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Speeding up Derivative Configuration from Product Platforms

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Abstract: To compete in the global marketplace, manufacturers try to differentiate their products by focusing on individual customer needs. Fulfilling this goal requires that companies shift from mass production to mass customization. Under this approach, a generic architecture, named product platform, is designed to support the derivation of customized products through a configuration process that determines which components the product comprises. When a customer configures a derivative, typically not every combination of available components is valid. To guarantee that all dependencies and incompatibilities among the derivative constituent components are satisfied, automated configurators are used. Flexible product platforms provide a big number of interrelated components, and so, the configuration of all, but trivial, derivatives involves considerable effort to select which components the derivative should include. Our approach alleviates that effort by speeding up the derivative configuration using a heuristic based on the information theory concept of entropy.

Keywords: entropy-based heuristic; mass customization; product configuration; software product line; feature model

1. Introduction

To increase variety, improve customer satisfaction, reduce lead-times, and shorten costs, many companies have shifted from mass production to mass customization [1]. This shift of paradigm enriches

the mass production economies of scale with custom manufacturing flexibility by developing families of related products instead of single products. From this perspective, designing a product family is the process of capturing and modeling multiple product variants to satisfy different market niches. A generic architecture, named product platform, is designed to support the creation of customized products called derivatives.

Product platforms usually support a high quantity of derivatives. For instance, the number of derivatives for product platforms in the automotive industry may range from 10^3 for the smallest Peugeot and Nissan car models, to 10^{16} or 10^{21} for the BMW 3-Series and Mercedes C-Class, respectively [2]. To achieve that flexibility, a number of configuration options are available. For example, the Peugeot 206 and Mercedes C-Class car models have 86 and 389 customer selectable options, respectively. Typically, not all option combinations are valid. There may be option incompatibilities (e.g., “manual transmissions are not compatible with V8 engines”), option dependencies (e.g., “sport cars require manual gearbox”), *etc.* Configuring a valid derivative implies ensuring that all constraints between its constituent components are satisfied. Checking by hand those constraints is infeasible for all, but the most trivial product platforms, so derivative configuration is usually assisted by automated configurators [3]. Some examples of commercial configurators are *Configit* [4], *SAP Product Configurator* [5], *Oracle Configurator* [6], *etc.* In addition, many automotive companies have their own configurators. For instance, *Volvo* uses *KOLA*, *Scania* uses *SPECTRA*, *Mercedes* uses *SMARAGD*, *etc.*

Our work enriches existing configurators by reducing the number of steps required to configure a valid derivative. It takes advantage of the fact that, due to the component composition constraints, some decisions may be automatically derived from other decisions previously made. Therefore, the order in which decisions are made has a strong influence on the number of decisions required to complete a derivative. For instance, given the constraint “sport cars require manual gearbox”, a customer might configure a sport car using two decision orderings: one requiring two steps (*i.e.*, Step 1, select “manual gearbox”, and Step 2, select “sport car”) or another one using just a single step (*i.e.*, select “sport car”, so that the decision to select “manual gearbox” is implicitly made).

As van Nimwegen *et al.* [7] note, customers sometimes prefer to first answer questions that are important to them, or easy to answer, before being led through the remaining questions [7]. In this sense, our approach respects customer preferences. Instead of imposing a fixed ordering, it suggests orderings dynamically, reacting to the customer decisions. In particular, the process to get a derivative is performed in successive steps. In each step, the customer gets a question ranking, selects one of the questions and answers it. In the next step, the question ranking is readjusted to account for the customer’s answer. The computation of the ranking is grounded on the information theory concept of entropy, which was introduced by Shannon [8] and measures the average uncertainty of a random variable.

At the first configuration step, the uncertainty is total. With no information at all, the configurator cannot figure out which derivative the customer desires. As the process advances, configuration options are eliminated according to the customer decisions, and so, the information about the final configuration increases (*i.e.*, the set of included/excluded components grows). Consequently, the entropy decreases. When the derivative is totally configured, there is no uncertainty, and the entropy is zero.

As we will see, not only our approach, but also Mazo *et al.*'s Heuristic 3 [9] and Chen *et al.*'s approach [10] require computing the probabilities of all variables in a Boolean formula. The usual way to perform such task is calling repeatedly a logic engine, e.g., a SATsolver or a binary decision diagram (BDD) library, one time for each variable [11]. Unfortunately, this approach has a high computational cost and, thus, imposes long response times, hindering customer-configurator interactivity. To overcome such a problem, this paper proposes an algorithm that computes efficiently variable probabilities using BDDs. Since more complex logics than the propositional one, which include integer arithmetic, transitive closure, *etc.*, can be reduced to Boolean functions [12,13] and, thus, encoded as BDDs, our algorithm is general enough to support most configuration model notations.

The validity of our approach has been tested on two benchmarks widely used by the configuration and software product line communities: the Renault Megane platform provided by the car manufacturing company Renault DVI [14] and the Electronic Shopping case study [17]. Results show that our approach requires less configuration steps than related work, and that our BDD algorithm gets short response times, supporting this way not only our approach but also other methods proposed in related work.

The remainder of this paper is structured as follows. Section 2 presents the running example that we will use to motivate and illustrate our work. Section 3 summarizes related work to our approach. Section 4 introduces the concept of entropy and describes how to compute it from a configuration model. Later, our entropy-driven approach is described in detail. Section 5 reports the experimental validation of our approach. Finally, Section 6 outlines the conclusions of our work.

2. Motivational Example

This section illustrates the problem our approach tackles using an example provided by [18], where derivatives are cars with different automated driving capabilities.

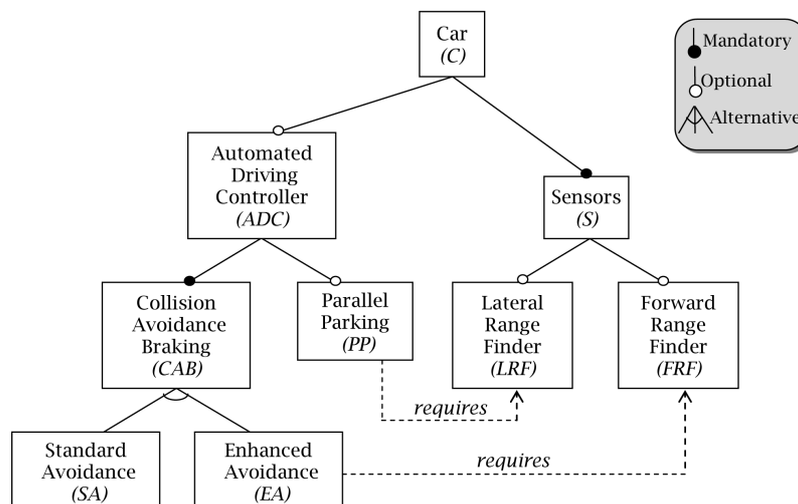
To model the configurable options of a product family, a number of different notations are available. For instance, feature diagrams (FDs) [19], decision diagrams [20], the Configit language, the SAP Product Configurator language, the Oracle Configurator language, *etc.* Interestingly, most of those notations are semantically equivalent [21,22]. In fact, automated configurators instead of processing configuration models directly usually translate them into a propositional logic representation, such as a logic formula in conjunctive normal form, a BDD, *etc.* That logic representation is then processed using off-the-shelf tools, such as SAT solvers, BDD engines, *etc.* (see Section 4.1.2. for an explanation of the configuration model to logic translation). The input to our approach is the logic representation of the configuration model, so it is independent of the original notation used to specify the model.

To show what a configuration model looks like, please refer to Figure 1, which models our running example as an FD (a hierarchically arranged set of features with different relations among them). This paper follows the generic semantics for FDs given by Schobbens *et al.* [22]. Figure 1 includes three kinds of hierarchical relations:

- optional, denoted by simple edges ending with an empty circle; e.g., cars may (or may not) include an automated driving controller (ADC).
- mandatory, denoted by simple edges ending with a filled circle; e.g., if a car has an ADC, it must include some kind of collision avoidance braking (CAB).

- alternative, denoted by edges connected by an arc; e.g., standard avoidance (SA) and enhanced avoidance (EA) are the mutually exclusive options for collision avoidance braking (CAB).

Figure 1. Feature diagram (FD) for car automated driving capabilities.



