Computing *k*-Bisimulations for Large Graphs: A Comparison and Efficiency Analysis

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Abstract. Summarizing graphs w.r.t. structural features is important to reduce the graph's size and make tasks like indexing, querving, and visualization feasible. Our generic parallel BRS algorithm efficiently summarizes large graphs w.r.t. a custom equivalence relation \sim defined on the graph's vertices V. Moreover, the definition of \sim can be chained $k \geq 1$ times, so the defined equivalence relation becomes a k-bisimulation. We evaluate the runtime and memory performance of the BRS algorithm for k-bisimulation with k = 1, ..., 10 against two algorithms found in the literature (a sequential algorithm due to Kaushik et al. and a parallel algorithm of Schätzle et al.), which we implemented in the same software stack as BRS. We use five real-world and synthetic graph datasets containing 100 million to two billion edges. Our results show that the generic BRS algorithm outperforms the respective native bisimulation algorithms on all datasets for all k > 5 and for smaller k in some cases. The BRS implementations of the two bisimulation algorithms run almost as fast as each other. Thus, the BRS algorithm is an effective parallelization of the sequential Kaushik et al. bisimulation algorithm.

Keywords: structural graph summarization \cdot bisimulation \cdot large labeled graphs.

1 Introduction

Storing, indexing, querying, and visualizing large graphs is difficult [7]. One way to mitigate this challenge is graph summarization [9]. Graphs can be summarized w.r.t. so-called graph summary models [6] that define structural features (e.g., incoming/outgoing paths), statistical measures (e.g., occurrences of specific vertices), or frequent patterns found in the graph [9]. This gives a summary graph, which is usually smaller than the original but contains an approximation of or exactly the same information as the original graph w.r.t. the selected features of the summary model. Tasks that were to be performed on the original graph can be performed on the summary but much faster. Use cases are optimizing database queries [22], data visualization [12], and OWL reasoning [28].

Blume, Richerby, and Scherp developed a generic structural summarization approach, here referred to as BRS [5, 6]. The BRS algorithm summarizes an

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input graph w.r.t. an arbitrary user-defined equivalence relation specified in its formal language FLUID [6]. The FLUID language supports all features of structural graph summarization found in the literature [6]. There are two groups of these features. The first comprises a vertex's *local* information, e.g., its label set, its direct neighbors, and the labels of its incoming or outgoing edges. The second group considers a vertex's *global* information at distance k > 1. This includes, e.g., local information about reachable vertices up to distance k or information about incoming or outgoing paths of length up to k. We use stratified k-bisimulations (formally described in Section 3.2) to summarize a graph w.r.t. global information and group together vertices that have equivalent structural neighborhoods up to distance k. Several existing approaches use k-bisimulations to incorporate global information into structural graph summarization [6, 9]. The BRS algorithm generalizes these approaches and can chain any definable equivalence relation k times, such that the resulting equivalence classes can be efficiently computed by global information up to distance k [4, 24].

However, it is not known if a general approach like the BRS algorithm sacrifices performance. We choose two representative algorithms as examples to demonstrate the capabilities of our generic BRS algorithm. First, we have reimplemented the efficient, parallel single-purpose k-bisimulation algorithm of Schätzle, Neu, Lausen, and Przjaciel-Zablocki [25] and investigate whether it outperforms our generic BRS algorithm. Second, we investigate the sequential algorithm for bisimulation by Kaushik, Shenoy, Bohannon, and Gudes [16]. Being sequential, it is naturally disadvantaged against parallel algorithms such as ours. However, we show in this work that the bisimulation of Kaushik et al. [16] can be declaratively specified and executed in the generic BRS algorithm. This effectively parallelizes the algorithm "for free". We evaluate the performance of the BRS-based parallelized computation of the Kaushik et al. graph summary model and compare it with their sequential native algorithm. We also compare both Kaushik et al. variants, native and BRS-based, with the parallel native algorithm of Schätzle et al. [25] and a BRS implementation of Schätzle et al. (see also [4]). Thus, we have four k-bisimulation algorithms. For a fair comparison, we reimplemented the existing native algorithms in the same graph processing framework as the BRS algorithm. We execute the four algorithms on five graph datasets – two synthetic and three real-world – of different sizes, ranging from 100 million edges to billions of edges. We evaluate the algorithms' performance for computing k-bisimulation for k = 1, ..., 10. We measure running time per iteration and the maximum memory consumption.

The questions we address are: Do the native bisimulation algorithms have an advantage over a generic solution? How well do the native and generic algorithms scale to large real-world and synthetic graphs? Is it possible to effectively scale a sequential algorithm by turning it into a parallel variant by using a general formal language and algorithm for graph summaries?

