How to Enumerate Your Keys Accurately and Efficiently After a Side Channel Attack

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Abstract. Side channels provide additional information to skilled adversaries that reduce the effort to determine an unknown key. If sufficient side channel information is available, identification of the secret key can even become trivial. However, if not enough side information is available, some effort is still required to find the key in the key space (which now has reduced entropy). To understand the security implications of side channel attacks it is then crucial to evaluate this remaining effort in a meaningful manner. Quantifying this effort can be done by looking at two key questions: first, how ‘deep’ (at most) is the unknown key in the remaining key space, and second, how ‘expensive’ is it to enumerate keys up to a certain depth?

We provide results for these two challenges. Firstly, we show how to construct an extremely efficient algorithm that accurately computes the rank of a (known) key in the list of all keys, when ordered according to some side channel attack scores. Secondly, we show how our approach can be tweaked such that it can be also utilised to enumerate the most likely keys in a parallel fashion. We are hence the first to demonstrate that a smart and parallel key enumeration algorithm exists.

1 Introduction

Side channel attacks have proven to be a hugely popular research topic, as the proliferation of new venues such as CHES, COSADE and HOST shows. Much of the published research is about key recovery attacks utilising side channel information. Key recovery attacks essentially take a number of side channel observations, colloquially referred to as ‘traces’, and apply a so-called distinguisher to traces that assigns scores to keys. An attack is considered (first-order) successful given a set of traces, if the actual secret key receives the highest score. Besides describing methods (i.e. the distinguishers) that recover the secret key from the available data, papers focus on the question of how many traces are required for successful first-order attacks.

The trade-off chosen in most work is, hence, to increase the number of traces to ensure that the secret key is recovered successfully with almost certainty. As observed by Veyrat-Charvillon et al. [16] in their seminal paper on optimal key enumeration, this might not be the trade-off that a well resourced adversary
would choose. Suppose that access to the side channel is scarce or difficult. In such a case the actual secret key might not have the highest score after utilising the leakage traces, but it might still have a higher score than many other keys. Now imagine that the adversary can utilise substantial computational resources. This implies that by searching through the key space (in order of decreasing scores; we call this smart key enumeration) the adversary would find the secret key much faster than by a naïve brute-force search (i.e. one that treats all keys as equally likely). Consequently, the true security level of an implementation cannot be judged solely by its security against first-order side channel attacks. Instead it is important to understand how the number of traces impacts on the effort required for a smart key enumeration.

We now illustrate this motivation by linking it to evaluating the impact of the most influential type of side channel attack: Differential Power Analysis.

1.1 Evaluating Resistance against Differential Power Analysis

Differential Power Analysis (DPA) [10] consists of predicting a so-called target function, e.g. the output of the Substitution Boxes, and mapping the output of this function to ‘predicted side channel values’ using a power model. For this process it is not necessary to know or guess the whole secret key, $SK$. One only needs to make guesses about ‘enough bits’. The predicted values for a key chunk are then ‘compared’ to the real traces (point-wise) using a distinguisher. Assuming enough traces are available, the value that represents the correct key guess will lead to a ‘higher’ distinguishing value. In Kocher et al.’s original paper [10] this was illustrated for the DES cipher, but most contemporary research uses AES as running example.

With respect to AES: Kocher’s attack consists of using a t-test statistic as a distinguisher to compute scores for the values of each 8-bit chunk of the 128-bit key; see Fig. 1 for a visual example. Here we have $m = 16$ chunks, each containing $n = 256$ values, with associated distinguishing scores as derived via a t-test statistic. In the graphical illustration, the secret key values are marked out in grey.

If sufficient side information is available, the values of the chunks that correspond to the secret key will have by far the highest distinguishing scores, such as the majority of key chunks in our graphical illustration. In this case the secret key can be trivially found (it is the concatenation of the chunks that lead to the uniquely highest score). However, if less side information is available, the scores may not necessarily favour a single key. Nevertheless, an adversary is still able to utilise these scores to smartly enumerate and then test keys (by using a known plaintext-ciphertext pair).

Security evaluations Considering the perspective of a security evaluator, it is obviously important to characterise the remaining security of an implementation after leakage. The evaluator (who can run experiments with a known key, and a varying number of traces) wants to compute its position in a ranked list of all
keys. Knowing this position allows the evaluator to assess the amount of effort required by an adversary performing a smart search (given some distinguishing vectors). Ideally, the evaluator is able to compute the ranks of arbitrarily deep keys.

**Accuracy and efficiency are key requirements:** Naturally, because the evaluator performs concrete statistical experiments, a single run of a single attack is not sufficient to gather sound evidence. In practice, any side channel experiment needs to be repeated multiple times by the evaluation lab, and many different attacks need to be carried out, utilising different amounts of side channel traces. Having the capability to determine the position of the key in a ranked list accurately (rather than just giving an estimation), and efficiently, is crucial to correctly assess the effort of a real world adversary. Previous works’ algorithms [17, 7] were capable of estimating the key rank within some bound. We demonstrate that we are accurate when enough precision is used, and importantly, we put forward the first approach for parallel and smart key enumeration.

### 1.2 Problem statement and notation

We use a bold type face to denote multi-dimensional entities. Indices in superscript refer to column vectors (we use the variable $j$ for this purpose), and indices in subscript refer to row vectors (we use $i$ for this purpose). Two indices $i, j$ refer to an element in row $i$ and column $j$. To maintain an elegant layout, we sometimes typeset column vectors ‘in line’, and then indicate transposition via a superscript $k = (\ldots)^T$.

We partition a key guess $k$ into $m$ chunks, each able to take one of $n$ possible values, i.e. $k = (k^0, \ldots, k^{m-1})$, and $k^j = (d_{0,j}, d_{1,j}, \ldots, d_{n-1,j})^T$. After exploiting some leakage $L$ all chunks $k^j$ have some corresponding score vectors, i.e. we know the score for each guess $k_{i,j}$ is $d_{i,j}$ after leakage. For convenience we use the variable $sk_j$ to refer to the indices (in each chunk) that correspond to the

\[
\begin{array}{cccc}
   0 & k^0 & k^1 & k^2 & k^{n-1} \\
   0 & 0.01 & 0.01 & 0.40 & 0.02 \\
   1 & 0.03 & 0.13 & 0.05 & 0.03 \\
   2 & 0.41 & 0.11 & 0.20 & \cdots & 0.31 \\
   3 & 0.38 & 0.27 & 0.07 & \cdots & 0.33 \\
   \vdots & \vdots & \vdots & \vdots & \vdots \\
   n-2 & 0.11 & 0.01 & 0.30 & 0.12 \\
   n-1 & 0.08 & 0.02 & 0.01 & 0.04 \\
\end{array}
\]

Fig. 1: Score vectors for $m$ key chunks. Each chunk can take values from 0 to $n-1$, and scores $d_{i,j}$ are on a scale that depends on the side channel distinguisher. The values that correspond to the (hypothetical) secret key are highlighted in grey.
correct key, i.e. \( SK = (k_{sk_1,1}, k_{sk_2,2}, \ldots, k_{sk_{m-1},m-1}) \). The score \( D \) of the secret key is then \( D = \sum_{j=0}^{m-1} d_{sk,j} \). We will later map scores to (integer) weights and the weight of the secret key will be \( W \).