To manage the complexity of modeling the similarities and differences among the derivatives of a product family, the FD notation follows a divide and conquer strategy. Derivative variabilities are modeled by progressively decomposing complicated and abstract features into simpler ones, until elemental features, which are directly implemented by physical components, are reached. The hierarchical structure of an FD graphically depicts such conceptual decomposition. From here on, derivatives will be expressed enumerating the final components they include, *i.e.*, using references to the terminal nodes of the FD. For example, {PP, LRF, FRF, \neg SA, EA} expresses the configuration of a car with components PP, LRF, FRF, EA and without SA.

The FD notation supports narrowing the configuration space by adding additional cross-tree constraints. For instance, Figure 1 represents as “PP $\xrightarrow{\text{requires}}$ LRF” the fact that cars with parallel parking need to include the lateral range finder component. Thus, a car derivative with components {PP, \neg LRF, \neg FRF, SA, \neg EA} complies with the FD relations, but is not valid because it violates the constraint “PP $\xrightarrow{\text{requires}}$ LRF”.

For a configuration model with n options and no component interdependencies, the number of possible configurations is 2^n . Due to the feature relations and additional cross-tree constraints, the number of valid configurations in the example is reduced from $2^5 = 32$ to the 13 ones summarized in Table 1.

To configure a car, the decision-maker needs to answer a sequence of questions. For example, the sequence (1) *is EA in the configuration?* no, (2) *FRF?* no, (3) *LRF?* yes, (4) *PP?* yes configures Car 11 in Table 1. Current automated configurators guarantee the derivation of valid products ensuring the satisfaction of all model constraints. When the first question is answered, the configurator deduces that the car being configured necessarily includes SA (otherwise, the alternative relation between EA and SA would not hold). This way, the configurator is indirectly saving the decision-maker from answering the irrelevant question, “is SA in the configuration?”

The goal of our work is to make the most of the configuration model constraints going beyond current configurators to minimize the number of questions required to specify a derivative. To do so, our approach tries to find an optimal question ordering that maximizes the number of decisions automatically derived from other questions previously answered.

Table 1. Valid derivatives for Figure 1. SA, standard avoidance; EA, enhanced avoidance.

Valid Derivatives	
1	¬PP, ¬LRF, ¬FRF, ¬SA, ¬EA
2	¬PP, ¬LRF, FRF, ¬SA, ¬EA
3	¬PP, LRF, ¬FRF, ¬SA, ¬EA
4	¬PP, LRF, FRF, ¬SA, ¬EA
5	¬PP, ¬LRF, ¬FRF, SA, ¬EA
6	¬PP, LRF, ¬FRF, SA, ¬EA
7	¬PP, ¬LRF, FRF, ¬SA, EA
8	¬PP, LRF, FRF, ¬SA, EA
9	¬PP, ¬LRF, FRF, SA, ¬EA
10	¬PP, LRF, FRF, SA, ¬EA
11	PP, LRF, ¬FRF, SA, ¬EA
12	PP, LRF, FRF, ¬SA, EA
13	PP, LRF, FRF, SA, ¬EA

A straightforward approach to get such optimal question ordering is computing for each valid product all possible orderings and, thus, finding the ordering with less questions on average for every product. Table 2 sums up the needed computations. For instance, the next-to-last column summarizes the number of questions needed for derivative {PP, LRF, FRF, SA, ¬EA}. Ordering PP < LRF < FRF < SA < EA needs three questions, LRF < PP < FRF < SA < EA needs four, and so on. Afterwards, the average number of questions for each ordering is computed. Using this approach in the previous example, ordering PP < LRF < FRF < SA < EA would be selected as an optimal one. As a result, the question sequence for Derivative 11 in Table 1 would be shortened to (1) is PP in the configuration? yes, (2) FRF? no, (3) SA? yes, removing the need for answering “if LRF is in the configuration”.

Unfortunately, this approach requires $m \cdot n!$ computations, where n is the number of components of the configuration model and m is a number $\leq 2^n$. Therefore, it is extremely expensive in computational terms and does not scale, except for the most trivial configuration models. To overcome the scalability limitations of the former approach, this paper proposes a heuristic solution grounded on the information theory concept of entropy.

Table 2. Brute force approach to compute the optimal ordering on average.

orderings ($n!$)	derivatives ($\leq 2^n$)				average number of questions
	{¬PP, ¬LRF, ¬FRF, ¬SA, ¬EA}	{¬PP, ¬LRF, FRF, ¬SA, ¬EA}	...	{PP, LRF, FRF, SA, ¬EA}	
PP < LRF < FRF < SA < EA	4	4	...	3	(4+4+...+3)/13
LRF < PP < FRF < SA < EA	4	4	...	4	(4+4+...+4)/13
⋮	⋮	⋮	⋮	⋮	⋮
LRF < FRF < SA < EA < PP	4	4	...	4	(4+4+...+4)/13

3. Related Work

Research on automated configurators is mainly focused on consistency checking and optimization [3,23]. For example, reasoning engines, such as BDD libraries, SAT solvers and logic-truth maintenance systems, have been used to detect invalid derivatives (*i.e.*, those which violate some option dependency or incompatibility) [24–26], to provide explanations for configuration flaws [27,28], to optimize configurations (*i.e.*, to find configurations whose cost is less or equal than a given one) [29–31], *etc.*

Despite the importance of the interactive question ordering problem that our work tackles, which was pointed out by Steinberg more than thirty years ago [32], there is little research on it. A recent approach that specifically deals with this problem is provided by Chen *et al.* [10], who propose to minimize the number of configuration steps by sorting the components according to their probability of being included in a derivative. Such a probability is computed by Equation (1).

$$\Pr(c) = \frac{\text{Number of valid derivatives that include } c}{\text{Total number of valid derivatives}} \quad (1)$$

In their original paper and with a fully equivalent meaning, Chen *et al.* [10] use the term selectivity instead of probability. As our approach follows an entropy driven heuristic and the information theory concept of entropy is defined in terms of probability (see Section 4.1.1.), we have preferred to use probability throughout this paper.

In addition to Chen *et al.*'s approach, Mazo *et al.* [9] proposes the following heuristics for ordering configuration questions:

Heuristic 1: Components with the smallest domain first. Choose first the component with the smallest domain. The domain of a component is the set of possible values that the component can take according to its domain definition and the constraints in which the component is involved.

Heuristic 2: The most constrained components first. Choose the component that participates in the largest number of constraints.

Heuristic 3: Components appearing in most products first. This heuristic is exactly the same as Chen *et al.*'s approach.

Heuristic 4: Automatic completion when there is no choice. This heuristic “provides a mechanism to automatically complete the configuration of variables where only one value of their domain is possible [...] it also works when a variable has several values on its domain but only one is valid”. In ascending order of computational cost and descending order of constraint propagation capacity, Mazo *et al.* [9] summarize three approaches to implement Heuristic 4 when the configuration model is encoded as a predicate logic formula: (i) forward-checking; (ii) partial look-ahead; and (iii) full look-ahead (*i.e.*, whereas forward-checking is the fastest algorithm, but produces the most limited form of constraint propagation during search, full look-ahead is the most expensive approach, but gets the best results). In our paper, configuration models are encoded as propositional logic formulas, where a full constraint propagation is computationally feasible. In particular, a configuration model will be encoded as a BDD ψ . To completely propagate a customer decision d , the BDD $\psi \wedge d$ will be computed. Fortunately, this computation only takes linear time on the size of the BDD. That is, the complexity of computing mixing two BDDs

ψ_1 and ψ_2 using the apply algorithm proposed by Bryant [33] is $O(|\psi_1||\psi_2|)$, so the complexity of $\psi \wedge d$ is $O(|\psi||d|) = O(|\psi| \cdot 1) = O(|\psi|)$. Instead of considering Heuristic 4 apart, we will use it as a complement to the remaining heuristics by running constraint propagation after every configuration step.

Heuristic 5: Components required by the latest configured component first. Choose the component that has the largest number of constraints with the past-configured components.

Heuristic 6: Components that split the problem space in two first. Set first the components that divide the problem space into two parts of approximately the same size. Unfortunately, Mazo *et al.* [9] do not provide a way to implement this heuristic, which takes into account all model constraints. In particular, Mazo *et al.* propose a simplification by just using the tree structure of an FD, or the variation points of an orthogonal variability model [34], but not processing the cross-tree constraints.

