$ of $E_{p}$ corresponds a 2-degree chain $c_{e}$ in $G_{r}$, with exactly nine vertices, from a vertex of the 3 -clique $p_{i}$ (associated to $v_{i}$ ) to a vertex of the 3 -clique $p_{j}$ (associated to $v_{j}$ ).
First, for every $v_{i} \in V^{\prime}$, add the 3 -clique $p_{i}$ in $\mathcal{P}$. Since $V^{\prime}$ is a covering set, every chain has now one of its extreme vertex inside a clique of $\mathcal{P}$. The 8 other remaining vertices of the chain can then easily be partitioned into 4 additional cliques. Add these cliques to $\mathcal{P}$.
Once all these additional cliques are added to $\mathcal{P}$, the latter is a partition of $G_{r}$ whose size is $k+4 \times m$.

(a) Edge of $G_{p}$.

(b) Rectilinear embedding view.

(c) Corresponding chain (subgraph of $G_{r}$ ).

Fig. 7: Transforming an edge of the cubic planar graph $G_{P}$ into, first, its rectilinear view, second, a rectangle graph $G_{r}$.

Backward implication: Assume $G_{r}$ can be partition into $k+4 \times m$ cliques and let $\mathcal{P}$ be such a partition. We shall call a free clique a clique that only contain vertices of the same given chain. Similarly, a shared clique is a clique involving (extreme) vertices of at least two different chains.
Every chain contains 9 vertices. One can easily see that it requires at least 5 cliques to be partitioned, 4 of which are free. For every chain $c$, remove 4 free cliques of $c$ from $\mathcal{P}$. Then there is still (at least) one clique left in $\mathcal{P}$ that involves a vertex of $c$. When this process is done, the number of remaining cliques in $\mathcal{P}$ is $k+4 \times m-4 \times m=k$.
Now, for every clique $\mathcal{C}$ of $\mathcal{P}$ : if $\mathcal{C}$ is shared, put the corresponding vertex of $G_{P}$ in $V^{\prime}$. If $\mathcal{C}$ is free, consider the edge $e$ of $G_{P}$ associated to the chain and put anyone of the two endpoint vertices of $e$ into $V^{\prime}$. We have $\left|V^{\prime}\right|=k$ and since every chain has a vertex in the remaining cliques of $\mathcal{P}$, every edge of $G_{P}$ is covered by $V^{\prime}$.

## 5 A Very First Polytime Contractor

We shall now propose a simple algorithm for computing bound consistency with respect to atmost_nvector. Let us denote by $\operatorname{dim}$ the dimension of the vectors.

The contractor is derived from the following implication:
atmost_nvector $\left(n,\left\{\mathbf{x}^{(1)}, \ldots, \mathbf{x}^{(k)}\right\}\right) \Longrightarrow$
$\operatorname{atmost\_ nvalue}\left(n,\left\{x_{1}^{(1)}, \ldots, x_{1}^{(k)}\right\}\right) \wedge \ldots \wedge \operatorname{atmost\_ nvalue}\left(n,\left\{x_{\text {dim }}^{(1)}, \ldots, x_{\text {dim }}^{(k)}\right\}\right)$.
Therefore, applying a contractor for atmost_nvalue with the projections of the boxes onto each dimension in turn gives a contractor for atmost_nvector. Since a contractor for atmost_nvalue enforcing GAC (hence, bound consistency) when domains are intervals already exists [1], we are done. This is achieved in $O(\operatorname{dim} \times$ $k \log (k))$. Although the purpose of this paper is not to describe an efficient contractor, we shall introduce right away a first (but significant) improvement of our naive algorithm.


Fig. 8: Illustration of what the at most_nvalue algorithm yields. Here, $k=7, \operatorname{dim}=2$, $n=3$ and the algorithm is run for the first projection (i.e., onto the horizontal axis). Three groups, represented by gray ellipsis, are identified, namely $G_{1}=\left\{x_{1}^{(1)}, x_{1}^{(2)}\right\}$, $G_{2}=\left\{x_{1}^{(4)}, x_{1}^{(5)}\right\}$ and $G_{3}=\left\{x_{1}^{(7)}\right\}$. The variables of the same group are proven to be all equal (otherwise, the constraint is violated). E.g., $x_{1}^{(1)}$ and $x_{1}^{(2)}$ must satisfy $x_{1}^{(1)}=x_{1}^{(2)}$. Domains for the variables of a given group can be replaced by the corresponding kernel (e.g., $\left[x_{1}^{(1)}\right]$ and $\left[x_{1}^{(2)}\right]$ can be set to $\left.\left[G_{1}\right]\right)$. Notice that the kernels are all disjoint. In our suggested improvement, the whole boxes are intersected instead of the first components only. Hence the domains of $\left[\mathbf{x}^{(1)}\right]$ and $\left[\mathbf{x}^{(2)}\right]$ are intersected, which gives one of the hatched rectangles. The other hatched rectangle is $\left[\mathbf{x}^{(4)}\right] \cap\left[\mathbf{x}^{(5)}\right]$.