Without loss of generality we assume that if several keys have the same score as the secret key, the secret key is ranked highest.

**Definition 1 (Key rank computation).** Given \( m \) vectors of \( n \) distinguishing scores, and the score \( D \) of the secret key \( SK \), count the number of keys with score strictly larger than \( D \).

**Definition 2 (Smart key enumeration).** Given \( m \) vectors of \( n \) distinguishing scores, list the \( B \) keys with the highest score.

### 1.3 Outline and our contributions

In a nutshell, we utilise an elegant mapping of the key rank computation problem to a knapsack problem, which can be simplified and expressed as (efficient) path counting. As a result, we can compute accurate key ranks, and importantly, this enables us to put forward the first algorithm that can perform smart key enumeration in a parallel manner.

Our contribution is structured in four main sections as follows:

**Casting the key enumeration as an integer knapsack.** In Sect. 2 we show how to cast the key enumeration problem as a solution to counting knapsack solutions. In particular, we develop the representation of key rank as a multi-dimensional knapsack, and discuss its resulting graph representation.

**An accurate key rank algorithm.** In Sect. 3 we map the multi-dimensional knapsack to a directed acyclic graph. We can therefore count solutions to the multi-dimensional knapsack problem by counting paths in the graph. The restriction of picking one item per chunk keeps the number of vertices in the directed acyclic graph small. As the graph is compact, and each node has at most two outgoing edges, the path counting problem can be solved efficiently in \( O(m^2 \cdot n \cdot W \cdot \log n) \).

**Smart key enumeration.** In Sect. 4, with the additional book-keeping of storing the vertices we visit, we can enumerate the \( B \) most likely keys with complexity \( O(m^2 \cdot n \cdot W \cdot B \cdot \log n) \). We then show several techniques to make this process as efficient as possible.

**Practical evaluation and comparison with previous work.** In Sect. 5 we discuss requirements around precision. The main factor that influences performance is the size of the key rank graph, which is determined by the precision of the initial mapping and the weight of the target key. We compare our work with previous works in terms of precision and speed with regards to the key rank algorithm, and in terms of speed with regards to smart key enumeration.
1.4 Previous work

**Key Rank** Veyrat-Charvillon *et al.* [17] were the first to demonstrate an algorithm to estimate the rank of the key without using full key enumeration. The search space can be represented as a multidimensional space, with each dimension corresponding to a key chunk, sorted by decreasing likelihoods. The space can be divided into two, those keys ranked above the target and those ranked below. Using the property that the ‘frontier’ between these two spaces is convex, they are able to ‘trim’ each space down until the key rank has been estimated to within 10 bits of accuracy.

Bernstein *et al.* [2] propose two key ranking algorithms. The first is based on [17] and adds a post processing phase which has been shown to tighten the bounds from 10 bits to 5 bits. The second algorithm ranks keys using techniques similar to those used to count all $y$-smooth numbers less than $x$. By having an accuracy parameter they are able to get their bounds arbitrarily tight (at the expense of run time).

Glowacz *et al.* [7] construct a novel rank estimation algorithm using convolution of histograms. Using the property that $S_1 + S_2 := \{x_1 + x_2 | x_1 \in S_1, x_2 \in S_2\}$ can be approximated by histogram convolution, by creating a histogram per key chunk and convoluting them all together, they are able to efficiently estimate the key rank to within 1 bit of precision.

Duc *et al.* [5] perform key rank using a method inspired by Glowacz *et al.* [7]. They repeatedly ‘merge’ the data in one column at a time (as the histograms were convoluted in one at a time). Each piece of information is downsampled to one of a series of discrete values (similar to putting into a histogram bin). The major difference is that instead of just downsampling the originial data, they also downsample after each key chunk is merged in.

**Key Enumeration** Veyrat-Charvillon *et al.* [16] propose a deterministic algorithm to enumerate keys based on a divide-and-conquer approach. Using a tree-like recursion (starting with two subkeys, then four, all the way to sixteen) and keeping track of what they call the frontier set (similarities can be drawn to the frontier of Veyrat-Charvillon *et al.* [17]), they are able to efficiently enumerate keys.

Ye *et al.* [18] present what they describe as a Key Space Finding algorithm. A Key Space Finding algorithm takes in the distinguishing score vector and returns two outputs: the minimum verification complexity to ensure a desired success probability, along with the optimal effort distributor which achieves this lower bound. Given this it is straightforward to run a (probabilistic) key enumeration algorithm. The distinguisher intuitively moves the boundary of which subkeys to enumerate until the desired probability is achieved.

**Reflecting on the approaches taken by previous work** All of the previous work has treated key rank and key enumeration as two disjoint problems and hence approached them using different techniques. For instance, it is unclear how
to extend the existing key rank algorithms to enumerate keys, and conversely, it is
not apparent how to simplify the enumeration algorithms to compute key ranks
efficiently (i.e. without just counting as you enumerate). We however believe
that both of these problems are highly similar in nature and by maintaining
some structure within the key rank it should be possible to enumerate without
making the ranking inefficient. In the remainder of the paper we explain how to
do just that: efficient key ranking with enough structure to make (fully parallel)
enumeration possible.

2 Casting the key enumeration as a knapsack

We now explain how the key enumeration problem can be formulated as a variant
of a knapsack problem. In its most basic form a knapsack problem takes a set
of $n$ items that have a profit $p_i$ and a weight $w_i$. A binary variable $x_i$ is used
to select items from the set. The objective is then to select items such that the
profit is maximised whilst the total weight of the items does not exceed a set
maximum $W$:

\[
\text{maximize: } \sum_{i=0}^{n-1} p_i \cdot x_i \\
\text{subject to: } \sum_{i=0}^{n-1} w_i \cdot x_i \leq W \\
x_i \in \{0, 1\}, \forall i
\]

The counting knapsack (#knapsack) problem is then understood to be the
associated counting problem: given a knapsack definition, count how many so-
lutions there are to the knapsack problem.