As will be discussed in Section 4.3, our approach may be thought of as an implementation of Heuristic 6 that, in addition, takes into account all configuration model constraints. In Section 5, it will be shown that our approach provides better outcomes than Heuristics 1, 2, 3 (*i.e.*, Chen's approach [10]) and 5.

4. Entropy-Based Approach to Sort Configuration Questions

This section presents our heuristic to minimize the number of steps required to configure a derivative from a configuration model. Section 4.1.1. introduces the theoretical background of our approach. As we will see, our heuristic, as other ones summarized in Section 3, requires computing the component probabilities. Section 4.1.2. discusses the scalability limitations of the approach commonly used to compute those probabilities. To overcome such limitations, in Section 4.2, we propose an algorithm that provides an efficient probability computation. Finally, Section 4.3 describes our heuristic.

4.1. Preliminaries

4.1.1. Information Theory

The following definitions were originally introduced by Shannon [8]. Let us start with the concept of entropy.

Definition 1 Let X be a discrete random variable with alphabet X and probability mass function $\Pr(x) = \Pr\{X = x\}$, $x \in X$; the entropy H of X is defined by Equation (2):

$$H(X) = - \sum_{x \in X} \Pr(x) (\log_2 \Pr(x)) \quad (2)$$

Let us present the concept of conditional entropy, which is the entropy of a random variable conditional on the knowledge of another random variable.

Definition 2 Let X and Y be two discrete random variables. The conditional entropy $H(X|Y)$ of X given Y is defined by Equation (3):

$$H(X|Y) = \sum_{y \in \mathcal{Y}} \Pr(y) H(X|Y = y) \quad (3)$$

Finally, let us introduce the concept of mutual information, also called information gain, which represents the reduction in a variable uncertainty due to another random variable.

Definition 3 Consider two random variables X and Y with a joint probability mass function $\Pr(x, y)$ and marginal probability mass functions $\Pr(x)$ and $\Pr(y)$. The mutual information $I(X; Y)$ is defined by Equation (4) as the relative entropy between the joint distribution and the product distribution $\Pr(x)\Pr(y)$:

$$I(X; Y) = \sum_{x,y} \Pr(x, y) \log_2 \frac{\Pr(x, y)}{\Pr(x)\Pr(y)} \tag{4}$$

Entropy and mutual information satisfy the following properties that will be used throughout this paper:

- (1) $H(X) \geq 0$
- (2) $H(X) \leq \log_2 \#\mathcal{X}$, with equality if and only if X is distributed uniformly over \mathcal{X} (in this paper, the number of elements of a set S is denoted as $\#S$)
- (3) $I(X; Y) = H(X) - H(X|Y) = H(Y) - H(Y|X) = I(Y; X)$

4.1.2. Straightforward Approach to Compute Component Probabilities

A widespread approach to support the automated management of configuration models is translating them to propositional logic formulas [3,25], which are processed using off-the-self tools, such as SAT solvers [35] or BDD engines [33]. Table 3 summarizes the translations needed to encode our running example into propositional logic. A more detailed explanation of how to translate feature models into logic may be found in [36]. Equation (5) is the equivalent logic formula to Figure 1.

$C \neg\circ ADC$	$\xrightarrow{\text{is translated to}}$	$(\neg ADC \vee C) \wedge$
$ADC \rightarrow\bullet CAB$	$\xrightarrow{\text{is translated to}}$	$(\neg ADC \vee CAB) \wedge (\neg CAB \vee ADC) \wedge$
CAB	$\xrightarrow{\text{is translated to}}$	$(\neg SA \vee CAB) \wedge (\neg EA \vee CAB) \wedge (\neg CAB \vee SA \vee EA) \wedge (\neg SA \vee \neg EA) \wedge$
$SA \quad EA$		
$ADC \neg\circ PP$	$\xrightarrow{\text{is translated to}}$	$(\neg PP \vee ADC) \wedge$
$C \rightarrow\bullet S$	$\xrightarrow{\text{is translated to}}$	$(\neg S \vee C) \wedge (\neg C \vee S) \wedge$
$S \neg\circ LRF$	$\xrightarrow{\text{is translated to}}$	$(\neg LRF \vee S) \wedge$
$S \neg\circ FRF$	$\xrightarrow{\text{is translated to}}$	$(\neg FRF \vee S) \wedge$
$PP \xrightarrow{\text{requires}} LRF$	$\xrightarrow{\text{is translated to}}$	$(\neg PP \vee FRF)$

Once a configuration model is encoded into a logic formula ψ :

- the total number n_1 of valid derivatives is equivalent to the number of satisfying assignments of ψ (i.e., those that evaluate ψ to be true).
- the number n_2 of valid derivatives that include component c is equivalent to the number of satisfying assignments of $\psi \wedge c$.

Table 3. Equivalence between configuration models and propositional logic formulas.

Type of relationship	Feature model representation	Translation to propositional logic
mandatory	$A \dashv\vdash B$	$(\neg A \vee B) \wedge (\neg B \vee A)$
optional	$A \circ B$	$\neg B \vee A$
alternative	A	$\bigwedge_{i=1}^n (\neg B_i \vee A) \wedge (\neg A \vee \bigvee_{i=1}^n B_i) \wedge \bigwedge_{i < j} (\neg B_i \vee \neg B_j)$
requires	$ \begin{array}{cccc} B_1 & B_2 & \dots & B_n \\ A \xrightarrow{\text{requires}} & & & B \end{array} $	$\neg A \vee B$

Since the probability of a component is $\frac{n_2}{n_1}$ and the computation of the number of satisfying assignments of a Boolean formula is supported by most BDD engines and SAT solvers (in particular, #SAT counters are a type of SAT solver specifically oriented to compute such a number), a straightforward approach to compute the component probabilities is calling repeatedly a logic engine using $\psi \wedge c_i$ as the input [11]. Unfortunately, this approach has a high computational cost and does not scale for all, but trivial configuration models. While the SAT problem is known to be NP-complete [37], it is widely believed that the #SAT problem is even harder [35]. If n is the number of components, computing the component probabilities requires calling a #SAT solver n times, which is extremely time-consuming. Similarly, computing the number of satisfying assignments with a BDD has computational complexity $O(m)$ [33], where m is the number of nodes of the BDD. Hence, the complexity of computing the component probabilities by calling repeatedly the BDD engine is $O(n \cdot m)$, which is excessively time-consuming for most configuration models.

For instance, it is well known by the car manufacturing community that the first issue of car configurators is performance [38]. Thus, as soon as customers make a configuration choice, they want to find out what the consequences of the choice are. From a marketing perspective, it is unpleasant for customers to wait for several seconds to know whether their requirements are correct or not in terms of configuration. As it will be shown experimentally in Section 5, computing the component probabilities by calling repeatedly a BDD may force the customer to wait for more than 600 seconds for just a single configuration step!

To overcome the aforementioned scalability limitations, in the following section, we propose a BDD algorithm that computes component probabilities in almost linear time to m .

4.2. Efficient Computation of the Probabilities of the Variables of a Boolean Formula from a BDD

BDDs are a way of representing Boolean functions. They are rooted, directed, acyclic graphs, which consist of several decision nodes and terminal nodes [33]. There are two types of terminal nodes, called 0-terminal and 1-terminal. Each decision node v_i is labeled by a Boolean variable x_k and has two child nodes called low and high (which are usually depicted by dashed and solid lines, respectively). The edge from node v_i to a low (or high) child represents an assignment of v_i to 0 (resp. 1). Such a BDD is called

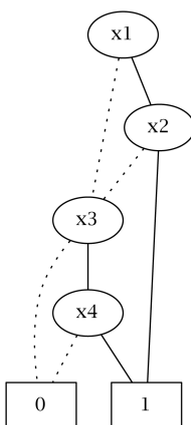
ordered if different variables appear in the same order on all paths from the root. A BDD is said to be reduced if the following two rules have been applied to its graph: (i) isomorphic subgraphs are merged; and (ii) nodes whose two children are isomorphic are eliminated.

