

# Clone Detection in Repositories of Business Process Models

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**Abstract.** Over time, process model repositories tend to accumulate duplicate fragments (also called *clones*) as new process models are created or extended by copying and merging fragments from other models. This phenomenon calls for methods to detect clones in process models, so that these clones can be refactored as separate subprocesses in order to improve maintainability. This paper presents an indexing structure to support the fast detection of clones in large process model repositories. The proposed index is based on a novel combination of a method for process model decomposition (specifically the Refined Process Structure Tree), with established graph canonization and string matching techniques. Experiments show that the algorithm scales to repositories with hundreds of models. The experimental results also show that a significant number of non-trivial clones can be found in process model repositories taken from industrial practice.

## 1 Introduction

It is nowadays common for organizations engaged in long-term business process management programs to deal with collections of hundreds or even thousands of process models, with sizes ranging from dozens to hundreds of elements per model [15, 14]. While highly valuable, such collections of process models raise a maintenance challenge [14]. This challenge is amplified by the fact that process models are generally maintained and used by stakeholders with varying skills, responsibilities and goals, sometimes distributed across independent organizational units.

One problem that arises as repositories grow is that of managing overlap across models. In particular, process model repositories tend to accumulate duplicate fragments over time, as new process models are created by copying and merging fragments from other models. Experiments conducted during this study show that a well-known process model repository contains several hundred clones. This situation is akin to that observed in source code repositories, where significant amounts of duplicate code fragments – known as *code clones* – are accumulated over time [10].

Cloned fragments in process models raise several issues. Firstly, clones make individual process models larger than they need to be, thus affecting their comprehensibility. Secondly, clones are modified independently, sometimes by different stakeholders, leading to unwanted inconsistencies across models. Finally, process model clones hide

potential efficiency gains. Indeed, by factoring out cloned fragments into separate subprocesses, and exposing these subprocesses as shared services, companies may reap the benefits of larger resource pools.

In this setting, this paper addresses the problem of retrieving all clones in a process model repository that can be refactored into shared subprocesses. Specifically, the contribution of the paper is an index structure, namely the *RPSDAG*, that provides operations for inserting and deleting models, as well as an operation for retrieving all clones in a repository that meet the following requirements:

- All retrieved clones must be single-entry, single-exit (SESE) fragments, since subprocesses are invoked according to a call-and-return semantics.
- All retrieved clones must be *exact* clones so that every occurrence can be replaced by an invocation to a single (shared) subprocess. While identifying *approximate* clones could be useful in some scenarios, approximate clones cannot be directly refactored into shared subprocesses, and thus fall outside the scope of this study.
- Maximality – once we have identified a clone, every SESE fragment strictly contained in this clone is also a clone, but we do not wish to return all such sub-clones.
- Retrieved clones must have at least two nodes (no “trivial” clones).

Identifying clones in a process model repository boils down to identifying fragments of a process model that are isomorphic to other fragments in the same or in another model. Hence, we need a method for decomposing a process model into fragments and a method for testing isomorphism between these fragments. Accordingly, the *RPSDAG* is built on top of two pillars: (i) a method for decomposing a process model into SESE fragments, namely the *Refined Process Structure Tree (RPST)* decomposition; and (ii) a method for calculating canonical codes for labeled graphs. These canonical codes reduce the problem of testing for graph isomorphism between a pair of graphs, to a string equality check. Section 2 introduces these methods and discusses how they are used to address the problem at hand. Next, Section 3 describes the *RPSDAG*. Section 4 presents an evaluation of the *RPSDAG* using several process model repositories. Finally, Section 5 discusses related work while Section 6 draws conclusions.

## 2 Background

This section introduces the two basic ingredients of the proposed technique: the Refined Process Structure Tree (RPST) and the code-based graph indexing.

### 2.1 RPST

The RPST [18] is a parsing technique that takes as input a process model and computes a tree representing a hierarchy of SESE fragments. Each fragment corresponds to the subgraph induced by a set of edges. A SESE fragment in the tree contains all fragments at the lower level, but fragments at the same level are disjoint. As the partition is made in terms of edges, a single vertex may be shared by several fragments.

Each SESE fragment in an RPST is of one of four types [13]. A *trivial* (T) fragment consists of a single edge. A *polygon* (P) fragment is a sequence of fragments. A *bond* corresponds to a fragment where all child fragments share a common pair of vertices. Any other fragment is a *rigid*.

Figure 1a presents a sample process model for which the RPST decomposition is shown in the form of dashed boxes. A simplified view of the RPST is shown in Fig. 2.