Intuitively, we should be able to frame the key rank computation problem as
a knapsack variant. In contrast to a basic knapsack, however, we have classes of
items (these are the distinguishing vectors $k^j$), profits can be dropped since we
are counting the number of solutions, and we must take exactly one item from
each class. The weight $w_{i,j}$ for each item can be derived\(^1\) from the distinguishing
score $w_{i,j} = \text{MapToWeight}(d_{i,j})$ in such a way that higher distinguishing scores
lead to lower weights\(^2\). We define the maximum weight $W$ as the sum of the
weights associated with the secret key chunks, i.e. $W = \sum_{j=0}^{m-1} w_{sk_{j,j}}$. Recall
that we assume if several keys have weight $W$ the secret key (which must be
among those) is listed first. This enforces $W$ as a strict upper bound in the
knapsack definition.

\(^1\) For the sake of readability, we do not discuss the implications of needing to map
distinguishing scores (which are floating point values) to weights at this point, but
refer the reader to Sect. 5.1 for a discussion.

\(^2\) This ensures compatibility with knapsack notation.
The multiple-choice knapsack problem that identifies keys with weight lower than $W$ is then defined as follows:

$$
\sum_{j=0}^{m-1} \sum_{i=0}^{n-1} w_{i,j} \cdot x_{i,j} < W
$$

$$
\sum_{i=0}^{n-1} x_{i,j} = 1, \forall j
$$

$$
x_{i,j} \in \{0,1\}, \forall i,j
$$

The first constraint ensures that all keys (i.e. selections of items) have a weight lower than the secret key. The second constraint ensures that only one item per distinguishing vector is selected. The counting version of this multiple-choice knapsack equals to computing the key rank.

Counting solutions to knapsack problems in general is known to be a computationally hard problem, and known classical solutions [6] rely on combinations of dynamic programming and rejection sampling to construct an FPRAS. Gopalan et al. [8] more recently utilise branching programs for efficient counting, and we took inspiration from this paper to approach the solution to our counting problem.

To illustrate our solution, we have to slightly modify the knapsack representation. It will be convenient to express the multiple-choice knapsack as a multi-dimensional knapsack variation as follows. Each key chunk corresponds to ‘one dimension’. Each item $k_{i,j}$ has an associated weight vector $w_{i,j}$ of length $m + 1$ of the form $(w_{i,j}, 0, \ldots, 1, 0, \ldots, 0)$, where the 1 is in position $j$. The global weight is also expressed as a vector $W = (W, 2, \ldots, 2)$ of length $m + 1$.

The key rank problem is then to count the number of solutions (that satisfy all constraints simultaneously) to

$$
\sum_{j=0}^{m-1} \sum_{i=0}^{n-1} w_{i,j} \cdot x_{i,j} < W
$$

$$
x_{i,j} \in \{0,1\}, \forall i,j
$$

The constraint $W$ ensures that all keys that are counted have a strictly lower weight than the secret key. If the weight vector has a 1 in position $j$, it means that this is a value for the $j$th key chunk. Since the weight limit is 2 in the constraint vector $W$, it means that only a single value for any key chunk can be chosen. We now illustrate this by a simple example.

**Example 1.** Our illustrative example, which will run throughout the paper, consists of two distinguishing vectors with three elements each: $k^0 = (0, 1, 3)^T$, and $k^1 = (0, 2, 3)^T$. We assume that the secret key, $SK$, is $(2, 1)$. First we derive the global weight constraint vector. In this case it has length $m + 1 = 3$ and
contains the maximum weight \( W = w_{0.2} + w_{1.1} = 3 + 2 = 5 \), which results in \( W = (5, 2, 2) \). The weight vectors of the \( k_{i,j} \) are:

\[
\begin{align*}
  w_{0,0} &= (0, 1, 0), \quad w_{0,1} = (1, 1, 0), \quad w_{0,2} = (3, 1, 0) \\
  w_{1,0} &= (0, 0, 1), \quad w_{1,1} = (2, 0, 1), \quad w_{1,2} = (3, 0, 1)
\end{align*}
\]

Given that \( W = 5 \), all except two of the combinations are below this threshold. Hence the solutions to the knapsack are:

\[
\begin{align*}
  (k_{0,0}, k_{0,1}), \quad (k_{0,0}, k_{1,1}), \quad (k_{0,0}, k_{2,1}), \\
  (k_{1,0}, k_{0,1}), \quad (k_{1,0}, k_{1,1}), \quad (k_{1,0}, k_{2,1}), \\
  (k_{2,0}, k_{0,1})
\end{align*}
\]

Notice that the knapsack solution will never contain the secret key itself, as it returns all keys with weight strictly less than the weight of the secret key. For the ranking problem this would give us that our secret key has rank 8.

For standard knapsack problems it is well known [4] that solutions can be found via finding longest paths on a directed acyclic graph. In the following section we will show that such a graph exists also for our knapsack, and importantly, that the resulting graph allows for a particularly efficient path counting solution, which gives us a solution to the Key Rank problem.

### 3 An accurate key rank algorithm

In this section we first define the graph and illustrate how it relates to the multi-dimensional knapsack via intuition and a working example. We then explain our fast path counting algorithm for a compact representation of the graph.

#### 3.1 Key rank graph

Recall that our multi-dimensional knapsack has \( m \cdot n \) elements, and for each element we have a weight vector. Also, a correct solution to the multi-dimensional knapsack must have a weight that is strictly smaller than \( W \). Since we need to be able to represent all permissible solutions we need \( W \) extra vertices (per element). This means that we ‘encode’ all solutions to the knapsack in a graph with \( m \cdot n \cdot W \) vertices (plus an extra two for accept and reject nodes). The vertices corresponding to item \( k_{i,j} \) are labelled \( V_{w_{i,j}} \), where the variable \( w \) denotes the ‘current weight’. The key rank graph contains a start node \( S \), an accept node \( A \) and a reject node \( R \). The edges are constructed as follows:

- \((V_{w_{i,j}}^{w_{i,j}}, V_{w_{i,j}+1})\) which corresponds to the item not being chosen in this set
- \((V_{w_{i,j}}^{w_{i,j}+w_{i,j}}, V_{w_{i,j}+w_{i,j}})\) if the item is chosen for this set and \( w + w_{i,j} < W \)
- \((V_{n-1,j}^{w_{i,j}}, R)\) if no elements are chosen from the set
– \((V_{w_{i,m-1}}^w, A)\) if the item is chosen for the last set and \(w + w_{i,m-1} < W\)
– \((V_{i,j}^w, R)\) if the item is chosen for this set and \(w + w_{i,j} \geq W\)
– \(S = V^0_{0,0}\) to set up the start node