In popular usage, the term BDD almost always refers to a reduced ordered binary decision diagram [12]. In this paper, we will follow that convention, as well. Let us use formula $\psi \equiv (x_1 \wedge x_2) \vee (x_3 \wedge x_4)$ as a running example for this subsection. Table 4 is the truth table for ψ . Figure 2 is its BDD representation using the variable ordering $x_1 < x_2 < x_3 < x_4$. Note that a logic formula may be encoded with different BDDs according to the variable ordering used to synthesize the BDD. Obviously, our algorithm produces the same results for equivalent BDDs (*i.e.*, BDDs that encode the same formula).

Table 4. Truth table for $\psi \equiv (x_1 \wedge x_2) \vee (x_3 \wedge x_4)$.

x_1	x_2	x_3	x_4	ψ
0	0	0	0	0
0	0	0	1	0
0	0	1	0	0
0	0	1	1	1
0	1	0	0	0
0	1	0	1	0
0	1	1	0	0
0	1	1	1	1
1	0	0	0	0
1	0	0	1	0
1	0	1	0	0
1	0	1	1	1
1	1	0	0	1
1	1	0	1	1
1	1	1	0	1
1	1	1	1	1

Figure 2. The binary decision diagram (BDD) for ψ according to the variable ordering $x_1 < x_2 < x_3 < x_4$.



The remainder of this subsection is structured as follows. Firstly, some definitions required to understand our algorithm are given. Next, the data structures that the algorithm uses are described from a theoretical perspective. Then, the algorithm is presented. Finally, the algorithm computational cost is discussed.

4.2.1. Definitions

Definition 4 The satisfying set of a Boolean formula $\psi(x_1, \dots, x_n)$, denoted S_ψ , is defined by Equation (6).

$$S_\psi = \{(x_1, \dots, x_n) | \psi(x_1, \dots, x_n) = \text{true}\} \tag{6}$$

Definition 5 The satisfying set of the variable x_i of a Boolean formula $\psi(x_1, \dots, x_{i-1}, x_i, x_{i+1}, \dots, x_n)$, denoted $S_{\psi|x_i=\text{true}}$, is defined by Equation (7).

$$S_{\psi|x_i=\text{true}} = \{(x_1, \dots, x_{i-1}, \text{true}, x_{i+1}, \dots, x_n) | \psi(x_1, \dots, x_{i-1}, \text{true}, x_{i+1}, \dots, x_n) = \text{true}\} \tag{7}$$

For instance, according to Table 4, $\#S_\psi = 7$ since there are seven rows where ψ evaluates as true (throughout this paper, 0/1 and false/true are used interchangeably), and $\#S_{\psi|x_4} = 5$, because $x_4 = 1$ in five of the seven rows where $\psi = 1$.

Definition 6 The satisfying probability of a Boolean formula $\psi(x_1, \dots, x_n)$, denoted $\text{Pr}(\psi)$, is defined by Equation (8).

$$\text{Pr}(\psi) = \frac{\#S_\psi}{2^n} \tag{8}$$

Definition 7 The satisfying marginal probability of a variable x_i in a Boolean formula $\psi(x_1, \dots, x_{i-1}, x_i, x_{i+1}, \dots, x_n)$, denoted $\text{MPr}(\psi|x_i=\text{true})$, is defined by Equation (9).

$$\text{MPr}(\psi|x_i=\text{true}) = \frac{\#S_{\psi|x_i=\text{true}}}{2^n} \tag{9}$$

Definition 8 The satisfying probability of a variable x_i in a Boolean formula $\psi(x_1, \dots, x_{i-1}, x_i, x_{i+1}, \dots, x_n)$, denoted $\text{Pr}(\psi|x_i=\text{true})$, is defined by Equation (10).

$$\text{Pr}(\psi|x_i=\text{true}) = \frac{\#S_{\psi|x_i=\text{true}}}{\#S_\psi} \tag{10}$$

For instance, looking at Table 4, it is easy to see that $\text{Pr}(\psi) = \frac{7}{2^4}$, $\text{MPr}(\psi|x_4=\text{true}) = \frac{5}{2^4}$, and $\text{Pr}(\psi|x_4=\text{true}) = \frac{5}{7}$. For convenience, in the remainder of the paper, we denote $\text{Pr}(\psi|x_i=\text{true})$ and $\text{MPr}(\psi|x_i=\text{true})$ as $\text{Pr}(x_i)$ and $\text{MPr}(x_i)$, respectively.

4.2.2. Data Structures

Let us represent a BDD that has m nodes and encodes a Boolean formula with n variables by using the following data structures:

- The variable ordering used to synthesize the BDD is represented by an array declared as follows:

```
var_ordering: array[0..n-1] of string
```

- Each node is represented by a record declared as follows:

```
type node = record
  index: 0..n
  low, high: node
  mark: Boolean
end
```

where:

- (1) index is the index of the variables in the ordering. The terminal nodes of the BDD (i.e., 0 and 1) have index n .
- (2) low and high are the low and high node successors
- (3) mark is used to mark which nodes have been visited during a traversal of the graph. As we will see, our algorithm is called at the top level with the root node as the argument and with the mark fields of the nodes being either all true or all false. It then systematically visits every node in the graph by recursively visiting the subgraphs rooted by the two children, low and high. As it visits a node, it complements the value of the mark field, so that it can later determine whether a child has already been visited by comparing the two marks.

- The BDD is represented by an array declared as follows:

```
bdd: array[0..m] of node
```

The terminal nodes of the BDD, 0 and 1, are stored at Positions 0 and 1 of the bdd array, respectively.

For instance, Tables 5 and 6 represent the content of bdd and *var_ordering* for the BDD in Figure 2, respectively.

Table 5. Content of the bdd array for Figure 2.

position	index	low	high	mark
0	4	nil	nil	false
1	4	nil	nil	false
2	3	0	1	false
3	2	0	2	false
4	1	3	1	false
5	0	3	4	false

Table 6. Content of the *var_orderingarray* for Figure 2.

position	content
0	“x ₁ ”
1	“x ₂ ”
2	“x ₃ ”
3	“x ₄ ”

4.2.3. Algorithm

Pr(*x_i*) is computed jointly by Algorithms 1, 2 and 3. Figure 3 summarizes the computations for the BDD in Figure 2. Let us examine how our approach proceeds:

Algorithm 1 computes Pr(*x_i*) as $Pr(x_i) = \frac{MPr(x_i)}{Pr(\psi)}$ by calling the auxiliary Algorithms 2 and 3.

Algorithm 2 computes Pr(ψ).

A nice mental picture to understand Algorithm 2 is thinking about pouring one liter of water from the BDD root to the terminal nodes. One liter goes through the root, then half a liter goes through the low branch and half a liter through the high branch. This procedure advances, until the water reaches the leaves. Hence, MPr(*x_i*) is the amount of water that Node 1 has.

According to Tables 5 and 6, the root node has label *v₅*, and it is in Position 5 of the bdd array. In Figure 2, through node *v₅* goes one liter (*i.e.*, formula_sat_prob[5] = 1). Half of it goes to *v₃* and the other half to *v₄*. Whereas, through *v₄* passes $\frac{1}{2}$ liters, through *v₃* goes the $\frac{1}{2}$ liters that come from *v₅* and half of the water that comes from *v₄* (*i.e.*, formula_sat_prob[3] = $\frac{1}{2} + \frac{1}{2} = \frac{3}{4}$).

Algorithm 3 computes MPr(*x_i*).

In particular, let us examine how it computes MPr(*x₂*). In the truth Table 4, ψ evaluates as true when *x₂* is true five times:

- (1) In four of them, *x₁* is true. When the call get_marginal_prob(4, ...) is made, Lines 10–23 compute the marginal probability of *x₂* for the explicit path *v₅* → *v₄*. The probabilities due to the low and high branches of *v_i* are stored into the prob_low and prob_high variables, respectively. As bdd[4].low ≠ 0, a recursive call is made to compute the total probability due to the low descendants of *v₄* (*i.e.*, get_marginal_prob(3, ...)). As a result:

$$total_prob[3] = prob_low_{v_3} + prob_high_{v_3} = 0 + \frac{3}{16} = \frac{3}{16}$$

Notice that prob_low_{*x₂*} is not simply equal to total_prob[3], because total_prob[3] depends also on the probability that comes from the link *v₅* → *v₃*. To get just the probability due to the link *v₄* → *v₃*, prob_low has to be adjusted using the formula_sat_prob array as:

$$prob_low = \frac{total_prob[3] \cdot \frac{formula_sat_prob[4]}{2}}{formula_sat_prob[3]} = \frac{\frac{3}{16} \cdot \frac{1}{2}}{\frac{3}{4}} = \frac{1}{16}$$

Since $\text{bdd}[4].\text{high} = 1$, prob_high is directly computed as:

$$\text{prob_high} = \frac{\text{formula_sat_prob}[4]}{2} = \frac{\frac{1}{2}}{2} = \frac{1}{4}$$

Finally:

$$\text{prob}[\text{bdd}[4].\text{index}] = \text{prob_high} = \frac{1}{4}$$

(2) In one of them, x_1 is false. The two following implicit paths have been removed from the reduced BDD, (i) $v_5 \dashrightarrow v_4 \dashrightarrow v_3$; and (ii) $v_5 \dashrightarrow v_4 \rightarrow v_3$. Nevertheless, path $v_5 \dashrightarrow v_4 \rightarrow v_3$ should be considered to compute the marginal probability of x_2 . Lines 24–31 account for that kind of implicit path, adjusting the marginal probability with the variables omitted in the paths. For instance, when the algorithm is called for v_5 , the marginal probability of x_2 is updated with half the prob_low of v_5 .