The reader must simply admit that the contractor of atmost_nvalue $\left(n,\left\{x_{i}^{(1)}\right.\right.$, $\left.\ldots, x_{i}^{(k)}\right\}$ ) works in two steps (see Figure 8). First, it builds groups of variables. If the number of groups turns to be $n$, it is then proven that for all $x_{i}^{(j)}$ and $x_{i}^{(l)}$ that belong to the same group, $x_{i}^{(j)}$ and $x_{i}^{(l)}$ must satisfy $x_{i}^{(j)}=x_{i}^{(l)}$. Hence, the domains of all the variables of the same group $G$ can be shrunk to their
common intersection denoted by $[G]$. Furthermore, this process results in a set of $n$ disjoint intervals $\left[G_{1}\right], \ldots,\left[G_{n}\right]$ that we will call kernels.
In the favorable case of $n$ groups, the last step can be improved. If two variables $x_{i}^{(j)}$ and $x_{i}^{(l)}$ belong to the same group $G$, the whole vector variables $\mathbf{x}^{(j)}$ and $\mathbf{x}^{(l)}$ are actually constrained to be equal. Indeed, assume that $\mathbf{x}^{(j)}$ and $\mathbf{x}^{(l)}$ could take two different vectors. The $k-2$ other variables necessarily share $n-1$ different vectors because their $i^{\text {th }}$ components must "hit" the $n-1$ (disjoint) kernels [ $\left.G^{\prime}\right]$ with $G^{\prime} \neq G$. This means that the overall number of distinct vectors is at least $(n-1)+2>n$.
Hence, the algorithm of atmost_nvalue can be modified to intersect boxes (instead of just one component). This multiplies the complexity by $\operatorname{dim}$ (which is small in practice).

## 6 Experimental Evaluation: the SLAM Problem

This section shows how the atmost_nvector constraint allows to improve the modeling and resolution of the SLAM problem. We propose an extension of the original model given in Section 2, and provide a graphical validation.
We introduce $k 3$-dimensional variables $\mathbf{d}^{(1)}, \ldots, \mathbf{d}^{(k)}$ related to all the detections. A distance constraint involving each of the latters is added into the model, as well as a nvector constraint capturing the fact that only $n$ landmarks exist.

$$
\left(\mathcal{P}^{\prime \prime}\right)\left\{\begin{array}{l}
\text { additional variables: } \\
\mathbf{d}^{(1)} \in\left[\mathbf{d}^{(1)}\right], \ldots, \mathbf{d}^{(k)} \in\left[\mathbf{d}^{(k)}\right] \\
\text { domains: } \\
{\left[\mathbf{d}^{(1)}\right]:=(-\infty,+\infty), \ldots,\left[\mathbf{d}^{(k)}\right]:=(-\infty,+\infty)} \\
\text { additional constraints: } \\
\operatorname{dist}\left(\mathbf{p}^{\tau(i)}, \mathbf{d}^{(i)}\right)=r_{i} \quad(i=1 . . k) \\
\text { atmost }-\operatorname{nvector}\left(n,\left\{\mathbf{d}^{(1)}, \ldots, \mathbf{d}^{(k)}\right\}\right)
\end{array}\right.
$$

The introduction of the atmost-nvector constraint has provided the expected results. Let us first explain how the benefits of this constraint can be quantified. In the context of differential equations, we are dealing with a very large number of variables and a sparse system with an identified structure. Moreover, the system is subject to many uncertainties. Under these conditions, the quality of the result is more relevant than the computation time, as we justify now.

No choice point. In applications, the number of variables can be huge (more than 1140000 in $[7,8]$ in order to represent the three coordinates of the robot, the Euler angles, the speed, the altitude and the components of the rotation matrix at each time step). This prevents us for making choice points. Note that the situation is even worse since the solution set is not a thin trajectory (formed by isolated points). The numerous uncertainties make the best solution ever a thick
beam of trajectories (as depicted in Figure 9). Continuums of solutions usually dissuade one from making choice points, even for much smaller problems.