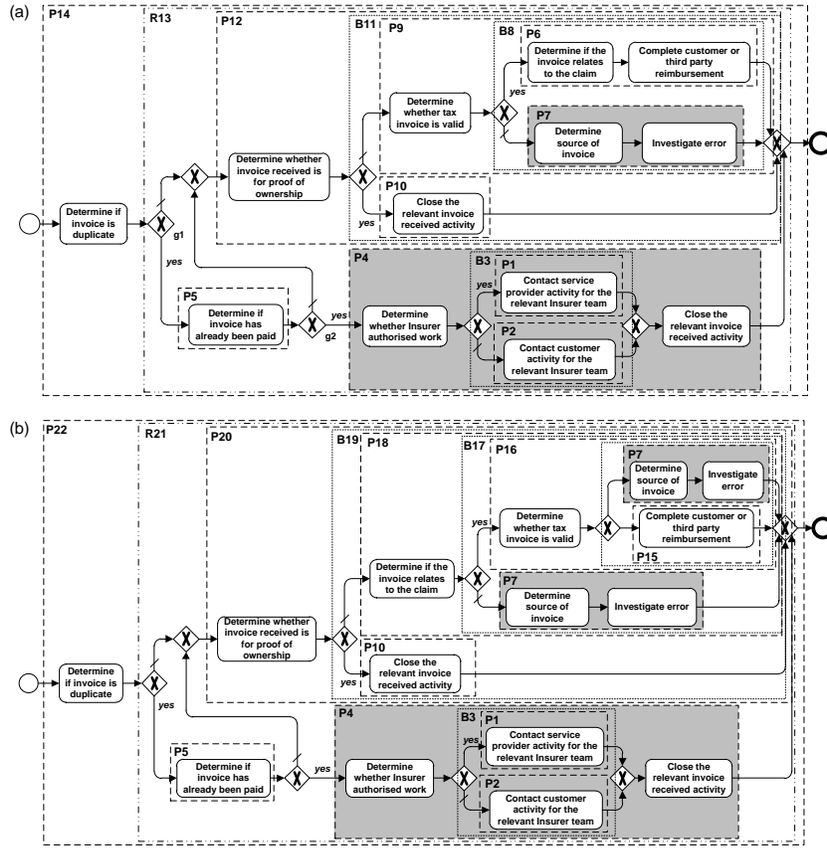


Fig. 1. Excerpt of two process models of an insurance company.

We observe that the root fragment corresponds to polygon P14, which in turn contains rigid R13 and a set of trivial fragments (simple edges) that are not shown in order to simplify the figure. R13 is composed of polygons P12, P5, P4, and so forth. Polygons P4, P5, P7 and P10 have been highlighted to ease the comparison with the similar subgraphs found in the process model shown in Figure 1b. In particular, we note that polygon P4 occurs in both process models. This is an example of a clone that we wish to detect.

Although the models used as examples are encoded in BPMN, the proposed technique can be applied to other graph-based modeling notations. To achieve this notation-independence, we adopt the following graph-based representation of process models:

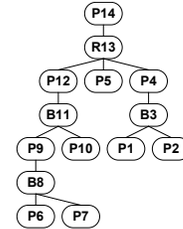


Fig. 2. RPST of process model in Fig. 1a.

**Definition 1 (Process Graph).** A *Process Graph* is a labelled connected graph  $G = (V, E, l)$ , where:

- $V$  is the set of vertices.
- $E \subseteq V \times V$  is the set of directed edges (e.g. representing control-flow relations).

- $l : V \rightarrow \Sigma^*$  is a labeling function that maps each vertex to a string over alphabet  $\Sigma$ . We distinguish the following special labels:  $l(v) = \text{“start”}$  and  $l(v) = \text{“end”}$  are reserved for start events and end events respectively;  $l(v) = \text{“xor-split”}$  is used for vertices representing xor-split (exclusive decision) gateways; similarly  $l(v) = \text{“xor-join”}$ ,  $l(v) = \text{“and-split”}$  and  $l(v) = \text{“and-join”}$  represent merge gateways, parallel split gateways and parallel join gateways respectively. For a task node  $t$ ,  $l(t)$  is the label of the task.

This definition can be extended to capture other types of BPMN elements by introducing additional types of nodes (e.g. a type of node for inclusive gateways, data objects etc.). Organizational information such as lanes and pools can be captured by attaching dedicated attributes to each node (e.g. each node could have an attribute indicating the pool and lane to which it belongs). In this paper, we do not consider sub-processes, since each sub-process can be indexed separately for the purpose of clone identification.

## 2.2 Canonical labeling of graphs

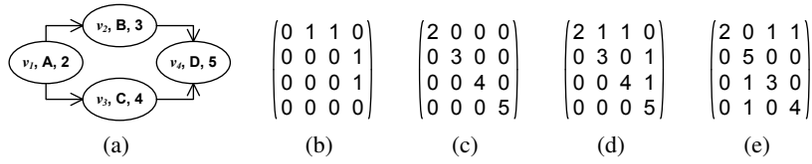
Our approach for graph indexing is an adaptation of the approach proposed in [20]. The adaptations we make relate to two specificities of process models that differentiate them from the class of graphs considered in [20]: (i) process models are directed graphs; (ii) process models can be decomposed into an RPST.

Following the method in [20], our starting point is a matrix representation of a process graph encoding the vertex adjacency and the vertex labels, as defined below.

**Definition 2 (Augmented Adjacency Matrix of a Process Graph).** Let  $G = (V, E, l)$  be a Process Graph, and  $v = (v_1, \dots, v_{|V|})$  a total order over the elements of  $V$ . The adjacency matrix of  $G$ , denoted as  $A$ , is a  $(0, 1)$ -matrix such that  $A_{i,j} = 1$  if and only if  $(v_i, v_j) \in E$ , where  $i, j \in \{1 \dots |V|\}$ . Moreover, let us consider a function  $h : \Sigma^* \rightarrow \mathbb{N} \setminus \{0, 1\}$  that maps each vertex label to a unique natural number greater than 1. The *Augmented Adjacency Matrix*  $M$  of  $G$  is defined as:  $M = \text{diag}(h(l(v_1)), \dots, h(l(v_{|V|}))) + A$

Given the augmented adjacency matrix of a process graph (or a SESE fragment therein), we can compute a string (hereby called a *code*) by concatenating the contents of each cell in the matrix from left to right and from top to bottom. For illustration, consider graph  $G$  in Figure 3(a), which is an abstract view of fragment B3 of the running example (cf. Figure 1a). For convenience, next to each vertex we show the unique vertex identifier (e.g.  $v_1$ ), the corresponding label (e.g.  $l(v_1) = \text{“A”}$ ), and the numeric value associated with the label (e.g.  $h(l(v_1)) = 2$ ). Assuming the order  $v = (v_1, v_2, v_3, v_4)$  over the set of vertices, the matrix shown in Figure 3(b) is the adjacency matrix of  $G$ . Figure 3(c) is the diagonal matrix built from  $h(l(v))$  whereas Figure 3(d) shows the augmented adjacency matrix  $M$  for graph  $G$ . It is now clear why 0 and 1 are not part of the codomain of function  $h$ , i.e. to avoid clashes with the representation of vertex adjacency. Figure 3(e) shows a possible permutation of  $M$  when considering the alternative order  $v' = (v_1, v_4, v_2, v_3)$  over the set of vertices.