To sum up:

$$\text{MPr}(x_2) = \text{MPr}(v_5 \dashrightarrow v_4 \rightarrow v_3) + \text{MPr}(v_5 \rightarrow v_4) = \frac{\text{prob_low}_{v_5}}{2} + \text{prob}[\text{bdd}[4].\text{index}] = \frac{\frac{1}{8}}{2} + \frac{1}{4} = \frac{5}{16}$$

Algorithm 1: get_prob

```

1 Input bdd and var_ordering arrays
2 Output an array which stores  $\text{Pr}(x_i)$  in position  $i$ 
3 var formula_sat_prob, total_prob: array[0..length(bdd)-1] of float;
4   prob: array[0..length(var_ordering)-1] of float; i: int;
5 begin
6   for ( $i=0$ ;  $i < \text{length}(\text{bdd})$ ;  $i++$ ) do
7      $\text{total\_prob}[i] = 0.0$ 
8   for ( $i=0$ ;  $i < \text{length}(\text{var\_ordering})$ ;  $i++$ ) do
9      $\text{prob}[i] = 0.0$ 
10  formula_sat_prob = get_formula_sat_prob(bdd)
11  get_marginal_prob(length(bdd)-1, total_prob, formula_sat_prob, prob, bdd, var_ordering)
12  for ( $i=0$ ;  $i < \text{length}(\text{var\_ordering})$ ;  $i++$ ) do
13     $\text{prob}[i] = \frac{\text{prob}[i]}{\text{formula\_sat\_prob}[1]}$ 
14  return prob

```

Algorithm 2: get_formula_sat_prob

```

1 Input bdd array
2 Output an array which in position  $i$  stores  $\text{Pr}(\psi)$ 
3 var formula_sat_prob: array[0..length(bdd)-1] of float; i: int
4 begin
5   for ( $i=0$ ;  $i < \text{length}(\text{bdd})-1$ ;  $i++$ ) do
6     formula_sat_prob[i] = 0.0 // non-root nodes prob is initialized to 0
7   formula_sat_prob[i] = 1.0 // root node prob is 1
8    $i = \text{length}(\text{bdd})-1$ 
9   while  $i > 1$  do // for all non-terminal nodes
10    formula_sat_prob[bdd[i].low] +=  $\frac{\text{formula\_sat\_prob}[i]}{2.0}$ 
11    formula_sat_prob[bdd[i].high] +=  $\frac{\text{formula\_sat\_prob}[i]}{2.0}$ 
12     $i--$ 
13  return formula_sat_prob

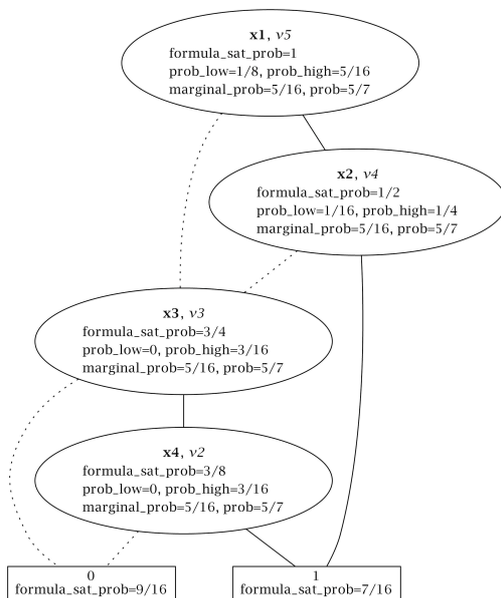
```

Algorithm 3: get_marginal_prob

```

1 Input v: 0..length(bdd)-1; total_prob, formula_sat_prob: array[0..length(bdd)-1] of float;
2   prob: array[0..length(var_ordering)-1] of float; bdd and var_ordering arrays
3 Output prob is passed by reference and, at the end of the algorithm execution,
4   it stores MPr(xi) in position i
5 var prob_low, prob_high: float; i: int
6 begin
7   prob_low = 0.0
8   prob_high = 0.0
9   bdd[v].mark = not bdd[v].mark
  // explicit path recursive traversal
10  if bdd[v].low == 1 then
11    | prob_low =  $\frac{\text{formula\_sat\_prob}[v]}{2.0}$ 
12  else if bdd[v].low ≠ 0 then
13    | if bdd[v].mark ≠ bdd[bdd[v].low].mark then
14      | | get_marginal_prob(bdd[v].low, total_prob, formula_sat_prob, prob, bdd, var_ordering)
15    | prob_low =  $\frac{\text{total\_prob}[\text{bdd}[v].\text{low}] \cdot \frac{\text{formula\_sat\_prob}[v]}{2.0}}{\text{formula\_sat\_prob}[\text{bdd}[v].\text{low}]}$ 
16  if bdd[v].high == 1 then
17    | prob_high =  $\frac{\text{formula\_sat\_prob}[v]}{2.0}$ 
18  else if bdd[v].high ≠ 0 then
19    | if bdd[v].mark ≠ bdd[bdd[v].high].mark then
20      | | get_marginal_prob(bdd[v].high, total_prob, formula_sat_prob, prob, bdd, var_ordering)
21    | prob_high =  $\frac{\text{total\_prob}[\text{bdd}[v].\text{high}] \cdot \frac{\text{formula\_sat\_prob}[v]}{2.0}}{\text{formula\_sat\_prob}[\text{bdd}[v].\text{high}]}$ 
22  total_prob[v] = prob_low + prob_high
23  prob[bdd[v].index] += prob_high
  // implicit path iterative traversal
24  i = bdd[v].index + 1
25  while i < bdd[bdd[v].low].index do
26    | prob[i] +=  $\frac{\text{prob\_low}}{2.0}$ 
27    | i += 1
28  i = bdd[v].index + 1
29  while i < bdd[bdd[v].high].index do
30    | prob[i] +=  $\frac{\text{prob\_high}}{2.0}$ 
31    | i += 1
  
```

Figure 3. Probability computation for BDD 2.



4.2.4. Computational Cost

Let m be the number of nodes of the BDD and n the number of variables of the Boolean formula. Algorithm 2 requires traversing all of the nodes, so its computational complexity is $O(m)$. Algorithm 3 also traverses all of the BDD nodes. In addition, to account for the implicit paths removed from the reduced BDD, the variables omitted on the edges that come from each node need to be traversed (which is done by Lines 24–31). Table 7 summarizes those traversals for Figure 2. For instance, when v_4 is recursively traversed, the variables x_3 and x_4 need to be iteratively traversed, because the edge $v_4 \rightarrow 1$ omits them (*i.e.*, the variable encoded by node v_4 , x_2 , jumps directly to one, omitting the intermediate variables x_3 and x_4 in the ordering $x_1 < x_2 < x_3 < x_4$). Table 7 helps with noticing the savings our algorithm provides compared to the straightforward approach described in Section 4.1.2., which requires traversing all nodes for all variables (which in computational cost terms is equivalent to traversing all variables for every node). Therefore, Algorithm 3 does not traverse $m \cdot n$ elements, but $m \cdot n'$, where n' is strictly less than n . Otherwise, if $n' = n$, all nodes in the BDD should go directly to zero or one, jumping over all of the variables. Nevertheless, as BDDs are organized in hierarchical levels according to the variable ordering, this is impossible (*i.e.*, the nodes that encode a variable with position k in the ordering can only jump over the variables with positions $k + 1 \dots n$).