We discuss related work next. Section 3 defines preliminaries, while the algorithms are introduced in Section 4. Section 5 outlines the experimental apparatus. Section 6 describes the results, and these are discussed in Section 7.

2 Related Work

Summary graphs can be constructed in several ways. Čebirić et al. [9] classify existing techniques into structural, pattern-mining, statistical, and hybrid approaches. In this paper, we consider only structural approaches based on quotients. Other structural summarization techniques, not based on quotients, are extensively discussed by Čebirić et al. [9]. Structural approaches summarize a graph G w.r.t. an equivalence relation $\sim \subseteq V \times V$ defined on the vertices V of G [?,9]. The resulting summary graph SG consists of vertices VS, each of which corresponds to a equivalence class of the equivalence relation \sim .

One can observe that k-bisimulation is a popular feature for structural graph summarization [6]. Bisimulation comes in three forms: backward k-bisimulation classifies vertices based on incoming paths of length up to k, forward bisimulation considers outgoing paths, and backward-forward bisimulation considers both. Bisimulation may be based on edge labels, vertex labels, or both, but this makes no significant difference to the algorithms. A notion of k-bisimulation w.r.t. graph indices is introduced by seminal works such as the k-RO index [21] and the Tindex summaries [20]. Milo and Suciu's T-index [20], the A(k)-Index by Kaushik et al. [16], and others are examples that summarize graphs using backward kbisimulation. We chose as representative the sequential algorithm by Kaushik et al. [16], which uses vertex labels, as described in Section 4.2. Conversely, the k-RO index, the Extended Property Paths of Consens et al. [11], the SemSets model of Ciglan et al. [10], Buneman et al.'s RDF graph alignment [8], and the work of Schätzle et al. [25] are based on forward k-bisimulation. We note that Schätzle et al. use edge labels, as described in Section 4.1. Tran et al. compute a structural index for graphs based on backward-forward k-bisimulation [27]. Moreover, they parameterize their notion of bisimulation to a forward-set L_1 and a backward-set L_2 , so that only labels $l \in L_1$ are considered for forwardbisimulation and labels $l \in L_2$ for backward-bisimulation.

Luo et al. examine structural graph summarization by forward k-bisimulation in a distributed, external-memory model [18]. They empirically observe that, for values of k > 5, the summary graph's partition blocks change little or not at all. Therefore they state, that for summarizing a graph with respect to kbisimulation, it is sufficient to summarize up to a value of k = 5 [17]. Finally, Martens et al. [19] introduce a parallel bisimulation algorithm for massively parallel devices such as GPU clusters. Their approach is tested on a single GPU with 24 GB RAM, which limits its use on large datasets. Nonetheless, their proposed blocking mechanism could be combined with our vertex-centric approach to further improve performance.

Each of the works proposes a single algorithm for computing a single graph summary based on a bisimulation. Some of the algorithms have bisimulation parameters such as the height and label parameterization in Tran et al. [27]. We suggest a generic algorithm for computing k-bisimulation and show its advantages. Also, our approach allows the bisimulation model to be specified in a declarative way and parallelizes otherwise sequential computations like in Kaushik et al. [16] into a parallel computation.



Fig. 1: An example graph G displaying two universities and three employees. Vertices are denoted by ellipses and edges by arrows. Vertex labels are marked with rectangles and edge labels are written on the edge.

3 Preliminaries

3.1 Data Structures

The algorithms operate on multi-relational, labeled graphs $G = (V, E, l_V, l_E)$, where $V = \{v_1, v_2, \ldots, v_n\}$ is a set of vertices and $E \subseteq V \times V$ is a set of directed edges between the vertices in V. Each vertex $v \in V$ has a finite set of labels $l_V(v)$ from a set Σ_V and each edge has a finite set of labels $l_E(u, v)$ from a set Σ_E .

Figure 1 shows an example graph representing two universities and three employees. Vertices are represented by ellipses and edges are labeled arrows. Vertex labels are shown in rectangles. For example, the edges (pr837, xuni) and (st143, xuni) labeled with *worksAt* together with edges (pr837, "Bob"), (st143, "Alice") and (xuni, "X University") labeled with *name* and vertex labels *Professor* (pr837), *Student* (st143) and *Organization* (xuni) state that professor Bob and student Alice both work at the organization X University.

In a graph $G = (V, E, l_V, l_E)$ the *in-neighbors* of a vertex $v \in V$ are the set of vertices $N^-(v) = \{u \mid (u, v) \in E\}$ from which v receives an edge. Similarly, v's *out-neighbors* are the set $N^+(v) = \{w \mid (v, w) \in E\}$ to which it sends edges. For a set $S \subseteq V$, let $N^+(S) = \bigcup_{v \in S} N^+(v)$ be the set of out-neighbors of S.