When visualising the key rank graph it will be convenient to think of the indices \(i, j\) as though they are ‘flattened’ (i.e. they are topologically sorted and occur in a linear order). In this representation the graph is \(n \cdot m\) deep, \(W\) wide, where the width of the key rank graph essentially tracks the current weight (of the partial keys). Each vertex has exactly two edges coming out of it (with the exception of \(A\) and \(R\)): either the vertex was ‘included’ (this corresponds to the choice of selecting the corresponding value of the key chunk to become part of the key) or not. If the answer is yes then the edge must point to the first item in the next key chunk, as we can only choose one item per key chunk, and the weight must be incremented by the weight of this key item. If the item/vertex is not chosen then the edge must go to the next item in the chunk (or reject if this was the last item) and the weight must not be incremented as the item was not selected. For any partial key, if a new key chunk is added and the new weight exceeds \(W\) then this is not a valid key and thus the path will go to the reject node \(R\).

**Graph Example** To illustrate the working principle we provide in Fig. 2 the process of constructing the graph for the example provided above.

Initially the graph is constructed (top right) and the start node is initialised based on the rule \(S = V^0_{0,0}\). The width of the graph is set to be 5 (0-4) as this matches our maximum weight (\(W = 5\)). The depth of the graph becomes 6 as each chunk contains 3 items.

Next (middle left) the first two children from the start node are created. The edge that denotes the chunk is, in fact, selected (the right child) is built following the rule \((V_{0,0}^0, V_{0,1}^0)\), which creates the edge from the start node, to an element in the next chunk. The edge that denotes the chunk is not selected (the left child) is built following the rule \((V_{0,0}^0, V_{1,0}^0)\), which creates the edge from the start node to the next item within the same chunk.

Moving onto the following step (middle right), children continue to be created through the same set of rules. However, note that at the point a link is created to the accept node based on the rule \((V_{0,1}^0, A)\), this demonstrates that the selection of key chunk 0 followed by 0 is a valid solution to the problem.

In the following steps links continue to be created based on the rules until all paths in the graph are created. Please note that throughout the construction of the graph, the last item in each chunk will have a left child that points to reject (as obviously there are no further chunks to select) but these have been omitted from the example diagram for the sake of clarity.

All the greyed out nodes also have their children calculated. However, as they do not alter the path count, we have excluded them from the example figures to aid clarity.

Each path from \(S\) to \(A\) if, corresponds to a key with lower weight than our secret key. Thus, counting these paths will yield the rank of the secret key. While
in general path counting is hard [15], we explain how our graph structure, having
at most two outgoing edges per node, lends itself to efficient counting.

3.2 Counting valid paths

Clearly our key rank graph is a directed acyclic graph. We have already men-
tioned that it is convenient to ‘flatten’ the graph (as it has been presented in
the example). This ‘flat’ graph is also more suited for an efficient counting al-
gorithm. Hence, from this point onwards we will now assume that the graph
is topologically sorted\(^3\). The start node will be labelled 1 and the final node
will be labelled A. There are \(n \cdot m \cdot W + 2\) vertices in the graph, we have that
\(A = n \cdot m \cdot W + 2\) and \(R = A - 1\).

We also assume two constant time functions \(LC(\cdot), RC(\cdot)\) which return the
index of the left child and right child, respectively. The algorithmic descriptions
of these functions can be found in App. B for our particular graph. We therefore
have the following recurrence relation, where \(PC\) is a vector and \(PC[i]\) stores
the number of accepting paths from \(i\) to \(A\):

\[
PC[c] = \begin{cases} 
1, & \text{if } c = A, \\
0, & \text{if } c = A - 1, \\
PC[LC(c)] + PC[RC(c)], & \text{otherwise.}
\end{cases}
\]

The total number of paths between 1 (our start node) and A (our accept
node) is then simply \(PC[1]\). This recurrence relation forms the algorithm given
in Fig. 1, which assumes that \(LC, RC\) are globally accessible functions.

**Algorithm 1** The key rank algorithm

\[
\begin{align*}
PC[A] & \leftarrow 1 \\
\text{for } c = A - 1 \text{ to } i = 1 \text{ do} & \\
PC[c] & \leftarrow PC[LC(c)] + PC[RC(c)] \\
\text{end for} \\
\text{return } PC[1]
\end{align*}
\]

For an example of this, see Fig. 3. This figure shows that the vector is tra-
versed from the end back to the start, and cells are filled by summing the values
in their left and right children cells. For clarity in the figure, we only show
example links between two cells, whereas in practise they are present on all.

\(^3\) It turns out that because the path counting for the key rank graph is already ex-
tremely fast and memory efficient, the choice of sorting is irrelevant, bar the excep-
tion that \(S\) must be the first node and \(A\) must be the final node. However, when it
comes to key enumeration this will be an important consideration and thus will be
discussed in further detail in the corresponding section.
Fig. 2: An example showing the construction of the graph for the small example instance provided
Correctness The base case of $PC[A] = 1$ is self explanatory; there is exactly one path from $A$ to $A$, the path involving no edges. From an arbitrary node $c$, it is possible to traverse the edge to the left child (and thus take however many paths start there) or traverse the edge to the right child (and do the same) and we conclude that $PC[c] = PC[LC(c)] + PC[RC(c)]$. We can iterate over all nodes starting at the final node $A$ and working backwards until we reach the start node, and since our graph is topologically sorted when we are operating on a node, the values for the nodes’ children will already have been calculated as they come later in the topological sorting.

We note at this point that this counting algorithm is exact. However, as we pointed out before, we need to convert floating point distinguishing scores to integer weights, and this conversion may incur a loss of precision and hence could cause a loss of accuracy. We discuss this in Sect. 5.

Time complexity The time complexity of the key rank algorithm depends on the number of vertices in the key rank graph and their size. Our graph contains $A = m \cdot n \cdot W$ vertices. The integers stored in the vertices could be up to $O(2^A)$ (and thus be of size $O(A)$) because in the worst case each value can be double the previous value (if $PC[LC(i)] = PC[LC(i)]$). Hence, given that we have $A$ vertices and perform an integer addition with an $A$-bit variable for each vertex, we have worst-case time complexity of $O(A^2) = O(m^2 \cdot n^2 \cdot W^2)$.