It follows that Algorithm 1 has computational complexity $O(m \cdot n')$. As will be shown in Section 5, in practice, n' is usually much smaller than n , and thus, variable probabilities can be efficiently computed.

Table 7. Variables iteratively traversed for BDD in Figure 2.

node	arcs	omitted varsthat are traversed
v_5	$v_5 \dashrightarrow v_3$	x_2
	$v_5 \rightarrow v_4$	none
v_4	$v_4 \dashrightarrow v_3$	none
	$v_4 \rightarrow 1$	x_3, x_4
v_3	$v_3 \dashrightarrow 0$	x_4
	$v_3 \rightarrow v_2$	none
v_5	$v_5 \dashrightarrow 0$	none
	$v_5 \rightarrow 1$	none

4.3. Entropy-Driven Configuration

Let us return to the original problem this paper tackles. Given a set of questions Q , our goal is to sort it in such a way that the user has to answer as few questions as possible to complete the configuration. To find the optimal order of Q , we propose to rank each question, q , according to its expected information gain, *i.e.*, measuring how much uncertainty can be reduced on average when the engineer answers it. Such an information gain is modeled as the mutual information $I(C; q)$, where C is the set of all valid configurations (*i.e.*, the ones that satisfy all asset interdependencies).

When a configuration is completed, the entropy of every question q is zero. Since q has been answered, $H(q|C) = 0$. Thus, it follows that $I(C; q) = H(q)$, as Equation (11) demonstrates (see Property 3 in Subsection 4.1.1.).

$$I(C; q) = H(q) - H(q|C) = H(q) \quad (11)$$

When we ask “is component c in the configuration?”, the entropy of the question $H(q)$ is computed by Equation (12), where $\Pr(c)$ is the probability that c is included in the configuration.

$$\begin{aligned} H(q) &= -\Pr(c)\log_2\Pr(c) - \Pr(\neg c)\log_2\Pr(\neg c) \\ &= -\Pr(c)\log_2\Pr(c) - (1 - \Pr(c))\log_2(1 - \Pr(c)) \end{aligned} \quad (12)$$

Our approach to guide the configuration of a derivative may be thought of as a binary search for the user-desired configuration (remember Heuristic 6 in Section 3). To successively divide the search space into subspaces of approximately the same size (*i.e.*, where the pursued configuration is approximately of the same probability), the user answers the question that provides more information about the configuration (*i.e.*, the question with the highest entropy). Thus, the configuration process advances iteratively, by performing the following activities, until the entropy of all components becomes zero:

- (1) Computing the component probabilities from the input configuration model. As the process advances, the configuration space gets narrower and, consequently, the component probabilities change.
- (2) Computing the entropy value for each question.
- (3) Sorting the questions in descending order of entropy.
- (4) Asking the user to answer a question with entropy greater than zero. Note that when a question has zero entropy, it is because it has been answered in a previous step directly or indirectly (*i.e.*, because of the question interdependencies).
- (5) Updating the set of answers and the configuration model (e.g., if the customer answers a question q negatively, the Boolean formula ψ that encodes the configuration model is updated to $\psi \wedge \neg f$).

Entropy may also be used to measure how hard it is to configure a given model. From the “point of view” of an automated configurator, when the configuration process starts, the derivative desired by the customer is any c in C with the same probability. Therefore, the configuration model uncertainty is calculated by Equation (13) (see Property 2 in Section 4.1.1.).

$$H(C) = \log_2\#C \quad (13)$$

4.3.1. Example

Coming back to the running example introduced in Section 2, let us see how our approach works. Figure 4 sums up the steps required to configure the derivative {PP, LRF, \neg FRF, SA, \neg EA} using the entropy heuristic. In the first step, EA is the component with the highest entropy. Therefore, the system asks the user if SA is included in the derivative. Once the user answers affirmatively, the probabilities of the components are recomputed and, so, the entropies (e.g., the inclusion of SA implies the exclusion of EA, so $\Pr(\text{EA})=0$ and thus $H(\text{EA})=0$).

Figure 4. Configuring derivative {PP, LRF, -FRF, SA, -EA} using component entropy.

Component	H
PP	0.78
LRF	0.96
FRF	0.96
SA	0.99
EA	0.78

SA ? Yes

→

Component	H
PP	0.92
LRF	0.92
FRF	1
SA	0
EA	0

FRF ? No

→

Component	H
PP	0.92
LRF	0.92
FRF	0
SA	0
EA	0

PP ? Yes

We remark here that our approach does not force the user to follow a fixed sequence of questions. In each configuration step, the user may decide not to answer the best entropy-ranked question, but the one she thinks is more convenient. After the question is answered, the entropies are recomputed and, thus, our approach adjusts to the user preferences in an interactive way.

5. Experimental Evaluation

To test the validity of our approach, we have used two case studies:

- (1) The configuration model provided by the car manufacturing company Renault DVI [14], which deals with the configuration of a family of cars named Renault Megane and is freely available at [15]. We have selected this model because it illustrates the practical applicability of our approach (*i.e.*, instead of using an example made up for academic purposes, our work is tested on a real configuration model that comes from the industry). In addition, the Renault Megane problem is a benchmark of widespread use by the configuration community [39–49].
- (2) The Electronic Shopping model provided by Lau [17], which deals with an electronic commerce platform that allows consumers to directly buy goods or services from a seller over the Internet using a web browser. This benchmark is widely used by the software product line community [26,50,51] and is freely available at [16].

5.1. Experimental Design

The goal of this section is to check if:

- Our approach produces better results than related work.
- The algorithm presented in Section 4.2 provides reasonable response times and, thus, supports customer interactivity during the configuration process.

To do so, we have created a test bed composed of 1,000 random derivatives for every configuration model. As we will see, a sample of 1,000 derivatives is big enough to get results with high statistical power and significance.

To generate valid derivatives that satisfy all constraints, we have encoded the models as propositional logic formulas (see Subsection 4.1.2.) and then as BDDs. To get efficient BDD sizes, the directions given by Narodytska *et al.* [43] have been followed. The BuDDy package [52] has been used to guarantee the generation of valid derivatives (*i.e.*, derivatives that conform to the BDD).

The test bed is used to compare the following methods:

- (1) Mazo *et al.*'s Heuristics 1, 2 and 5. [9] Remember that, strictly speaking, Mazo *et al.*'s Heuristic 4 is not a heuristic, but a propagation mechanism that all configuration systems should support. Therefore, we have included such mechanism in all of the methods tested in this paper.
- (2) The probability-driven approach, *i.e.*, the method proposed by Chen *et al.* [10] and Mazo *et al.* [9] (Heuristic 3).
- (3) The entropy-driven approach, *i.e.*, the method we propose in this paper.

To compute the option probabilities, which are required by the entropy and probability approaches, an implementation of the algorithm presented in Section 4.2 has been included into the BuDDy package.

5.2. Case Study 1: Renault Megane

5.2.1. Results

Table 8 summarizes the results of the experiments for the Renault Megane configuration model. Histograms in Figure 5a represent the number of steps needed to configure the 1,000 derivatives using Mazo *et al.*'s [9] Heuristics 1, 2 and 5 and the entropy and probability approaches. Figure 5b complements the histogram representation of the results with a box plot. In Figure 5b, “whiskers” start from the edge of the box and extend to the furthest data point that is within 1.5 times the inter-quartile range (*i.e.*, the range that goes from the 25th percentile to the 75th percentile). Points that are past the ends of the whiskers have been considered outliers and are displayed with dots.

Figure 5. The number of configuration steps according to the used approach for Renault Megane. (a): Histograms; (b): Box plots.