Next, we transform the augmented adjacency matrix into a code by scanning from left-to-right and top-to-bottom. For instance, the matrix in Figure 3(d) can be represented as “2.1.1.0.0.3.0.1.0.0.4.1.0.0.0.5”. This code however does not uniquely represent the graph. If we chose a different ordering of the vertices, we would obtain a different code. To obtain a unique code (called a *canonical code*), we need to pick the



**Fig. 3.** (a) a sample graph, (b) its adjacency matrix, (c) its diagonal matrix with the vertex label codes, (d) its augmented adjacency matrix, and (e) a permutation of the augmented matrix

code that lexicographically “precedes” all other codes that can be constructed from an augmented adjacency matrix representation of the graph. Conceptually, this means that we have to consider all possible permutations of the set of vertices and compute a code for each permutation, as captured in the following definition.

**Definition 3 (Graph Canonical Code).** Let  $G$  be a process graph,  $M$  the augmented adjacency matrix of  $G$ . The *Graph Canonical Code* is the smallest lexicographical string representation of any possible permutation of matrix  $M$ , that is:

$$\text{code}(M) = \text{str}(P^T M P) \mid P \in \Pi \wedge \forall Q \in \Pi, P \neq Q : \text{str}(P^T M P) \prec \text{str}(Q^T M Q)$$

where:

- $\Pi$  is the set of all possible permutations of the identity matrix  $I_{|M|}$
- $\text{str}(N)$  is a function that maps a matrix  $N$  into a string representation.

Consider the matrices in Figures 3(d) and 3(e). The code of the matrix in Figure 3(e) is “2.0.1.1.0.5.0.0.1.3.0.0.1.0.4”. This code is lexicographically smaller than the code of the matrix in Figure 3(d) (“2.1.1.0.0.3.0.1.0.0.4.1.0.0.5”). If we explored all vertex permutations and constructed the corresponding matrices, we would find that “2.0.1.1.0.5.0.0.1.3.0.0.1.0.4” is the canonical code of the graph in Figure 3(a).

Enumerating all vertex permutations is unscalable (factorial on the number of vertices). Fortunately, optimizations can be applied by leveraging the characteristics of the graphs at hand. Firstly, by exploiting the nature of the fragments in an RPST, we can apply the following optimizations:

- The code of a polygon is computed in linear time by concatenating the codes of its contained vertices in the (total) order implied by the control flow.
- The code of a bond is computed in linear time by taking the entry gateway as the first vertex, the exit gateway as the last vertex and all vertices in-between are ordered lexicographically based on their labels. Since these vertices in-between do not have any control-flow dependencies among them, duplicate labels do not pose a problem.

In the case of a rigid, we start by partitioning its vertices into two subsets: vertices with distinct labels, and vertices with duplicate labels. Vertices with distinct labels are deterministically ordered in lexicographic order. Hence we do not need to explore any permutations between these vertices. Instead, we can focus on deterministically ordering vertices with duplicate labels. Duplicate labels in process models arise in two cases: (i) there are multiple tasks (or events) with the same label; (ii) there are multiple gateways of the same type (e.g. multiple “xor-splits”) that cannot be distinguished from one another since gateways generally do not have labels. To distinguish between multiple

gateways, we pre-process each gateway  $g$  by computing the tasks that immediately precede it and the tasks that immediately follow it within the same rigid fragment, and in doing so, we skip other gateways found between gateway  $g$  and each of its preceding (or succeeding) tasks. We then concatenate the labels of the preceding tasks (in lexicographic order) and the labels of the succeeding tasks (again in lexicographic order) to derive a new label  $s_g$ . The  $s_g$  labels derived in this way are used to order multiple gateways of the same type within the same rigid. Consider for example  $g_1$  and  $g_2$  in Figure 1b and let  $s_1 = \text{“Determine if invoice has already been paid”}$ ,  $s_2 = \text{“Determine whether invoice received is for proof of ownership”}$  and  $s_3 = \text{“Determine whether Insurer authorized work”}$ . We have that  $s_{g_1} = \text{“s1.s2”}$  while  $s_{g_2} = \text{“s1.s3.s2”}$ . Since  $s_3$  precedes  $s_2$ , gateway  $g_2$  will always be placed before  $g_1$  when constructing an augmented adjacency matrix for R13. In other words, we do not need to explore any permutation where  $g_1$  comes before  $g_2$ . Even if task “Determine if invoice is duplicate” precedes  $g_1$  this is not used to compute  $s_{g_1}$  because this task is outside rigid R13. To ensure unicity, vertices located outside a rigid should not be used to compute its canonical code.

A similar approach is used to order tasks with duplicate labels within a rigid. This “label propagation” approach allows us to considerably reduce the number of permutations we need to explore. Indeed, we only need to explore permutations between multiple vertices if they have identical labels and they are preceded and followed by vertices that also have the same labels. The worst-case complexity for computing the canonical code is still factorial, but on the size of the largest group of vertices inside a rigid that have identical labels and identical predecessors and successors’ labels.

### 3 Clone Detection Method

#### 3.1 Index structure and clone detection

The RPSDAG is a directed acyclic graph representing the union of the RPSTs of a collection of process models. The key feature of the RPSDAG is that each SESE fragment is represented only once. When a new process model is inserted into an RPSDAG, its RPST is computed and traversed bottom-up. For each fragment, it is checked if this fragment already exists in the RPSDAG. If it does, the fragment is not inserted, but instead, the existing fragment is reused. For instance, consider an RPSDAG constructed from the process model shown in Figure 1a. Imagine that the process model in Figure 1b is inserted into this RPSDAG, meaning that the fragments in its RPST are inserted into the RPSDAG one by one from bottom to top. When the RPST node corresponding to polygon P4 is reached, we detect that this fragment already exists. Rather than inserting the fragment, an edge is created from its parent fragment (R21) to the existing node P4.