However, whilst we touch each vertex once, we know that there are at most $O(n^m)$ keys. Consequently, we need no more than $O(m \log n)$ bits to store the path count (in contrast to the $O(A) = O(m \cdot n \cdot W)$ bits for the worst case). Hence the time complexity for computing the key rank via the key rank graph is $O(m^2 \cdot n \cdot W \cdot \log n)$.

It is worth noting that the key depth does not factor into the time complexity and the following example will help to clarify this. Consider the target key which has weight 1 in every column (which gives $W = 16$ and a grid with 65536 nodes). If all other key chunks have weight 0 then the target key will have rank $2^{128}$ since all other keys have a lower weight. However, if all other key chunks have weight...
Algorithm 2 The key enumeration algorithm

\[ KL[A] \leftarrow \emptyset \]

\textbf{for} \( c = A \) to 1 \textbf{do}

\[ KL[c] \leftarrow (value(c), KL[RC(c)]) \cup KL[LC(c)] \]

\textbf{end for}

\textbf{return} \( KL[1] \)

2, then our target key will have rank 0 because all other keys have a higher weight. None of the other values affect the size of the graph and thus it is clear that the runtime is not changed by the key depth.

In fact for AES-128 the values of \( m, n \) are also fixed and thus we get that the algorithm runs in \( O(W) \), that is to say it is linear in the weight of the secret key. See Sec. 5.1 for experiments supporting this.

4 Parallelisable key enumeration algorithm

We are able to further modify our algorithm such that with minor (standard book-keeping) adjustments, we are able to list all valid paths, as opposed to just counting them, with reasonable efficiency. The algorithm is given in Alg. 2 and requires an additional (constant time) function call \textit{value} which, given an index \( c \), returns the value of a vertex. We write \( (a, \{x_c\}_c) \) to mean \( \{(a, x_c)\}_c \). That is to say if we concatenate an item \( a \) with a set, we are really concatenating the item \( a \) with every item in the set to form a new set.

It is now easy to use this for key enumeration. Assume we have the resources to enumerate/test up to \( B \) keys. Then, we choose some weights (which correspond to key guesses) and use the key rank algorithm to determine their ranks and compare them to \( B \). This allows us to quickly select the appropriate \( W \) for the given \( B \). Then, Alg. 2 proceeds as follows: for any valid path (in the key rank graph), every time a right child is taken (this can be determined by the node indices) the corresponding value for the respective key chunk is chosen. A left child means that we are not taking a particular value for key chunk. In this manner the keys are effectively reconstructed from the key rank graph.

If one wanted to enumerate the keys in a smart order, this would simply be a case of altering the construction of the tree which stores the valid key chunks for enumeration. Currently the valid key chunks are stored in \textit{numerical} order within the tree, however if this was changed such that they were stored in order of scores, the keys would be rebuilt in a near optimal order.

In the rest of this section we discuss run time and memory requirements. Whilst the run time is bounded by the number of keys that we want to enumerate, we show there are different strategies to improve the memory performance. Finally, we show that with a further simple observation, we can parallelise the key enumeration algorithm.
4.1 Time complexity

We begin with a worst case analysis considering a general graph. In this case, the enumeration algorithm would be exponential in the length of the number of vertices, because to generate all paths (each vertex has two children) the algorithm clearly must take $O(2^n)$ time.

However, in our key rank graph, each path corresponds to a valid key with weight lower than $W$. Considering this, the run time of this algorithm is relative to the rank (which is determined by $W$) and not to the total number of keys; hence this algorithm can be used to enumerate keys for a given workload in time $O(m^2 \cdot n \cdot W \cdot B \cdot \log n)$. This is because all $O(n \cdot m \cdot W)$ nodes are touched once, and $B$ keys are reconstructed which are of length $m \cdot \log n$.

4.2 Memory efficiency

How we topologically sort the key rank graph has a major impact on the memory efficiency of the key enumeration. While there are a variety of explicit topological sorting algorithms in the literature [9, 14], we are able to avoid explicit sorting because we know our graph structure in advance. Hence, we show that our graph can be sorted implicitly by how the nodes are numbered within the calculation of the left and right child functions. The remaining question is what method of sorting is the most desirable.

In Fig. 4 we give the three options of topologically sorting the example graph previously considered in Fig. 2. Appendix B gives the pseudo-code for the left and right children functions for each of the three topological sorts given below. We discuss the pros and cons of each of the sorting methods. We also discuss how to improve memory efficiency further by appropriately storing the generated keys.

**Wide Sort** In this sorting the graph is numbered one chunk at a time, one item at a time, along the weight in increasing order (see Fig. 4a). Formally given a chunk, item and weight $(x, y, z)$ the index is $i = x \cdot W \cdot n + y \cdot W + z$. This is a valid topological sorting of the graph, since a nodes' children will be either one item lower in the same chunk (for the left child) or the the first item in the next chunk (right child) both of which have a higher number.

This is the topological sort we described for key rank. Note that, since key rank is extremely fast we describe the most intuitive sort since it did not have an impact on performance, while with enumeration this is no longer the case and must now be taken into consideration.

The advantage of this sorting is that it is due to the fact an element will only need to look at the item below and the item at the top of the next chunk; these are the only things needing to be stored in memory. This makes it very memory efficient requiring $O(W)$ memory.

The disadvantage of this method is that it is highly serial and it does not seem possible to (easily) parallelise.
Long Sort In this sorting the graph is numbered one weight at a time, one chunk at a time, along the item in increasing order (see Fig. 4b). Formally given a chunk, item and weight \((x, y, z)\) the index is \(i = z \cdot n \cdot m + x \cdot n + y\). This is a valid topological sorting of the graph, since the left child will have the same weight but higher item number and thus will have a higher number, while the right child will have either the same weight or a higher weight for the next chunk and thus will have a higher number.

The advantage with this sorting is that (with careful synchronisation) it can be made parallel along the weights. Since the left child is always in the same weight this requires no synchronisation as it will be computed by the same process. The only time that synchronisation is needed is when the right child is calculated because this could have larger weight and thus would be calculated by a different process. Therefore it is required that for a process to calculate a right child all processes running with higher weights have finished with the previous chunk before this process can begin.

The disadvantage with this method is that no clever memory tricks can be performed and thus the whole thing needs to be stored in memory at once, requiring \(O(n \cdot m \cdot W)\) space.