3.2 Bisimulation

A bisimulation is an equivalence relation on the vertices of a directed graph [16, 27]. Informally, a bisimulation groups vertices with equivalent structural neighborhoods, i. e., the neighborhoods cannot be distinguished based on the vertices' sets of labels and/or edges' labels. Forward bisimulation (fw) considers outgoing edges; backward bisimulation (bw) uses incoming edges. Vertices u and v are forward-bisimilar if, every out-neighbor u' of u has a corresponding out-neighbor v' of v, and vice versa; furthermore, the two neighbors u' and v' must be bisimilar [16, 27]. Backward-bisimulation is defined similarly but using in-neighbors.

k	Partition blocks
0	${pr837}, {cs902}, {st143}, {uoy, xuni}, literals$
1	${pr837}, {cs902}, {st143}, {uoy}, {xuni}, {"Alice"}, {"Charlie"},$
	{"Bob"}, {"X University", "Univ. of Y"}
2	${pr837}, {cs902}, {st143}, {uoy}, {xuni}, {"Alice"}, {"Charlie"},$
	{"Bob"}, {"X University"}, {"Univ. of Y"}

Table 1: Vertex-labeled backward 2-bisimulation partition of the example graph according to Definition 2.

This definition corresponds to a *complete* bisimulation. For the neighbors u' and v' to be bisimilar, their in-/out-neighbors must be bisimilar as well. A k-bi-simulation is a bisimulation on G that considers features a distance at most k from a vertex when deciding whether it is equivalent to another.

The algorithms of Schätzle et al. [25] and Kaushik et al. [16] compute versions of forward and backward k-bisimulation on labeled graphs $G = (V, E, \Sigma_V, \Sigma_E)$.

Definition 1. The edge-labeled forward k-bisimulation $\approx_{\text{fw}}^k \subseteq V \times V$ with $k \in \mathbb{N}$ of Schätzle et al. [25] is defined as follows:

 $\begin{array}{l} - \ u \approx^0_{\mathrm{fw}} v \ \text{for all } u, v \in V, \\ - \ u \approx^{k+1}_{\mathrm{fw}} v \ \text{iff } u \approx^k_{\mathrm{fw}} v \ \text{and, for every edge } (u, u'), \ \text{there is an edge } (v, v') \ \text{with} \\ l_E(u, u') = l_E(v, v') \ \text{and} \ u' \approx^k_{\mathrm{fw}} v', \ \text{and vice-versa.} \end{array}$

For the graph in Figure 1, \approx_{fw}^{0} has the single block V (as for all graphs, all vertices are initially equivalent) and \approx_{fw}^{1} has three blocks, i.e., sets of equivalent vertices: {pr837, cs902, st143}, {uoy, xuni}, and the literals. The vertices xuni and uoy are not 1-bisimilar to st143, cs902 and pr837, as they have no outgoing edge labeled *worksAt*. For $k \geq 2$, k-bisimulation in this case makes no more distinctions than 1-bisimulation. Note that Schätzle et al. compute full bisimulations. We have modified their algorithm to stop after k iterations.

Definition 2. The vertex-labeled backward k-bisimulation $\approx_{\text{bw}}^{k} \subseteq V \times V$ with $k \in \mathbb{N}$ of Kaushik et al. [16] is defined as follows:

 $\begin{array}{l} - \ u \approx_{\mathrm{bw}}^{0} v \ i\!f\!f \ l_{V}(u) = l_{V}(v), \\ - \ u \approx_{\mathrm{bw}}^{k+1} v \ i\!f\!f \ v \approx_{\mathrm{bw}}^{k} u \ and, \ f\!or \ every \ (u', u) \in E, \ there \ is \ (v', v) \in E \ with \\ u' \approx_{\mathrm{bw}}^{k} v', \ and \ vice \ versa. \end{array}$

Table 1 shows the vertex-labeled backward 2-bisimulation partitions of the graph in Figure 1. 0-bisimulation partitions by label. Then, xuni, uoy and the literals are split by their parents' labels. No vertex is 2-bisimilar to any other: every block is a singleton.

3.3 Graph Summaries for Bisimulation

The BRS algorithm summarizes graphs with respect to a graph summary model (GSM), a mapping from graphs G = (V, E) to equivalence relations $\sim \subseteq V \times V$.

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The equivalence classes of \sim partition G. A simple GSM is label equality, i.e., two vertices are equivalent iff they have the same label. Depending on the application, one might want to summarize a graph w.r.t. different GSMs. Therefore, the algorithm works with GSMs defined in our formal language FLUID [6]. To flexibly and quickly define GSMs, the language provides simple and complex schema elements, along with six parameterizations, of which we use two (for details we refer to [6]).