The RPSDAG is designed to be implemented on top of standard relational databases. Accordingly, the RPSDAG consists of three tables: Codes(Code, Id, Size, Type), Roots(GraphId, RootId) and RPSDAG(ParentId, ChildId). Table Codes contains the canonical code for each RPST fragment of an indexed graph. Column “Id” assigns a unique identifier to each indexed fragment, column Code gives the canonical code of a fragment, column Size is the number of vertices in the fragment, and column Type denotes the type of fragment (“p” for polygon, “r” for rigid and “b” for bond). The “Id” is auto-generated by incrementing a counter every time that a new code is inserted into the Codes table. Strictly speaking, the “Id” is redundant given that the code uniquely

identifies each fragment. However, the “id” gives us a shorter way of uniquely identifying a fragment. For optimization reasons, when the canonical code of a new RPST node is constructed, we use the short identifiers of its child fragments as labels (in the diagonal of the augmented adjacency matrix), instead of using the child fragment’s canonical codes, which are longer. For example consider computing the canonical code for fragment P9 in Figure 1a, let  $s1 = \text{“Determine whether tax invoice is valid”}$ . Instead of extracting the fragment B8 we include the fragment identifier to the code – resulting the canonical code “s1.B8”.

Table Roots contains the Id of each indexed graph and the Id of the root fragment for that graph. Table RPSDAG is the main component of the index and is a combined representation of the RPSTs of all the graphs in the repository. Since multiple RPSTs may share fragments (these are precisely the clones we look for), the RPSDAG table represents a Directed Acyclic Graph (DAG) rather than a tree. Each tuple of this table is a pair consisting of a parent fragment id and a child fragment id. Figure 4 shows an extract of the tables representing the two graphs in Figure 1. Here, the fragment Ids have been derived from the fragment names in the Figure (e.g. the id for P4 is 4) and the codes have been hidden for the sake of readability.

Looking at the RPSDAG, we can immediately observe that the maximal clones are those child fragments that have more than one parent. Accordingly, we can retrieve all clones in an RPSDAG by means of the following SQL query.

```
SELECT RPSDAG.ChildId , Codes.Size , COUNT(RPSDAG.ParentId)
FROM RPSDAG, Codes
WHERE RPSDAG.ChildId = Codes.Id AND Codes.Size >= 2
GROUP BY RPSDAG.ChildId , Codes.Size
HAVING COUNT(RPSDAG.ParentId) >= 2;
```

This query retrieves the id, size and number of occurrences of fragments that have at least two parent fragments. Note that if a fragment appears multiple times in the indexed graphs, but always under the same parent fragment, it is not a maximal clone. For example, fragment P2 in Figure 1 always appears under B3, and B3 always appears under P4. Thus, neither P2 nor B3 are maximal clones, and the above query does not retrieve them. On the other hand, the query identifies P4 as a maximal clone since it has two parents (cf. tuples (13,4) and (21,4) in table RPSDAG in Fig. 4). Also, as per the requirements spelled out in Section 1, the query returns only fragments with at least 2 vertices. For example, even if there are two tuples with ChildId 5 and 10 in the example RPSDAG, the query will not return clones P5 and P10 as they are single nodes.

The correctness and completeness of the RPSDAG for clone detection follow from the fact that the RPST decomposition separates the process graph into SESE subgraphs that are either disjoint or have a containment relationship. Thus, if two models share a common SESE fragment S, then fragment S will be indexed exactly once in the RPSDAG and its parents will be the smallest SESE fragments that contain S.

Code	Id	Size	Type
-	1	1	p
-	2	1	p
-	3	4	b
-	4	6	p
-	5	1	p
-	6	2	p
...	...	...	...
-	13	20	r
-	14	21	p
...	...	...	...
-	22	24	p
...	...	...	...

ParentId	ChildId
3	1
3	2
4	3
8	6
8	7
...	...
13	4
13	5
...	...
14	13
16	7
16	15
17	7
21	4
21	5
...	...

GraphId	RootId
1	14
2	22

**Fig. 4.** RPST of process model in Fig. 1a.

### 3.2 Insertion and Deletion

Algorithm 1 describes the procedure for inserting a new process graph into an indexed repository. Given an input graph, the algorithm first computes its RPST with function `ComputeRPST()` which returns the RPST's root node. Next, procedure `InsertFragment` is invoked on the root node to update tables `Codes` and `RPSDAG`. This returns the id of the root fragment. Finally, a tuple is added to table `Roots` with the id of the inserted graph and that of its root node.

---

#### Algorithm 1: Insert Graph

---

```

procedure InsertGraph(Graph  $m$ )
  RPST  $root \leftarrow$  ComputeRPST( $m$ )
   $rid \leftarrow$  InsertFragment( $root$ )
   $Roots \leftarrow Roots \cup \{(NewId(), rid)\}$  // NewId() generates a fresh id

```

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#### Algorithm 2: Insert Fragment

---

```

procedure InsertFragment(RPST  $f$ ) returns RPSDAGNodeId
   $\{RPST, RPSDAGNodeId\} C \leftarrow \emptyset$ 
  foreach RPST  $child$  in GetChildren( $f$ ) do
  |  $C \leftarrow C \cup \{(child, InsertFragment(child))\}$ 
   $code \leftarrow$  ComputeCode( $f, C$ )
   $(id, type) =$  InsertNode( $code, f$ )
  foreach  $(cf, cid)$  in  $C$  do
  |  $RPSDAG \leftarrow RPSDAG \cup \{(id, cid)\}$ 
  if  $type = "p"$  then InsertSubPolygons( $id, code, f$ )
  return  $id$ 