Wiggly Sort In this sorting the graph is numbered one chunk at a time, one weight at a time, along the item in increasing order (see Fig. 4c). Formally given a chunk, item and weight \((x, y, z)\) the index is \(i = x \cdot W \cdot n + z \cdot n + y\). This is a valid topological sorting because the left child will be one further down within the same weight and within a chunk we are calculating items down the same weight. The right child will be in the next chunk and we are calculating one chunk at a time.

This sorting combines the advantage of the previous two with none of the disadvantages. Within each chunk the individual weights can be run in parallel because it either relies on information in the same weight or information from item 0 in the previous chunk (which has already been calculated). Hence as long as all threads synchronise at item 0 in each chunk the rest can be calculated in parallel. Since we only require information in our current chunk and the item 0 from the previous chunk, memory can be reused giving a memory requirement of \(O(n \cdot W)\) which, while not as good as the first (serial) method, is better than the second (parallel) method.
Fig. 4: Different options for topological sorting for our previous example. Note that the deepest node in each chunk will be guaranteed to have a left child leading to $R$; for clarity these paths are omitted.
Fig. 5: The key tree for all possible three character keys containing ‘A’ or ‘B’

**Key Storage** The topological sorting of the graph is clearly a crucial factor for memory efficiency. The other factor is how keys are represented/stored within our graph.

In the algorithm as described all (partial) keys are stored at each point in the algorithm. This will become very inefficient. Consider, for example, the case where you want to enumerate all keys. There are $2^{120}$ keys which have the first key chunk set to zero (hence this chunk would be duplicated $2^{120}$ times). Clearly, one needs to choose an appropriate data structure, and we use a tree, see Fig. 5. This key tree is passed to a separate algorithm that converts it into a series of keys for testing. The advantage of this is threefold. First, it greatly speeds up the enumeration. Second, the conversion of the key tree into a list of keys is trivially parallelisable, and third, the actual testing (in our case checking the AES encryption using a given plaintext/ciphertext pair) can be amortised into this cost.

### 4.3 Parallelisation

We can achieve parallelisation with a simple observation: by adjusting the graph such that instead of vertices with a weight lower than $W$ going to the accept state, we only allow vertices with weight in the range between the two weights $W_1$ and $W_2$ to reach the accept state. The width of the graph is defined by $W_2$; $W_1$ has no impact on the graph size. This results in an algorithm that enumerates ‘batches’ of likely keys. Hence, one can run multiple instances of the key enumeration algorithm in parallel, where each instance enumerates a unique batch of keys.

All ranges of keys can be computed in parallel and require no communication between threads except for the initial passing of the distinguishing scores and a notification when the key has been found. It is hence trivial to utilise multiple machines (or cores).

*Further speed optimisations.* Currently the algorithm operates on every node of the graph. However, some of the nodes are not even reachable from the start node (for example the greyed out nodes in Fig. 2). Hence any computation done
on these nodes is wasted because it will never be combined into a solution. By precalculating the number of valid paths from \( S \) to all other nodes in the graph (a reasonably cheap operation compared to a large key enumeration – this is done using the key rank algorithm), we can skip over a node if the number of paths from the start node to here is 0 because any work here will not be combined with the final solution.

5 Practical evaluation and comparison with previous work

Our key enumeration and key rank algorithms are both based on a graph representation of a multi-dimensional knapsack. To define this multi-dimensional knapsack it is necessary to map distinguishing scores, which typically are floating point values, to integer weights. This requirement has implications for the performance of our algorithms, as the time complexities for both algorithms strongly depend on the parameters \( m \) (the number of key chunks), \( n \) (the number of items per chunk), and \( W \) (the maximum weight). In particular, for any fixed key size (and number of chunks) the size of the graph (i.e. the width) grows with \( W \), and \( W \) grows with the precision that we allow for in the conversion from floating point values to integers.

We hence focus our practical evaluation on the impact of the precision\(^4\), on accuracy\(^5\), and on performance. First, we discuss the precision requirements for practical DPA outcomes. Second, we explore the practical impact on the performance of key rank when we increase the precision. Third, assuming we allow for sufficient precision, we ask what are the best performance results that we can achieve on single but also many core platforms for the key enumeration.

It is clear that to answer these questions we need to be able to generate many practically relevant distinguishing vectors in a manner that is comparable to previous work. We hence decided to adopt the simulator used by Veyrat-Charvillon et al. [16], which we explain in App. A.1 in more detail. In a nutshell, Veyrat-Charvillon et al. create distinguishing vectors based on attacking the AES SubBytes output, assuming noisy Hamming weight leaks, and using the Hamming weight as power model. Their DPA simulator allows to manipulate the level of noise, and the number of measurements. They output ‘additive’ scores (by taking the logarithm of the raw matching scores), which we pass directly to our \( \text{MapToWeight} \) function (see A.2).

5.1 Evaluating and comparing precision

In practical DPA attacks the combination of measured power traces, model values, number of traces and distinguisher will influence the effective precision of the

\(^4\) Precision is the ability to reproduce a measurement result, \( i.e. \) if several measurements of a variable give very close values then the measurement is precise.

\(^5\) Accuracy is the closeness of a measurement to a true value, \( i.e. \) this relates to the ‘trueness’ of a measurement.
distinguisher scores\textsuperscript{6} We discuss the mentioned factors briefly. Then we experimentally determine the necessary level of precision for our key rank algorithm, and compare this to the number of bins for the method by Glowacz et al. [7].

**Precision in factors influencing DPA outcomes.**

**Leakage traces:** Modern scopes offer between 8 and 12 bits of vertical resolution. Dedicated AC/DC converters may achieve up to 16 bits of vertical resolution; however, they can only achieve this at low sampling frequencies (hence modern scopes opt for lower resolution but allowing higher sampling frequencies). This vertical resolution, however, is for all the points in a trace. Consequently, the resolution in a single point will be much lower\textsuperscript{7}. The loss of precision can be compensated for by averaging over several measurements: this is effectively utilised by DPA attacks because typically many more power traces are measured than there are power models. In a well-known paper, Aziz et al. [1] derive that to increase the precision of some data by $b$ bits, the oversampling rate needs to be $4^b$.

**Power model:** Many practical DPA attacks use the Hamming weight or Hamming distance to map the target values to predicted power consumption values. In the case of profiled attacks, the literature reports so-called stochastic models [13] (which model power consumption as a linear function in the bits of the target) mainly for 8-bit targets, and template attacks [3]. Template attacks can be run in different ways [11] utilising between $t$, $2^t$ or even $2^t \times 2^t$ distinct power models for a $t$-bit target. Consequently, practical template attacks typically remain restricted to 8-bit targets, or do not utilise the full set of templates. This brief overview shows that the precision of the power model is actually quite low.