A complex schema element $CSE := (\sim_s, \sim_p, \sim_o)$ combines three equivalence relations [6]. Vertices v and v' are equivalent, iff $v \sim_s v'$; and for all $w \in N^+(v)$ there is a $w' \in N^+(v')$ with $l_E(v, w) \sim_p l_E(v', w')$ and $w \sim_o w'$, and vice versa.

The chaining parameterization enables computing k-bisimulations by increasing the neighborhood considered for determining vertex equivalence [6]. It is defined by nesting CSEs. Given a complex schema element $CSE := (\sim_s, \sim_p, \sim_o)$ and $k \in \mathbb{N}_{>0}$, the chaining parameterization CSE^k defines the equivalence relation that corresponds to recursively applying CSE to a distance of k hops. $CSE^1 := (\sim_s, \sim_p, \sim_o)$ and, inductively for k > 1, $CSE^k := (\sim_s, \sim_p, CSE^{k-1})$. This results in a summary graph that has one vertex for each equivalence class in each equivalence relation defined within the CSE. Summary vertices v and w are connected via a labeled edge, if all vertices in the input graph represented by v have an edge with this label to a vertex in w. For full details, see [6].

To model backward k-bisimulations, we need to work with incoming edges, whereas the schema elements consider only outgoing edges. In FLUID, this is done with the direction parameterization [6] but, here, we simplify notation. We write SE^{-1} for the schema element defined analogously to SE but using the relation $E^{-1} = \{(y, x) \mid (x, y) \in E\}$ in place of the graph's edge relation E, and the edge labeling $\ell_E^{-1}(y, x) = \ell_E(x, y)$.

Following Definitions 1 and 2 of Schätzle et al. and Kaushik et al., we define CSE_{Sch} and CSE_{Kau} as follows. Here, $id = \{(v, v) \mid v \in V\}$ and $T = V \times V$, and vertices are equivalent in OC_{type} iff they have the same labels [6].

$$CSE_{\rm Sch} := (T, id, T)^k \tag{1}$$

$$CSE_{\text{Kau}} := \left((OC_{\text{type}}, T, OC_{\text{type}})^{-1} \right)^{\kappa}.$$
⁽²⁾

4 Algorithms

We introduce the single-purpose algorithms of Schätzle et al. [25] in Section 4.1 and Kaushik et al. [16] in Section 4.2. Finally, we introduce the generic BRS algorithm in Section 4.3. The first two algorithms compute summaries in a fundamentally different way to the BRS algorithm. At the beginning of the execution, BRS considers every vertex to be in its own equivalence class. During execution, vertices with the same vertex summary are merged. Therefore, BRS can be seen as a *bottom-up* approach. In contrast, Schätzle et al. consider all vertices to be equivalent at the beginning, and Kaushik et al. initially consider all vertices with the same label to be equivalent. The equivalence relation is then successively refined. These two algorithms can be seen as *top-down* approaches. Here, it is convenient to consider set partitions. A partition of a set V is a set $\{B_1, \ldots, B_\ell\}$ such that: (i) $\emptyset \subsetneq B_i \subseteq V$ for each i, (ii) $\bigcup_i B_i = V$, and (iii) $B_i \cap B_j = \emptyset$ for each $i \neq j$. The sets B_i are known as *blocks*. The equivalence classes of an equivalence relation over V partition the graph's vertices. A key concept is *partition refinement*. A partition $P_i = \{B_{i1}, B_{i2}, \ldots\}$ refines $P_j = \{B_{j1}, B_{j2}, \ldots\}$ iff every block B_{ik} of P_i is contained in a block $B_{j\ell}$ of P_j .

4.1 Native Schätzle et al. Algorithm

This algorithm [25] is a distributed MapReduce approach for reducing labeled transition systems. Two fundamental concepts are the *signature* and *ID* of a vertex v with respect to the current iteration's partition P_i .

The signature of a vertex v w.r.t. a partition $P_i = \{B_{i1}, B_{i2}, \ldots\}$ of V is given by $\operatorname{sig}_{P_i}(v) = \{(\ell, B_{ij}) \mid (v, w) \in E \text{ with } l_E(v, w) = \ell \text{ and } w \in B_{ij}\}$. That is, v's signature w.r.t. the current iteration's partition P_i is the set of outgoing edge labels to blocks of P_i . By Definition 1, $u \approx_{\mathrm{fw}}^{k+1} v$, iff $\operatorname{sig}_{P_k}(u) = \operatorname{sig}_{P_k}(v)$. Therefore signatures identify the block of a vertex and represent the current bisimulation partition, and the signature of v w.r.t. P_i can be represented as $\operatorname{sig}_{P_{i+1}}(v) = \{(l_E(v,w),\operatorname{sig}_{P_i}(w)) \mid (v,w) \in E\}$. The nested structure of vertex signatures means they can become very large. Thus, we compute a recursively defined hash value proposed by Hellings et al. [13], which is also used by Schätzle et al. [25]. We use this hash function to assign $\operatorname{sig}_{P_i}(v)$ an integer value which we denote by $\operatorname{ID}_{P_i}(v)$. Now the signature of a vertex v w.r.t. the current partition P_i can be represented as $\operatorname{sig}_{P_{i+1}}(v) = \{(l_E(v,w), \operatorname{ID}_{P_i}(w)) \mid (v,w) \in E\}$.