```

---

Procedure `InsertFragment` (Algorithm 2) inserts an RPST fragment in table `RPSDAG`. This algorithm performs a depth-first search traversal of the RPST. Nodes are visited in postorder, i.e. a node is visited after all its children have been visited. When a node is visited, its canonical code is computed – function `ComputeCode` – based on the topology of the RPST fragment and the codes of its children (except of course for leaf nodes whose labels are their canonical codes). Next, procedure `InsertNode` is invoked to insert the node in table `Codes`, returning its id and type. This procedure (Algorithm 3) first checks if the node already exists in `Codes`, via function `GetIdSizeType()`. If it exists, `GetIdSizeType()` returns the id, size and type associated with that code, otherwise it returns the tuple  $(0, 0, "")$ . In this latter case, a fresh id is created for the node at hand, then its size and type are computed via functions `ComputeSize` and `ComputeType`, and a new tuple is added in table `Codes`. Function `ComputeSize` returns the sum of the sizes of all child nodes or 1 if the current node is a leaf. Once the node id and type have been retrieved, procedure `InsertFragment` adds a new tuple in table `RPSDAG` for each children of the visited node.

If the node is a polygon, procedure `InsertSubPolygons()` is invoked in the last step of `InsertFragment`. This procedure is used to identify common subpolygons and factor

**Algorithm 3:** Insert Node

---

```

procedure InsertNode(String code, RPST f) returns (RPSDAGNodeId, String)
  (id, size, type)  $\leftarrow$  GetIdSizeType(code)
  if (id, size, type) = (0, 0, "") then
    id  $\leftarrow$  NewId()
    size  $\leftarrow$  ComputeSize(code)
    type  $\leftarrow$  ComputeType(f)
    Codes  $\leftarrow$  Codes  $\cup$  {(code, id, size, type)}
  return (id, type)

```

---

**Algorithm 4:** Insert SubPolygons

---

```

procedure InsertSubPolygons(RPSDAGNodeId id, String code, RPST f)
  foreach (zcode, zid, zsize, ztype) in Codes such that ztype = "p" and zid  $\neq$  id do
    {String} LCS  $\leftarrow$  ComputeLCS(code, zcode)
    foreach lcode in LCS do
      (lid, ltype) = InsertNode(lcode, f)
      if lid  $\neq$  id then RPSDAG  $\leftarrow$  RPSDAG  $\cup$  {(id, lid)}
      if lid  $\neq$  zid then RPSDAG  $\leftarrow$  RPSDAG  $\cup$  {(zid, lid)}
      foreach sid in GetChildrenIds(lid) do
        RPSDAG  $\leftarrow$  RPSDAG  $\setminus$  {(id, sid), (zid, sid)}  $\cup$  {(lid, sid)}
        InsertSubPolygons(lid, lcode)

```

---

them out as separate nodes. Indeed, two polygons may share one or more subpolygons which should also be identified as clones. To illustrate this scenario, let us consider the two polygons  $P_1$  and  $P_2$  in Fig. 5, where  $\text{code}(P_1) = B_2.a.B_1.w.z.a.B_1.c$  and  $\text{code}(P_2) = a.B_1.c.d.a.B_1.w.z$ .<sup>1</sup> These two polygons share bond  $B_1$  as common child, while bond  $B_2$  is a child of  $P_1$  only. However, at a closer look, their canonical codes share three Longest Common Substrings (LCS), namely  $a.B_1.c$ ,  $a.B_1$  and  $w.z$ .<sup>2</sup> These common substrings represent common *subpolygons*, and thus clones that may be refactored as separate subprocesses.

Assume  $P_1$  is already stored in the RPSDAG with children  $B_1$  and  $B_2$  (first graph in Fig. 5) and we now want to store  $P_2$ . We invoke procedure  $\text{InsertFragment}(P_2)$  and add a new node in the RPSDAG, with  $B_1$  as a child (second graph in Fig. 5). Since  $P_2$  is a polygon, we also need to invoke  $\text{InsertSubPolygons}(P_2)$ . This procedure (Algorithm 4) retrieves all polygons from *Codes* that are different than  $P_2$  (in our case there is only  $P_1$ ). Then, for each such polygon, it computes all the non-trivial LCSs between its code and the code of the polygon just being inserted. This is performed by function  $\text{ComputeLCS}()$  which returns an ordered list of LCSs starting from the longest one (there are efficient algorithms to compute the LCSs of two strings. This can be done

<sup>1</sup> For the sake of readability, here we use the node/fragment label as canonical code, instead of the numeric value associated with its label.

<sup>2</sup> An LCS of strings  $S_1$  and  $S_2$  is a substring of both  $S_1$  and  $S_2$ , which is not contained in any other substring shared by  $S_1$  and  $S_2$ . LCSs of size one are not considered for obvious reasons.

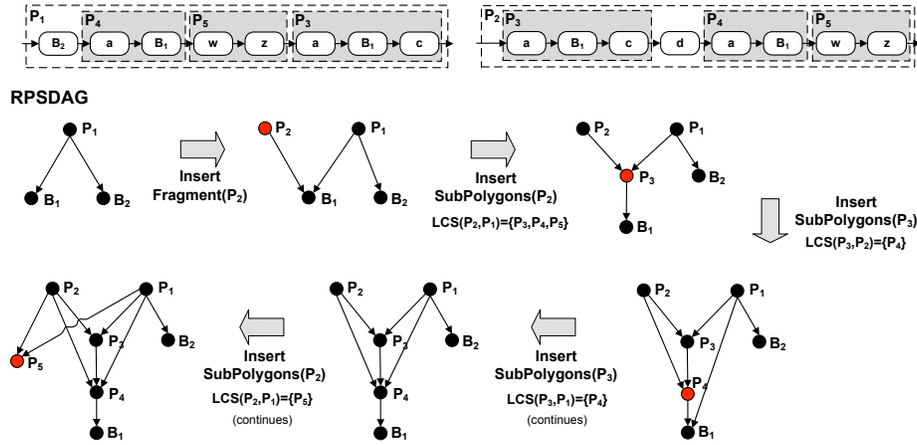


Fig. 5. Two subpolygons and the corresponding RPSDAG.