**Distinguisher:** We speculate that distinguishers that require ‘simple operations’, such as additions, subtractions, etc. of floating point values require less precision than distinguishers that e.g. include computing logarithms. Any confirmation for this hypothesis can only come from practical experiments, given the lack of work on this question.

**Experimentally measuring precision for key rank and Glowacz et al.**

We ran precision tests using Veyrat-Charvillon’s simulator (see A.1, using $N = 30$ and variance two) to determine the appropriate level of precision for further experiments. We plot the difference in ranking outcomes for increasing precision in Fig. 7 (left). In this figure, and in all figures that will follow, we plot outcomes of individual experiments in gray, and average outcomes in black. The $x$-axis show the precision in $w_{i,j}$. The $y$-axis refers to the change in ranking outcomes

\textsuperscript{6} Veyrat’s simulator stores values in variables with double precision (i.e. one has 53 bits of precision). But effectively, only a few of them are necessary to contain the effective precision.

\textsuperscript{7} In order to have the full resolution in a single point, the traces would need to be centred in this particular point.
Fig. 6: Impact of the distinguisher (left: correlation, right: Gaussian templates without log2) on the precision requirements when considering up to 16 bits of precision. 

when increasing the precision by 0.1 bits from the previous step. From 11 bits onwards the outcomes do not change anymore. Because our ranking method is exact with enough precision, we can infer that with 11 bits of precision in \( w_{i,j} \) we produce exact ranks. Already from 4 bits of precision (on average, as plotted in black) we are within five bits of accuracy from the real result. From about 8 bits onwards, increasing the precision changes the ranking outcomes by just under a bit for our algorithm.

We implemented the convolution based method by Glowacz et al. [7]. Their method is essentially based on building \( m \) histograms (one from each of the distinguishing vectors) and counting the keys by counting items in the ‘amalgamated’ histogram efficiently via convolution. Figure 7 (right) shows that they achieve very high average precision (plotted in black) from about 50,000 bins onward. We can therefore conclude that using 50,000 bins roughly corresponds to 11 bits of precision in \( w_{i,j} \). Glowacz et al. [7] actually recommend to use 500,000 bins in their paper.

Recall that we hypothesised that different distinguishers would lead to different precision requirements. To test this hypothesis we implemented two further distinguishers for the simulator: one distinguisher was based on correlation and one was based on Veyrat-Charvillon’s method but without applying the logarithm. Figure 6 shows the results for them, this time we allowed up to the 16 bit precision. The plots show that indeed, different distinguishers require different levels of precision, and that correlation has the least requirements.

To provide further evidence for the exactness of our ranking algorithm (provided enough precision), we considered the difference between the key rank output by our algorithm, and the key rank output by Glowacz et al.. In this experiment, we used 16 bits for our algorithm and 500,000 bins for Glowacz et al. Fig 8
Fig. 7: Bits of precision for Key Rank (left) and number of bins for Glowacz et al. (right).

shows the identical trend as Fig. 1 (right panel) of Glowacz et al. Hence the difference between our ranking outcomes and their ranking outcomes are identical to the rank estimation tightness that they measure, reinforcing the exactness of our ranking outcomes.

5.2 Evaluating and comparing run times for key rank

We explained in Sect. 3 that the run time of the Key Rank algorithm is independent of the actual depth of the key (see Table 1). The run time depends on the size of the graph, which is fixed for a certain choice of \( m \) and \( n \), and hence depends on the size of \( W \). Since \( W \) is derived from summing the weights of \( SK \), its precision will be determined by the precision that we allow during the conversion of the distinguishing scores.

We hence experimented with the relationship between run time and size of \( W \) and also precision. We did this by fixing all parameters for Veyrat-Charvillon’s simulator and only varying the precision allowed in the function \( \text{MapToWeight} \). As in the previous graph, we upper bounded the precision in \( W \) at 16 bits.

Table 1: Run times observed for calculating key depth with a fixed variance of 2, and 30 samples.

<table>
<thead>
<tr>
<th>Key Depth (log2)</th>
<th>Runtime (Seconds)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>0.2</td>
</tr>
<tr>
<td>20</td>
<td>0.2</td>
</tr>
<tr>
<td>30</td>
<td>0.2</td>
</tr>
<tr>
<td>40</td>
<td>0.2</td>
</tr>
</tbody>
</table>
Figure 9 shows how the run time increases for bigger $W$ (left) and more precision in $W$ (right). The run times for sufficient precision (i.e. 8 bits for $W$) are well below half a second. Even with 11 bits of precision (i.e. accurate ranking outcomes) our average run time is around 4 seconds. The plot shows that this average (black) is tracked well by the individual experiments (gray).

5.3 Evaluating and comparing run times for key enumeration

The run time of the key enumeration algorithm (as referred to by KEA in the graphs that will follow) is dominated by the depth to the key. Veyrat-Charvillon et al. [16] presented the current state of the art for smart key enumeration, and they kindly gave us access to the latest version of their implementation. We were hence able to run their code alongside ours. Therefore for all graphs that we provide in the following, the timings were obtained on identical platforms. Note that for all experiments, as the toolbox provided a known secret key on which the simulated attack was based, we knew at which point the enumeration had found the correct key without needing to perform an AES operation on a known plaintext/ciphertext pair (this is common within the literature).

Single core vs multi core comparisons. Figure 10 gives a comparison of run times of Veyrat-Charvillon et al.’s algorithm and our algorithm on a single core (left). We sampled multiple distinguishing scores for each key depth and ran our respective key rank algorithms. The graphs show that from key depths just under 30 bits onwards we clearly outperform Veyrat-Charvillon et al.’s algorithm, even on a single core. On the right, we provide some performance graphs when running our key enumeration algorithm on multiple cores. The graph shows that eight cores can enumerate $2^{40}$ keys in the same time as one core enumerates $2^{37}$, which is a vast difference. Also another result of note is a single core run enumerates $2^{38}$ keys in 13.9 hours and four cores performs the same enumeration in 6.4 hours.
A  Implementation Details

A.1  Veyrat-Charvillon’s distinguishing score simulation

Veyrat-Charvillon’s toolbox simulates a DPA attack using Hamming weight both as the power model and as the leakage function. It targets the SubBytes output (via exclusive-or between a plaintext byte and corresponding key byte, which is then used as input for SubBytes). For the leakage traces (each leakage trace consists of a single leakage value), random noise (i.e. sampled from a Gaussian distribution with zero mean and adjustable variance) is added to the Hamming weight of the SubBytes output (as produced with a known secret key). For the attack, the Hamming weight function is used again to map SubBytes outputs to predicted power values. The simulator then performs a standard DPA by utilising template matching as a distinguisher (this has been shown by Mangard et al. [12] to be equivalent to performing a correlation based DPA with a perfect model). Note that they use, as is common (see e.g. [11]), logarithms of matching outcomes to produce ‘additive’ scores. We kept the number of traces for attacks constant at 30 (which matches [16]) and changed the variance of the noise to create ‘deeper’ keys.