With $\operatorname{sig}_{P_i}(v)$ and $\operatorname{ID}_{P_i}(v)$, the procedure for computing an edge-labeled forward k-bisimulation partition is outlined in Algorithm 1. The initial partition is just V (line 2), as every vertex is 0-bisimilar to every other vertex. Next, the algorithm performs k iterations (lines 3–8). In the *i*th iteration, the information needed to construct a vertex's signature $\operatorname{sig}_{P_i}(v)$ is sent to every vertex v (line 5). This information is the edge label $l_E(v,w) \in \Sigma_E$ and the block identifier $\operatorname{ID}_{P_{i-1}}(w)$ for every $w \in N^+(v)$. The signature $\operatorname{sig}_{P_i}(v)$ is then constructed using the received information, and the identifiers $\operatorname{ID}_{P_i}(v)$ are updated for all v (line 6). At the end of each iteration, the algorithm checks if any vertex ID was updated, by comparing the number of distinct values in ID_{P_i} and $\operatorname{ID}_{P_{i-1}}$ (line 7). If no vertex ID was updated, we have reached full bisimulation [3, 25] and hence can stop execution early. At the end, the resulting k-bisimulation partition P_k is constructed by putting vertices v in one block if they share the same identifier value $\operatorname{ID}_{P_i}(v)$ (line 9).

4.2 Native Kaushik et al. Algorithm

This algorithm [16] sequentially computes vertex-labeled backward k-bisimulations. The following definitions are from [23], modified for backward bisimulation.

A subset $B \subseteq V$ is *stable* with respect to another subset $S \subseteq V$ if either $B \subseteq N^+(S)$ or $B \cap N^+(S) = \emptyset$. That is, vertices in a stable set B are indistinguishable

Algorithm 1: Bisimulation Algorithm by Schätzle et al. [25]

1 f	unction BISIMSCHÄTZLE $(G = (V, E, l_V, l_E), k \in \mathbb{N})$
	/* Initially, all $v \in V$ in same block with $\mathrm{ID}_{P_0}(v) = 0$ */
2	$P_0 \leftarrow \{V\};$
3	for $i \leftarrow 1$ to k do
	/* Map Job */
4	for $(v, w) \in E$ do
5	Send $(l_E(v, w), \mathrm{ID}_{P_{i-1}}(w))$
	/* Reduce Job */
6	Construct $\operatorname{sig}_{P_i}(v)$ and update $\operatorname{ID}_{P_i}(v)$;
	/* Check if full bisimulation is reached */
7	if $ ID_{P_i} = ID_{P_{i-1}} $ then
8	break;
9	Construct P_k from ID_{P_i} ;
10	return P_k ;

by their relation to S: either all vertices in B get at least one edge from S, or none do. If B is not stable w.r.t. S, we call S a *splitter* of B.

Building on this, partition P_i of V is stable with respect to a subset $S \subseteq V$ if every block $B_{ij} \in P_i$ is stable w.r.t. S. P_i is stable if it is stable w.r.t. each of its blocks B_{ij} . Thus, a partition P_i is stable if none of its blocks B_{ij} can be split into a set of vertices that receive edges from some B_{ik} and a set of vertices that do not. A stable partition corresponds to the endpoint of a bisimulation computation: no further distinctions can be made.

This gives an algorithm for bisimulation, due to Paige and Tarjan [23] who refer to it as the "naïve algorithm". The initial partition is repeatedly refined by using its own blocks or unions of them as splitters: if S splits a block B, we replace B in the partition with the two new blocks $B \cap N^+(S)$ and $B - N^+(S)$. When no more splitters exist, the partition is stable [23] and equivalent to the full backward bisimulation of the initial partition P_0 [15].