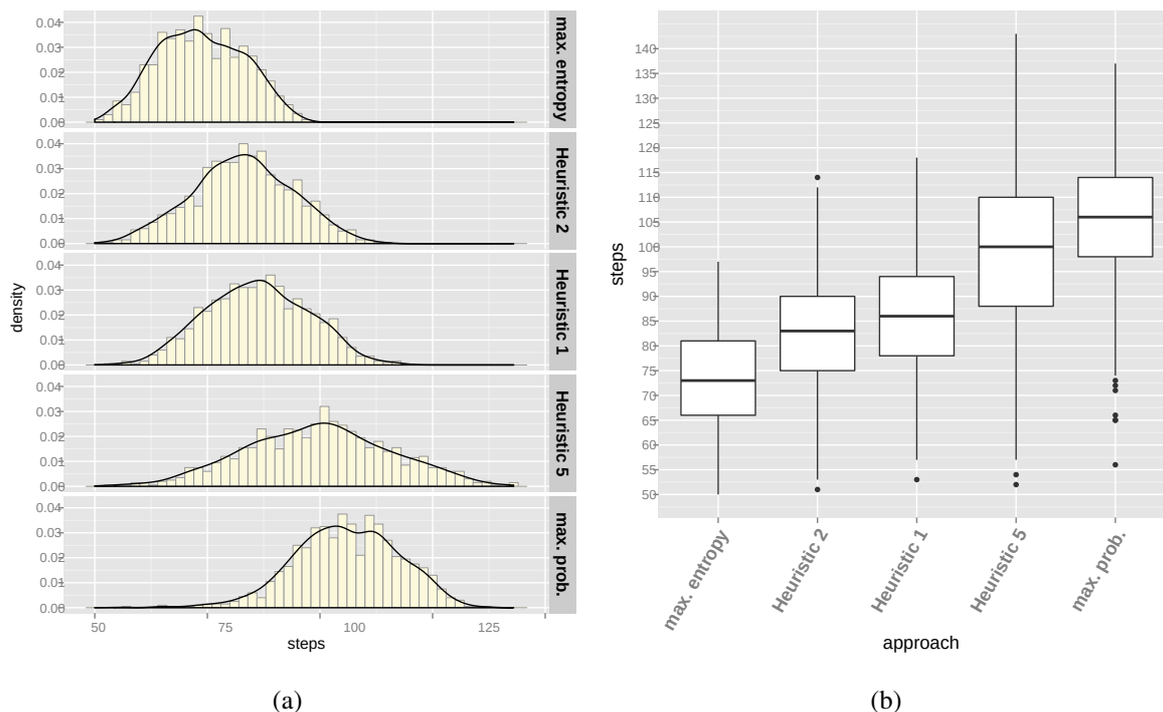


Table 8. Result of the experiments for Renault Megane.

approach	mean	SD	median	min	max	range
entropy	73.49	9.5	73	50	97	47
probability	105.79	11.54	106	56	137	81
Heuristic 1	86.04	11.26	86	53	118	65
Heuristic 2	82.74	11.12	83	51	114	63
Heuristic 5	99.39	15.95	100	52	143	91

Using the central limit theorem, the 95% confidence intervals (CI) of the population mean can be estimated (*i.e.*, the range where, with a 95% guarantee, the mean of the number of steps required to configure every derivative of the Megane model lies). Table 9 summarizes the CIs for each approach, which are estimated as $population\ mean\ CI = sample\ mean \pm t(std.\ error, 95\%, 999\ degrees\ of\ freedom)$, where t stands for the Student’s t -distribution.

Table 9. The 95% CIs of the population mean for Renault Megane.

entropy		probability		Heuristic 1		Heuristic 2		Heuristic 5	
std. error	95% CI	std. error	95% CI	std. error	95% CI	std. error	95% CI	std. error	95% CI
0.3	72.90-74.08	0.36	105.07-106.50	0.36	85.35-86.74	0.35	82.05-83.43	0.5	98.40-100.38

According to the summarized data, there is experimental evidence supporting that our approach produces better results than related work.

5.2.2. Statistical Significance

To check the statistical significance of the results, an analysis of variance (ANOVA) test has been run on the experimental data. Table 10 summarizes the ANOVA outcomes. Since the p -value is less than 0.001 (in particular, p -value $< 2 \cdot 10^{-16}$), the experimental results are statistically highly significant.

Table 10. ANOVA test for Renault Megane.

	Degrees of freedom	Sum of squares	Mean of squares	F-value	Pr(> F)
approaches	4	676,884	169,221	1,162	$< 2 \cdot 10^{-16}$
residuals	4,995	727,312	146		

Table 11 summarizes the power analysis of the ANOVA test. Given the sample size and the high effect size (*i.e.*, the high values of η^2 and Cohen’s f^2), the ANOVA test has high statistical power.

Finally, to check the statistical significance of the pairwise comparison between the approaches, a Tukey Honest Significant Differences (HSD) has been run. According to the results, summarized in Table 12, the difference between the number of steps required by any pair of approaches to configure a derivative is statistically highly significant. Whereas the ANOVA test rejects the null hypothesis: “there is no difference between the five approaches (*i.e.*, all of them produce approximately the same results)”,

the Tukey HSD test rejects ten null hypotheses separately: “there is no difference between Heuristic 2 and the entropy approach”, “there is no difference between Heuristic 1 and the entropy approach”, *etc.*

Table 11. Power analysis for Renault Megane.

Effect size		Power
eta squared η^2	Cohen’s f^2	
0.48	0.93	≈ 1

Table 12. Tukey HSD test for Renault Megane.

	Difference	95% CI	Adjusted <i>p</i> -value
Heuristic 2 vs. entropy	9.25	7.78–10.72	≈ 0
Heuristic 1 vs. entropy	12.56	11.08–14.02	≈ 0
Heuristic 5 vs. entropy	25.89	24.42–27.37	≈ 0
probability vs. entropy	32.29	30.82–33.77	≈ 0
Heuristic 1 vs. Heuristic 2	3.30	1.83–4.77	≈ 0
Heuristic 5 vs. Heuristic 2	16.65	15.17–18.12	≈ 0
probability vs. Heuristic 2	23.04	21.57–24.52	≈ 0
Heuristic 5 vs. Heuristic 1	13.34	11.87–14.82	≈ 0
probability vs. Heuristic 1	19.74	18.27–21.22	≈ 0
probability vs. Heuristic 5	6.40	4.93–7.87	≈ 0

5.3. Case Study 2: Electronic Shopping

Table 13 and Figure 6 summarize the results of the experiments for the Electronic Shopping configuration model. Table 14 summarizes the CIs for each approach. According to the outcomes, there is experimental evidence supporting that our approach produces better results than related work.

Table 13. Result of the experiments for Electronic Shopping.

approach	mean	SD	median	min	max	range
entropy	165.57	2.23	166	158	171	13
probability	193.67	6.06	194	164	207	43
Heuristic 1	187.38	5.69	188	168	201	33
Heuristic 2	189.36	5.7	190	170	203	33
Heuristic 5	169.33	3.1	169	153	178	25

Figure 6. The number of configuration steps according to the used approach for Electronic Shopping. (a): Histograms; (b): Box plots.

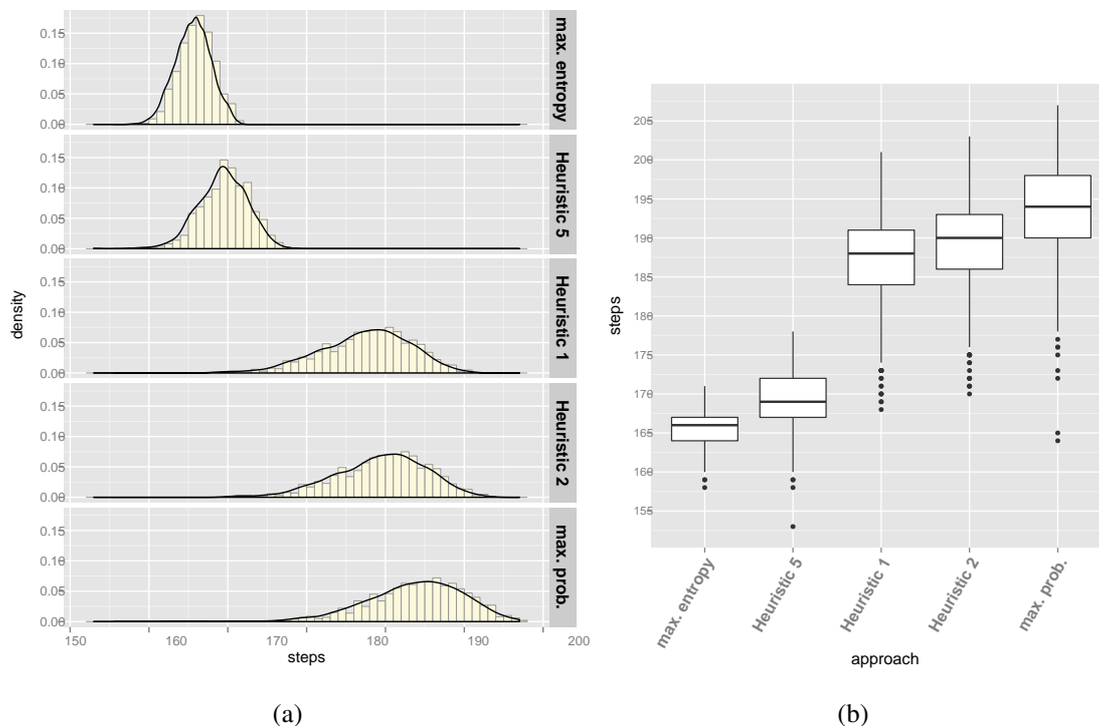


Table 14. The 95% CIs of the population mean for Electronic Shopping.