Their toolbox was originally written in C++ but was converted (by us) to Java to allow integration into our algorithms.

A.2  MapToWeight

As the algorithm requires integer weights, a mapping must be defined to convert the floating point numbers to integers. This is a very simple process of multiplying the raw score $d_{i,j}$, of value most $2^a$, in the distinguishing vector by
2^{p-\alpha}$ where $p$ is the bit value of precision we wish to maintain. Then performing an \texttt{abs} has the double effect of removing the negative sign, and making the most probable (the most negative numbers) the smallest, meaning they have the lightest weight which maps to our knapsack representation perfectly. Formally $w_{i,j} = \text{MapToWeight}(d_{i,j})$ where $\text{MapToWeight}(d_{i,j}) = \lfloor \text{abs}(d_{i,j} \cdot 2^{p-\alpha}) \rfloor$ for $p$ bits of precision.

### A.3 Computing environment

All code was implemented using Java 1.7, with the exception of the Glowacz et al.’s algorithm [7] which was implemented in Matlab to enable very fast convolution of the histograms. The language difference here was not an issue because key rank is so that fast we only ran accuracy comparisons and not timing comparisons. The implementation of Veyrat-Charvillon et al.’s key enumeration algorithm [16] was provided by the author, and translated into Java allowing for direct speed comparisons.

Running the single core enumeration tests, compared to Veyrat-Charvillon which are plotted in Fig. 10 (left), took place on a system running Arch Linux, with an Intel i7-4790S and 8GB of system memory.

Precision tests required larger memory capabilities and as such were carried out on a system running Ubuntu, with an Intel Xeon E5-1650 and 32GB of system memory.

Finally the multiple core tests plotted in Fig. 10 (right) were run on a cluster based environment, where each individual node provided 2 Intel E5-2670s and 64GB of memory.
B Left and Right Children Functions for Each Sort

In the following table are the left and right children function for each of the three topological sorts we considered. They all follow a similar pattern. The left child function sees if the element is the last item in the chunk and rejects if this is the case, else it returns the next item in the chunk. The right child calculates the \((x, y, z)\) coordinate for the index \(0 \leq c \leq A - 1\) given (it is trivial to convert to \(1 \leq c \leq A\) as in the rest of the paper but this greatly simplifies the code), using this it rejects if the new weight will exceed \(W\) else it returns the index of the first item in the next chunk with the weight incremented accordingly (or goes to an accept \(A\) if it is the last chunk).

<table>
<thead>
<tr>
<th>Left Child Wide</th>
<th>Right Child Wide</th>
</tr>
</thead>
<tbody>
<tr>
<td>if ((n \cdot W) - (c \mod (n \cdot W)) \leq W) then</td>
<td>(w' \leftarrow c \mod W)</td>
</tr>
<tr>
<td>return (R)</td>
<td>(\frac{(c - w') \mod (n \cdot W)}{nW})</td>
</tr>
<tr>
<td>else</td>
<td>(i \leftarrow \frac{c - w' - iW}{nW})</td>
</tr>
<tr>
<td>return (c + W)</td>
<td>if (w + w_{i,j} \geq W) then</td>
</tr>
<tr>
<td>end if</td>
<td>return (R)</td>
</tr>
<tr>
<td></td>
<td>else if (i \neq m - 1) then</td>
</tr>
<tr>
<td></td>
<td>return ((i + 1) \cdot n \cdot w' + W + w_{i,j})</td>
</tr>
<tr>
<td></td>
<td>else</td>
</tr>
<tr>
<td></td>
<td>return (A)</td>
</tr>
<tr>
<td>end if</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Left Child Long</th>
<th>Right Child Long</th>
</tr>
</thead>
<tbody>
<tr>
<td>(i \leftarrow c \mod n)</td>
<td></td>
</tr>
<tr>
<td>if (i = n - 1) then</td>
<td>(i \leftarrow c \mod n)</td>
</tr>
<tr>
<td>return (R)</td>
<td>(j \leftarrow \frac{(c-i) \mod m}{nW})</td>
</tr>
<tr>
<td>else</td>
<td>if (w + w_{i,j} \geq W) then</td>
</tr>
<tr>
<td>return (c + 1)</td>
<td>return (R)</td>
</tr>
<tr>
<td>end if</td>
<td>else if (j \neq m - 1) then</td>
</tr>
<tr>
<td></td>
<td>return (n \cdot (j + 1) + i \cdot m \cdot (w + w_{i,j}))</td>
</tr>
<tr>
<td></td>
<td>else</td>
</tr>
<tr>
<td></td>
<td>return (A)</td>
</tr>
<tr>
<td></td>
<td>end if</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Left Child Wiggly</th>
<th>Right Child Wiggly</th>
</tr>
</thead>
<tbody>
<tr>
<td>(i \leftarrow c \mod n)</td>
<td></td>
</tr>
<tr>
<td>if (i = n - 1) then</td>
<td>(i \leftarrow c \mod n)</td>
</tr>
<tr>
<td>return (R)</td>
<td>(w' \leftarrow \frac{(c-i) \mod n \cdot W}{n})</td>
</tr>
<tr>
<td>else</td>
<td>(j \leftarrow \frac{c - w' - iW}{nW})</td>
</tr>
<tr>
<td>return (c + 1)</td>
<td>if (w + w_{i,j} \geq W) then</td>
</tr>
<tr>
<td>end if</td>
<td>return (R)</td>
</tr>
<tr>
<td></td>
<td>else if (j \neq m - 1) then</td>
</tr>
<tr>
<td></td>
<td>return (n \cdot (w + w_{i,j}) + (j + 1) \cdot W \cdot n)</td>
</tr>
<tr>
<td></td>
<td>else</td>
</tr>
<tr>
<td></td>
<td>return (A)</td>
</tr>
<tr>
<td></td>
<td>end if</td>
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</table>

Table 2: test
Acknowledgements

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