The algorithm of Kaushik et al., Algorithm 2, modifies this naïve approach. The first difference is that in each iteration $i \in \{1, \ldots, k\}$ the partition is stabilized with respect to each of its own blocks (lines 7–16). This ensures that, after iteration *i*, the algorithm has computed the *i*-bisimulation [16], which is not the case in the naïve algorithm. Second, blocks are split as defined above, (lines 9–15). As a result, Algorithm 2 computes the *k*-backward bisimulation. To check if full bisimulation has been reached, the algorithm uses the Boolean variable wasSplit (lines 6 and 15). If this is false at the end of an iteration *i*, no block was split, so the algorithm stops early (line 17). Moreover, Algorithm 2 tracks which sets have been used as splitters (line 3), to avoid checking for stability against sets w.r.t. which the partition is already known to be stable. The algorithm provided by Kaushik et al. does not include this. If a partition *P* is stable w.r.t. *B* [23]. So after

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1 fu	unction $BISIMKAUSHIK(G = (V, E, l_V, l_E), k \in \mathbb{N})$
	/* $P := \{B_1, B_2, \dots, B_t\}$ */
2	$P \leftarrow \text{partition } V \text{ by label};$
3	usedSplitters $\leftarrow \emptyset$;
4	$\mathbf{for}\ i \leftarrow 1, \dots, k \ \mathbf{do}$
5	$P^{\operatorname{copy}} \leftarrow P;$
6	wasSplit \leftarrow false;
7	for $B^{\text{copy}} \in P^{\text{copy}} - \text{usedSplitters do}$
	/* Use blocks of copy partition to stabilize blocks of
	original partition */
8	$\mathbf{for}\;B\in P\;\mathbf{do}$
9	succ $\leftarrow B \cap N^+(B^{\text{copy}});$
10	nonSucc $\leftarrow B - N^+(B^{\text{copy}});$
	/* Split non-stable blocks */
11	if succ $\neq \emptyset$ and nonSucc $\neq \emptyset$ then
12	P.add(succ);
13	P.add(nonSucc);
14	P.delete(B);
15	wasSplit \leftarrow true;
16	usedSplitters.add $(B^{copy});$
17	$\mathbf{if} \neg \mathbf{wasSplit} \mathbf{then}$
18	break;
19	return P ;

Algorithm 2: Bisimulation Algorithm by Kaushik et al [16]

the partition P is stabilized w.r.t. a block copy B^{copy} (lines 7–16), we can add B^{copy} to the usedSplitters set and not consider it in subsequent iterations.

4.3 Generic BRS Algorithm

The parallel BRS algorithm is not specifically an implementation of k-bisimulation. Rather, one can define a graph summary model in a formal language FLUID (see Section 3.3). This model is denoted by \sim and input to the BRS algorithm, which then summarizes a graph w.r.t. \sim . In particular, the k-bisimulation models of Schätzle et al. and Kaushik et al. can be expressed in FLUID, as shown in Section 3.3. Thus, the BRS algorithm can compute k-bisimulation partitions. In other words, k-bisimulation can be incorporated into any graph summary model defined in FLUID. The BRS algorithm summarizes the graph w.r.t. \sim in parallel and uses the Signal/Collect paradigm. In Signal/Collect [26], vertices collect information from their neighbors, sent over the edges as signals. Details of the use of the Signal/Collect paradigm in our algorithm can be found in Blume et al. [5]. Briefly, the algorithm builds equivalence classes by starting with every vertex in its own singleton set and forming unions of equivalent vertices. Before outlining the algorithm, we give a necessary definition.

Definition 3. Suppose we have a graph summary GS of $G = (V, E, l_V, l_E)$ w.r.t. some $GSM \sim$. For each $v \in V$, the vertex summary vs is the subgraph of GS that defines v's equivalence class w.r.t. \sim .

We give the pseudocode of our version of the BRS algorithm in Algorithm 3 and briefly describe it below. [4] gives a step-by-step example run.

Initialization (lines 3–5). For each vertex $v \in V$, VERTEXSCHEMA computes the local schema information w.r.t. \sim_s and \sim_o of the graph summary model $(\sim_s, \sim_p, \sim_o)^k$. The method also takes into account \sim_p , i.e., the equivalence relation defined over the edge labels. Order-invariant hashes of this schema information are stored as the identifiers id_{\sim_s} and id_{\sim_o} (lines 4–5).

At the end of the initialization, every vertex has identifiers id_{\sim_s} and id_{\sim_o} . Two vertices v and v' are equivalent w.r.t. (\sim_s, \sim_p, \sim_o) , iff $v.id_{\sim_s} = v'.id_{\sim_s}$. Thus, this initialization step can be seen as iteration k = 0 of bisimulation.

Case of k = 1 bisimulation (lines 6–11). Every vertex v sends, to each inneighbor w, its id_{\sim_o} value and the label set $\ell(w, v)$ of the edge (w, v) (line 9).