by using a *suffix tree* [17] to index the canonical codes of the existing polygons, see e.g. [6] for a survey of LCS algorithms). In the example at hand, the longest substring is  $a.B_1.c$ . This substring can be seen as the canonical code of a “shared” subpolygon between  $P_2$  and  $P_1$ . To capture the fact that this shared subpolygon is a clone, we insert a new node  $P_3$  in the RPSDAG with canonical code  $a.B_1.c$  (unless such a node already exists, in which case we reuse the existing node) and we insert an edge from  $P_2$  to  $P_3$  and from  $P_1$  to  $P_3$ . In order to avoid redundancy in the RPSDAG, the new node  $P_3$  then “adopts” all the common children that it shares with  $P_2$  and with  $P_1$ . These nodes are retrieved by function `GetChildrenIds()`, which simply returns the ids of all child nodes of  $P_3$ : these will either be children of  $P_1$  or  $P_2$  or both. In our example, there is only one common child, namely  $B_1$ , which is adopted by  $P_3$  (see third graph in Fig. 5).

It is possible that the newly inserted node also shares subpolygons with other nodes in the RPSDAG. So, before continuing to process the remaining LCSs between  $P_1$  and  $P_2$ , we invoke procedure `InsertSubPolygons` recursively over  $P_3$ . In our example, the code of  $P_3$  shares the substring  $a.B_1$  with both  $P_2$  and  $P_1$  (after excluding the code of  $P_3$  from the codes of  $P_2$  and  $P_1$  since  $P_3$  is already a child of these two nodes). Thus we create a new node  $P_4$  with code  $a.B_1$  as a child of  $P_3$  and  $P_2$ , and we make  $P_4$  adopt the common child  $B_1$  (see fourth graph in Fig. 5). We repeat the same operation between  $P_3$  and  $P_1$  but since this time  $P_4$  already exists in the RPSDAG, we simply remove the edge between  $P_1$  and  $B_1$ , and add an edge between  $P_1$  and  $P_4$  (fifth graph in Fig. 5). Then, we resume the execution of `InsertSubPolygons(P_2)` and move to the second LCS between  $P_1$  and  $P_2$ , i.e.  $a.B_1$ . Since this substring has already been inserted into the RPSDAG as node  $P_4$ , nothing is done. This process of searching for LCSs is repeated until no more non-trivial common substrings can be identified. In the example at hand, we also add subpolygon  $w.z$  between  $P_1$  and  $P_2$  (last graph in Fig. 5). At this point, we have identified and factored out all maximal subpolygons shared by  $P_1$  and  $P_2$ , and we can repeat the above process for other polygons in the RPSDAG whose canonical code shares a common substring with that of  $P_1$ .

Sometimes there may be multiple overlapping LCSs of the same size. For example, given the codes of two polygons  $a.b.c.d.e.f.a.b.c$  and  $a.b.c.k.b.c.d$ , if we extract one substring (say  $a.b.c$ ), we can no longer extract the second one ( $b.c.d$ ). In these cases we locally choose based on the number of occurrences of an LCS within the two strings in question. If they have the same number of occurrences, we randomly choose one. In the example above,  $a.b.c$  has the same size of  $b.c.d$  but it occurs 3 times, so we pick  $a.b.c$ .

We do not explicitly discuss the cases where the polygon to be inserted is a sub-polygon or superpolygon of an existing polygon. These are just special cases of shared subpolygon detection, and they are covered by procedure `InsertSubPolygons`.

Algorithm 5 shows the procedure for deleting a graph from an indexed repository. This procedure relies on another procedure for deleting a fragment, namely `DeleteFragment` (Algorithm 6). The `DeleteFragment` procedure performs a depth-first search traversal of the RPSDAG, visiting the nodes in post-order. Nodes with at most one parent are deleted, because they correspond to fragments that appear only in the deleted graph. Deleting a node entails deleting the corresponding tuple in table `Codes` and deleting all tuples in the RPSDAG table where the deleted node corresponds to the parent id. If a fragment has two or more parents, the traversal stops along that branch since this node and its descendants must remain in the RPSDAG. At the end of the `DeleteGraph` procedure, the graph itself is deleted from table `Roots` through its root id.

If the node to be removed is a polygon, it has only two parents and both parents are polygons, it is a shared subpolygon. Before deleting such a node, we need to make sure all its children are adopted by the parent polygon that will remain in the RPSDAG. For space reasons, we omit this step in the deletion algorithm.

Also, for space reasons the above algorithms do not take into account the case where a fragment is contained multiple times in the same parent fragment, like for example a polygon that contains two tasks with identical labels. To address this case, we have to introduce an attribute “Weight” in the RPSDAG table, representing the number of times a child node occurs inside a parent node. Under this representation, a node is a clone if the sum of the weights of its incoming edges in the RPSDAG is greater than one.

*Complexity.* The deletion algorithm performs a depth-first search, which is linear on the size of the graph being deleted. Similarly, the insertion algorithm traverses the inserted graph in linear time. Then, for each fragment to be inserted, it computes its code. Computing the code is linear on the size of the fragment for bonds and polygons, while for rigids it is factorial on the largest number of vertices inside the rigid that share identical labels, as discussed in Section 2.2.<sup>3</sup> Finally, if the fragment is a polygon, we compute all LCSs between this polygon and all other polygons in the RPSDAG. Using a suffix tree, this operation is linear on the sum of the lengths of all polygons’ canonical codes.

## 4 Evaluation

We evaluated the RPSDAG using four datasets: the collection of SAP R3 reference process models [9], a model repository obtained from an insurance company under condition of anonymity and two collections from the IBM BIT process library [5], namely collections A and B3. In the BIT process library there are 5 collections (A, B1, B2, B3

<sup>3</sup> A tighter complexity bound for this problem is given in [1].