entropy		probability		Heuristic 1		Heuristic 2		Heuristic 5	
std. error	95% CI	std. error	95% CI	std. error	95% CI	std. error	95% CI	std. error	95% CI
0.07	165.43-165.71	0.19	193.29-194.04	0.18	187.03-187.73	0.18	189.01-189.72	0.1	169.14-169.52

5.3.1. Statistical Significance

Table 15 summarizes the ANOVA outcomes. Since the p -value is less than 0.001 (in particular, p -value $< 2 \cdot 10^{-16}$), the experimental results are statistically highly significant. Table 16 summarizes the power analysis of the ANOVA test. Given the sample size and the high effect size, the ANOVA test has high statistical power. Finally, Table 17 summarizes the outcomes of HSD, which show that the difference between the number of steps required by any pair of approaches to configure a derivative is statistically highly significant.

Table 15. ANOVA test for Electronic Shopping.

	Degrees of freedom	Sum of squares	Mean of squares	F -value	$\text{Pr}(> F)$
approaches	4	645,314	161,328	6,950	$< 2 \cdot 10^{-16}$
residuals	4,995	115,944	23		

Table 16. Power analysis for Electronic Shopping.

Effect size		Power
eta squared η^2	Cohen's f^2	
0.85	5.57	≈ 1

Table 17. Tukey HSD test for Electronic Shopping.

	Difference	95% CI	Adjusted p -value
Heuristic 5 vs. entropy	3.76	3.17–4.35	≈ 0
Heuristic 1 vs. entropy	21.81	21.22–22.40	≈ 0
Heuristic 2 vs. entropy	23.79	23.21–24.38	≈ 0
probability vs. entropy	28.09	27.51–28.68	≈ 0
Heuristic 1 vs. Heuristic 5	18.05	17.46–18.64	≈ 0
Heuristic 2 vs. Heuristic 5	20.03	19.45–20.62	≈ 0
probability vs. Heuristic 5	24.33	23.75–24.92	≈ 0
Heuristic 2 vs. Heuristic 1	1.98	1.39–2.57	≈ 0
probability vs. Heuristic 1	6.28	5.69–6.87	≈ 0
probability vs. Heuristic 2	4.30	3.71–4.89	≈ 0

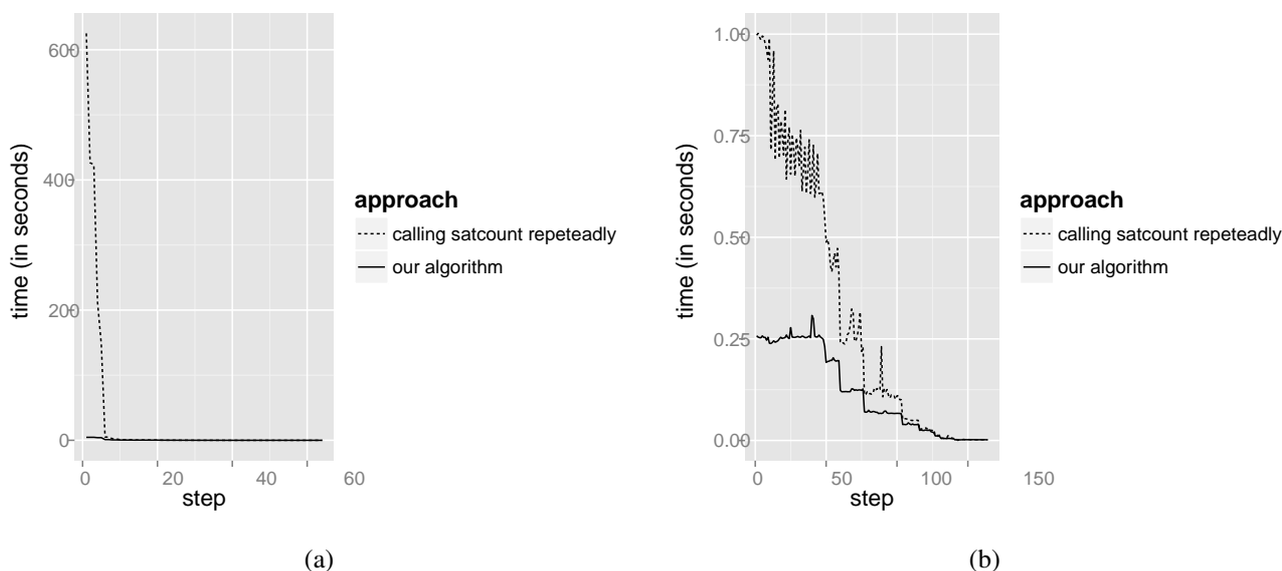
5.4. Threats to Validity

A threat to the validity of our approach is the time required to compute the component probabilities, which is also the Achilles' heel for Chen *et al.*'s [10] approach and Mazo *et al.*'s [9] Heuristic 3. For the sake of interactivity, configurators must provide customer guidance in a short time, and the usual way to compute the probabilities is highly time consuming (see Section 4.1.2.). To assess the response time of our algorithm (see Section 4.2), we determined the time needed to configure 1,000 randomly generated derivatives for Case Studies 1 and 2 using our entropy-driven approach. Figure 7 compares the average times needed to completely configure the derivatives by computing the component probabilities using our algorithm and calling repeatedly the BuDDy function, *satcount*. The performance tests were conducted on an Intel® Core™ 2 i3-4010U with 1.7 GHz and 4GB RAM (although, only one core was used).

As Figure 7 shows, our algorithm greatly improves the probability computation time. For instance, it requires 4.54 seconds on average to compute all component probabilities (and thus, their entropy values) for the first configuration step in the Renault Megane example. In contrast, calling *satcount* repeatedly consumes 625.18 seconds.

Note that the first configuration's steps are the most expensive in time. As the configuration process advances, the configuration space gets reduced and, so, the time needed to compute the probabilities. There is a point where both approaches converge and get response times close to zero.

Figure 7. Time required to compute component probabilities. (a): Renault Megane; (b) Electronic Shopping.



6. Conclusions

To satisfy a wide range of customers, product platforms must provide a high variety of optional components. For this reason, the configuration of all, but trivial, derivatives involves considerable effort in selecting which components they should include, while avoiding violations of the inter-component dependencies and incompatibilities. Our approach enriches existing automated configurators by reducing the number of steps required to configure a valid derivative.

Applying the information theory concept of entropy, our approach takes advantage of the fact that, due to the inter-component constraints, some decisions may be automatically derived from other decisions previously made. Therefore, the order in which decisions are made has a strong influence on the number of decisions required to complete a configuration. Moreover, our approach does not provide a static ordering that the customer is forced to follow. On the contrary, it suggests orderings dynamically, reacting to the customer decisions. In addition, we have proposed an algorithm that efficiently computes the variable probabilities of a Boolean formula, supporting in this way not only our approach, but also other methods proposed in related work.

The Renault Megane and Electronic Shopping configuration benchmarks have been used to test the applicability of our approach and its effectiveness. In particular, it has been shown that our approach needs less configuration steps than related work.

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Author Contributions

This is a joint effort based on an original idea conceived by Ruben Heradio and David Fernandez-Amoros. The development of the idea, its implementation and validation has been performed by Ruben Heradio, David Fernandez-Amoros, Hector Perez-Morago, and Antonio Adan.

Conflicts of Interest

The authors declare no conflict of interests.

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