Every vertex v sends, to each in-neighbor w, its id_{\sim_o} value and the label set $L = \ell(w, v)$ of the edge (w, v) (line 9). Thus, each vertex receives a set of schema $\langle L, id_{\sim_o} \rangle$ pairs from its out-neighbors, which are collated into the set M_o (line 10) and merged with an order-invariant hash to give v's new id_{\sim_s} (line 11). Here, the MERGEANDHASH (M_o) function first merges the elements of the schema message M_o received from vertex o and hashes it with an orderindependent hash function. This hash is then combined with the existing hash value $v.id_{\sim_s}$ using the xor (\oplus) operator.

Case of k > 1 bisimulation (lines 13–32). In the first iteration (lines 13–19), every vertex v sends a message to each of its out-neighbors w. The message contains v's id_{\sim_s} and id_{\sim_o} values, and the edge label set $\ell(w, v)$ (line 15). Subsequently, the incoming messages of the vertex are merged into a set of tuples with the received information $\langle \ell(w, v), id_{\sim_s} \rangle$ and $\langle \ell(w, v), id_{\sim_o} \rangle$ (lines 16 and 17). Finally, the identifiers id_{\sim_s} and id_{\sim_o} of v are updated by hashing the corresponding set (lines 18 and 19). Note that, whenever an update of an identifier value $v.id_{\sim_s}$ of vertex v is performed, the algorithm combines the old $v.id_{\sim_s}$ with the new hash value, indicated by \oplus .

In the remaining iterations, the algorithm performs the same steps (lines 20–27), but excludes the edge label set $\ell(w, v)$ when merging messages for id_{\sim_o} (line 25). When merging the messages in id_{\sim_o} , it is not necessary to consider $\ell(w, v)$, as in the iterations 2 to k-1 it is only needed to update the id_{\sim_o} values using the hash function as described above. This is possible as id_{\sim_o} by definition already contains the edge label set $\ell(w, v)$, computed in the first iteration.

In the final iteration, the identifiers are updated w.r.t. the final messages (lines 28–32). The final messages received are the values stored in the outneighbors' id_{\sim_o} values. Each vertex signals its id_{\sim_o} value to its in-neighbors (line 30). The messages a vertex receives are merged (line 31) and hashed to update the final id_{\sim_s} value (line 32). Equivalence between any two vertices v and v' can now be defined. Vertices with the same id_{\sim_s} value are merged (line 33), ending the computation.

Algorithm 3: Parallel BRS algorithm

1 fi 2	unction PARALLELSUMMARIZE $(G, (\sim_s, \sim_p, \sim_o)^k)$ returns graph symmary SG
	/* Initialization */
3	for all $v \in V$ do in parallel
4	$v.id_{\sim_s} \leftarrow hash(VERTEXSCHEMA(v, G, \sim_s, \sim_p));$
5	$v.id_{\sim_o} \leftarrow hash(VERTEXSCHEMA(v, G, \sim_o, \sim_p));$
	/* If $k=1$, only signal edge labels and $v.id_{\sim_o}$ */
6	if $k = 1$ then
7	for all $v \in V$ do in parallel
8	for all $w \in N^-(v)$ do
9	$ SENDMSGS(w, \langle \ell(w, v), 0, v.id_{\sim_o} \rangle);$
10	$M_o \leftarrow \{ \langle L, id_{\sim_o} \rangle \mid \langle L, id_{\sim_s}, id_{\sim_o} \rangle \text{ was received} \};$
11	$ v.id_{\sim_s} \leftarrow v.id_{\sim_s} \oplus \text{MergeAndHash}(M_o);$
12	else
	/* Signal initial messages. Update $v.id_{\sim_s}$ and $v.id_{\sim_o}$ */
13	for all $v \in V$ do in parallel
	/* Message each in-neighbor */
14	for all $w \in N^-(v)$ do
15	$SENDMSG(w, \langle \ell(w,v), v.id_{\sim_s}, v.id_{\sim_o} \rangle);$
	/* Collect all incoming messages of v */
16	$M_s \leftarrow \{ \langle L, id_{\sim_s} \rangle \mid \langle L, id_{\sim_s}, id_{\sim_o} \rangle \text{ was received} \};$
17	$M_o \leftarrow \{ \langle L, id_{\sim_o} \rangle \mid \langle L, id_{\sim_s}, id_{\sim_o} \rangle \text{ was received} \};$
	/* Update identifiers by hashing the messages */
18	$v.id_{\sim s} \leftarrow v.id_{\sim s} \oplus \text{MERGEANDHASH}(M_s);$
19	$v.id_{\sim_o} \leftarrow v.id_{\sim_o} \oplus \text{MERGEANDHASH}(M_o);$
	/* Signal messages $k-2$ times. As above, but we do not
	include L when updating $v.id_{\sim_o}.$ (See text.) */
20	for $i \leftarrow 2$ to $k - 1$ do
21	for all $v \in V$ do in parallel
22	for all $w \in N^-(v)$ do
23	$ \qquad \qquad$
24	$M_s \leftarrow \{ \langle L, id_{\sim_s} \rangle \mid \langle L, id_{\sim_s}, id_{\sim_o} \rangle \text{ received} \};$
25	$M_o \leftarrow \{ \langle id_{\sim_o} \rangle \mid \langle L, id_{\sim_s}, id_{\sim_o} \rangle \text{ received} \};$
26	$v.id_{\sim s} \leftarrow v.id_{\sim s} \oplus \text{MERGEANDHASH}(M_s);$
27	$ v.id_{\sim_o} \leftarrow v.id_{\sim_o} \oplus \text{MERGEANDHASH}(M_o);$
	/* Signal final messages. Update $v.id_{\sim_s}$ */
28	for all $v \in V$ do in parallel
29	for all $w \in N^{-}(v)$ do
30	SENDMSG $(w, \langle \emptyset, 0, v.id_{\sim_o} \rangle);$
31	$M_o \leftarrow \{ \langle id_{\sim_o} \rangle \mid \langle L, id_{\sim_s}, id_{\sim_o} \rangle \text{ was received} \};$
32	$v.id_{\sim_s} \leftarrow v.id_{\sim_s} \oplus \text{MergeAndHash}(M_o);$
33	$SG \leftarrow \text{FindAndMerge}(SG V)$
34	return SG .
04	roturn 50,