---

**Algorithm 5:** Delete Graph

---

```

procedure DeleteGraph(GraphId mid)
  RPSDAGNodeId rid  $\leftarrow$  GetRoot(mid)
  DeleteFragment(rid)
  Roots  $\leftarrow$  Roots  $\setminus$   $\{(mid, rid)\}$ 

```

---



---

**Algorithm 6:** Delete Fragment

---

```

procedure DeleteFragment(RPSDAGNodeId fid)
  if  $|\{(pid, cid) \in RPSDAG : cid = fid\}| \leq 1$  then
    foreach (pid, cid) in RPSDAG where pid = fid do
      DeleteFragment(cid)
      RPSDAG  $\leftarrow$  RPSDAG  $\setminus$   $\{(pid, cid)\}$ 
    Codes  $\leftarrow$   $\{(code, id, size, type) \in Codes : id \neq fid\}$ 

```

---

and C). We excluded collections B1 and B2 because they are earlier versions of B3, and collection C because it is a mix of models from different sources and as such it does not contain any clones. The SAP repository contains 595 models with sizes ranging from 5 to 119 nodes (average 22.28, median 17). The insurance repository contains 363 models ranging from 4 to 461 nodes (average 27.12, median 19). The BIT collection A contains 269 models ranging from 5 to 47 nodes (average 17.01, median 16) while collection B3 contains 247 models with 5 to 42 nodes (average 12.94, median 11).

*Performance evaluation.* We first evaluated the insertion times. Obviously inserting a new model into a nearly-empty RPSDAG is less costly than doing so in an already populated one. To factor out this effect, we randomly split each dataset as follows: One third of the models were used to construct an initial RPSDAG and the other two-thirds were used to measure insertion times. In the SAP repository, 200 models were used to construct an initial RPSDAG. Constructing the initial RPSDAG took 26.1s. In the insurance company repository, the initial RPSDAG contained 121 models and its construction took 7.4s. For the BIT collections A and B3, 90 and 82 models respectively were used for constructing the initial RPSDAG. Constructing the initial RPSDAGs took 4.2s for collection A and 3.5s for collection B3. All tests were conducted on a PC with a dual core Intel processor, 1.8 GHz, 4 GB memory, running Microsoft Windows 7 and Oracle Java Virtual Machine v1.6. The RPSDAG was implemented as a Java console application on top of MySQL 5.1. Each test was run 10 times and the execution times obtained across the ten runs were averaged.

Table 1 summarizes the insertion time per model for each collection (min, max, avg., std. dev., and 90<sup>th</sup> percentile). All values are in milliseconds. These statistics are given for two cases: without subpolygon clone detection and with subpolygon clone detection. We observe that the average insertion time per model is about 3-5 times larger when subpolygon clone detection is performed. This overhead comes from the step where we compare inserted polygon with each already-indexed polygon and compute the longest-common substring of their canonical codes. Still, the average execution

times remain in the order of tens of milliseconds across all model collections even with subpolygon detection. The highest average insertion time (125ms) is observed for the Insurance collection. This collection contains some models with large rigid components in their RPST. In particular, one model contained a rigid component in which 2 task labels appeared 9 times each – i.e. 9 tasks had one of these labels and 9 tasks had the other – and these tasks were all preceded by the same gateway. This label repetition affected the computation of the canonical code (4.4 seconds for this particular rigid). Putting aside this extreme case, all insertion times were under one second and in 90% of the cases ( $l_{90}$ ), the insertion times in this collection were under 50ms without subpolygon detection and 222ms with subpolygon detection. Thus we can conclude that the proposed technique scales up to real-sized model collections.

	min	max	avg	std	$l_{90}$	
SAP	4	85	20	14	40	No subpolygon
Insurance	5	1722	32	113	49	
BIT A	3	58	12	9	23	
BIT B3	5	467	14	9	28	
SAP	4	482	97	81	202	Subpolygon
Insurance	26	4402	126	291	222	
BIT A	18	128	41	20	59	
BIT B3	5	150	33	24	69	

**Table 1.** Model insertion times (in ms).

As explained in Section 3, once the models are inserted, we can find all clones with a SQL query. The query to find all clones with at least 2 nodes and 2 parents from the SAP reference models takes 75ms on average (90ms for the insurance models, 118 and 116ms for BIT collections A and B3).

*Refactoring gain.* One of the main applications of clone detection is to refactor the identified clones as shared subprocesses in order to reduce the size of the model collection and increase its navigability. In order to assess the benefit of refactoring clones into subprocesses, we define the following measure.

**Definition 4.** *The refactoring gain of a clone is the reduction in number of nodes obtained by encapsulating that clone into a separate subprocess, and replacing every occurrence of the clone with a task that invokes this subprocess. Specifically, let  $S$  be the size of a clone, and  $N$  the number of occurrences of this clone. Since all occurrences of a clone are replaced by a single occurrence plus  $N$  subprocess invocations, the refactoring gain is:  $S \cdot N - S - N$ .*

*Given a collection of models, the total refactoring gain is the sum of the refactoring gains of the clones of non-trivial clones (size  $\geq 2$ ) in the collection.*

Table 2 summarizes the total refactoring gain for each model collection. The first two columns correspond to the total number of clones detected and the total refactoring gain without subpolygon refactoring. The third and fourth column show number of clones and refactoring gain with subpolygon refactoring. The table shows that a significant number of clones can be found in all model collections, and that the size of these model collections could be reduced by 4-14% if clones were factored out into shared subprocesses. The table also shows that subpolygon clone detection adds significant value to the clone detection method. For instance, in the case of the insurance models, we obtain about 3 times more clones and 3 times more refactoring gain when subpolygon clone detection is performed.