Handcrafted propagation. The size of our problem also prevents us for using a general-purpose AC3-like propagation algorithm, since building an adjacency matrix for instance would simply require too much memory. Furthermore, we have a precise knowledge of how the network of constraints is structured. Indeed, the discretization of the motion yields a single (long) chain of constraints and cycles in the network only appear with detection constraints. An appropriate strategy is then to base propagation on the detection constraints, which are much fewer. Every time a detection reduces significantly the domain of a position $\mathbf{p}^{(i)}$, the motion constraints are propagated forward (from $\mathbf{p}^{(i)}$ to $\mathbf{p}^{(N)}$ ) and backward (from $\mathbf{p}^{(i)}$ downto $\mathbf{p}^{(0)}$ ). In a nutshell, propagation in this kind of problems is guided by the physics.

Irrelevance of computation time. When the application is run, it takes a couple of seconds to load data of the sensors (which amount to initialize the domains of variables representing input vectors) and to precalculate expressions (e.g., rotation matrices). With or without the nvector constraint, propagation takes a negligible time (less than 1\%) in comparison to this initialization. Therefore, focusing on computation time is not very informative.

Quality of the result. Our contribution is to make propagation for the SLAM problem automatic whereas part of it was performed by a human operator so far. This is a result in itself. However, one may wonder which between the automatic matching and the operator's is most competitive. This, of course, is hard to evaluate a priori. In the experiment of $[7,8]$, both have provided the optimal matching. The question that still remains is to know the extent to which our matching (the one provided by the algorithm in Section 5) improves the "quality" of the result, i.e., the accuracy of the trajectory.
The idea was to make the robot looping around the same initial point so that many detections would intersect. ${ }^{3}$ For this purpose, we have controlled the robot with a classical feedback loop which gives the expected cycloidal trajectory.
Four landmarks have been placed in the environment and we have basically considered that a landmark is detected everytime the distance between the robot and itself reaches a local minimum (if less than a reasonable threshold). The estimation of the landmark position is then calculated from this distance (with an additional noise) and a very rough initial approximation.
Figure 9 illustrates the effect of automatic matching on the estimation of the trajectory and the positions of the landmarks. All the results have been obtained in a couple of seconds by the Quimper system [5].

[^2]

Fig. 9: Comparing SLAM with the atmost-nvector contractor (Improved SLAM) and without (Basic SLAM). The trajectory is represented by gray filled boxes, detections by thick-border rectangles and landmarks by the little black squares. Figure 9(d) depicts the fixpoint of all the contractors: the four landmarks are very well localized and the trajectory is much thiner (after 4000 iterations, the largest diameter is 10 times less than in Figure 9(c)). Our algorithm has contracted many detections to optimal boxes around landmarks. We can also observe the weakness of our algorithm which has poorly reduced boxes whose projections on both axis encompass two projections of landmarks. This however exclude a detection which really encloses two landmarks: in this case, either there is a real ambiguity or it is the propagation to blame.

## 7 Conclusion

A somewhat natural generalization of the nvalue constraint, called nvector has been proposed. The nvector constraint can help modeling and solving many
localization problems where a bound on the number of landmarks to localized is known. This has been illustrated on the SLAM problem and applied on a real experiment. We have also analyzed the complexity of this global constraint and given a simple contractor.
The benefit of this constraint in terms of modeling has a direct impact on the way data of the experiments have to be processed. Indeed, the constraint allows to avoid requiring someone that matches landmarks by hand. Hence, it reduces considerably the amount of work and the probability of mistake this operation entails.
The field of application is not restricted to the SLAM problem. Ongoing works show that the nvector is as crucial for the passive location of vehicles using TDOA (time difference of arrival) in signal processing. All these problems involve real variables. Hence, as a side contribution, this paper also offsets the lack of activity about global continuous constraints.
Future works include the design of more sophisticated contractors with benchmarking. The nvector constraint also leads up to the study of other global constraints. As soon as several estimations of the same landmark position are matched by nvector, this position satisfies indeed a global constraint (namely, the intersection of several spheres if estimations result from distance equations).

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[^0]:    ${ }^{1}$ The overall accuracy may suffer from a lack of detections but the consistency of the model is always maintained (at any time, many landmarks are anyway out of the scope of the sensors and somehow "missed"). Besides, perception is based on very specific visual patterns that rule out any confusion with elements of the environment.

[^1]:    ${ }^{2}$ Allowing gaps inside intervals and applying GAC filtering leads to unacceptable space complexity [4].

[^2]:    ${ }^{3}$ The experiment of $[7,8]$ was not appropriate for this illustration because matching seamarks was actually too easy ( 6 seamarks and a rectangle graph of detections with 6 strongly connected components).