	No subpolygon		With subpolygon	
	Nr. clones	refactoring gain	Nr. clones	refactoring gain
SAP	204	1359 (10.3%)	479	1834 (13.8%)
Insurance	107	394 (4%)	279	883 (9%)
BIT A	57	195 (4.3%)	174	384 (8.4%)
BIT B3	19	208 (6.5%)	49	259 (6.6%)

**Table 2.** Total refactoring gain without and with subpolygon refactoring.

Table 3 shows more detailed statistics of the clones found with subpolygon detection. The first three columns give statistics about the sizes of the clones found, the next three columns refer to the frequency (number of occurrences) of the clones, and the last three correspond to refactoring gain. We observe that while the average clone size is relatively small (3-5 nodes), there are some large clones with sizes of 30+ nodes.

	Size			# occurrences			Refactoring gain		
	avg	max	std. dev.	avg	max	std. dev.	avg	max	std. dev.
SAP	4.79	41	4.05	2.33	8	0.77	3.83	44	5.68
Insurance	3.58	32	3.18	2.76	41	3.18	3.16	79	7.67
BIT A	3.02	16	2.08	2.75	9	1.29	2.21	15	3.12
BIT B3	3.18	9	1.58	3.82	20	3.70	5.29	37	8.87

**Table 3.** Statistics of detected clones (with subpolygon detection).

## 5 Related Work

Clone detection in software repositories has been an active field for several years. According to [3], approaches can be classified into: textual comparison, token comparison, metric comparison, abstract syntax tree (AST) comparison, and program dependence graphs (PDG) comparison. The latter two categories are close to our problem, as they use a graph-based representation. In [2], the authors describe a method for clone detection based on ASTs. The method applies a hash function to subtrees of the AST in order to distribute subtrees across buckets. Subtrees in the same bucket are compared by testing for tree isomorphism. This work differs from ours in that RPSTs are not perfect trees. Instead, RPSTs contain rigid components that are irreducible and need to be treated as subgraphs—thus tree isomorphism is not directly applicable. [11] describes a technique for code clone detection using PDGs. A subgraph isomorphism algorithm is used for clone detection. In contrast, we employ canonical codes instead of pairwise subgraph isomorphism detection. Another difference is that we take advantage of the RPST in order to decompose the process graph into SESE fragments.

Work on clone detection has also been undertaken in the field of model-driven engineering. [4] describes a method for detecting clones in large repositories of Simulink/-TargetLink models from the automotive industry. Models are partitioned into connected components and compared pairwise using a heuristic subgraph matching algorithm. Again, the main difference with our work is that we use canonical codes instead of subgraph isomorphism detection. In [12], the authors describe two methods for exact and approximate matching of clones for Simulink models. In the first method, they apply an incremental, heuristic subgraph matching algorithm. In the second approach, graphs are represented by a set of vectors built from graph features: e.g. path lengths, vertex in/out degrees, etc. An empirical study shows that this feature-based approximate matching approach improves pre-processing and running times, while keeping a high precision. However, this data structure does not support incremental insertions/deletions.

Our work is also related to graph database indexing and is inspired by [20]. Other related indexing techniques for graph databases include GraphGrep [16] – an index designed to retrieve paths in a graph that match a regular expression. This problem however is different from that of clone detection since clones are not paths (except for polygons). Another related index is the closure-tree [7]. Given a graph  $G$ , the closure-tree can be used to retrieve all indexed graphs in which  $G$  occurs as a subgraph. We could use the closure tree to index a collection of process graphs so that when a new graph is inserted we can check if any of its SESE regions appears in an already indexed graph. However, the closure tree does not directly retrieve the exact set of graphs where a given subgraph occurs. Instead, it retrieves a “candidate set” of graphs. An exact subgraph isomorphism test is then performed against each graph in the candidate set. In contrast, by storing the canonical code of each SESE region, the RPSDAG obviates the need for this subgraph isomorphism testing.

In [8], we described an index to retrieve process models in a repository that exactly or approximately match a given model fragment. In this approach, paths in the process models are used as index features. Given a collection of models, a B+ tree is used to reduce the search space by discarding those models that do not contain any path of the query model. The remaining models are tested for subgraph isomorphism.

In [19], eleven process model refactoring techniques are identified and evaluated. Extracting process fragments as subprocesses is one of the techniques identified. Our work addresses the problem of identifying opportunities for such “fragment extraction”.

## 6 Conclusion

We presented a technique to index process models in order to identify duplicate SESE fragments (clones) that can be refactored into shared subprocesses. The proposed index, namely the RPSDAG, combines a method for decomposing process models into SESE fragments (the RPST decomposition) with a method for generating a unique string from a labeled graph (canonical codes). These canonical codes are used to determine whether a SESE fragment in a model appears elsewhere in the same or in another model.

The RPSDAG has been implemented and tested using process model repositories from industrial practice. In addition to demonstrating the scalability of the RPSDAG, the experimental results show that a significant number of non-trivial clones can be found in industrial process model repositories. In one repository, ca. 480 non-trivial clones were found. By refactoring these clones, the overall size of the repository is reduced by close to 14%, which arguably would enhance the repository’s maintainability.

A standalone release of the RPSDAG implementation, together with sample models, is available at: <http://apromore.org/tools>. The current RPSDAG implementation can be further optimized in three ways: (i) by using suffix trees for identifying the longest common substrings between the code of an inserted polygon and those of already-indexed polygons; (ii) by storing the canonical codes of each fragment in a hash index in order to speed up retrieval; (iii) by using the Nauty library for computing canonical codes (<http://cs.anu.edu.au/~bdm/nauty>). Nauty implements several optimizations that could complement those described in Section 2.2.

Another avenue for future work is to extend the proposed technique in order to identify *approximate* clones. This has applications in the context of process standardization,

when analysts seek to identify similar but non-identical fragments and to replace them with standardized fragments in order to increase the homogeneity of work practices.

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