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Graph	V	E	$ \Sigma_V $	$r(l_V)$	$\mu(l_V(v))$	$ \Sigma_E $
Laundromat100M	$30\mathrm{M}$	$88\mathrm{M}$	33, 431	7,373	0.93 ± 44	5,630
BTC150M	$5\mathrm{M}$	$145\mathrm{M}$	69	137	1.04 ± 0.26	10,750
BTC2B	$80\mathrm{M}$	$1.92\mathrm{B}$	113, 365	576, 265	0.95 ± 1.82	38,136
BSBM100M	$18\mathrm{M}$	$90\mathrm{M}$	1,289	2,274	1.02 ± 0.13	39
BSBM1B	$172\mathrm{M}$	$941\mathrm{M}$	6,153	27,306	1.03 ± 0.18	39

Table 2: Statistics of the datasets.

We show in [4] that Algorithm 3 computes k-bisimulation of a graph with m edges in time O(km). As a modification of the algorithm in [6], the algorithm is correct, as long as hash collisions are avoided.

5 Experimental Apparatus

5.1 Datasets

We experiment with smaller and larger as well as real-world and synthetic graphs. Table 2 lists statistics of these datasets, where $r(l_V) = |\{l_V(v) \mid v \in V\}|$ is the number of different label sets (range) and $\mu(|l_V(v)|)$ is the average number of labels of a vertex $v \in V$.

Three real-world datasets were chosen. The Laundromat100M dataset contains 100 M edges of the LOD Laundromat service [1]. This service automatically cleaned existing linked datasets and provided the cleaned version on a publicly accessible website. The BTC150M and BTC2B datasets contain, respectively, around 150 million and 1.9 billion edges of the Billion Triple Challenge 2019 (BTC2019) dataset [14]. 93% of the total edges originate from Wikidata [29]. BTC150M is the first chunk of the 1.9 billion edges. For synthetic datasets, two versions of the Berlin SPARQL Benchmark (BSBM) [2] were used. The BSBM data generator produces RDF datasets that simulate an e-commerce use case. BSBM100M was generated with 284,826 products and has about 17.77 million vertices and 89.54 million edges. BSBM1B was generated with 2,850,000 products and has about 172 million vertices and 941 million edges.

5.2 Procedure

An experiment consists of the algorithm to run, the dataset to summarize, and the bisimulation degree k. In case of the BRS algorithm, it additionally consists of the graph summary model to use. We have two different graph summary models defined by Schätzle et al. [25] and Kaushik et al. [16]. Each model comes in two implementations, one native implementation as defined by the original authors and a generic implementation through our hash-based version of the BRS algorithm. We chose the bisimulation algorithms of Kaushik et al. and Schätzle et al. as they represent two typical variants of backward and forward k-bisimulation models as found in the literature (cf. Section 2). This choice of algorithms allows

us to demonstrate that our algorithm can be applied to different settings. We use the terms *BRS-Schätzle* and *BRS-Kaushik* to refer to our implementations of the two GSMs BRS algorithm; we refer to our single-purpose implementations of these two GSMs as *native Schätzle* and *native Kaushik*.

The four algorithms are applied on the five datasets, giving 20 experiments. Each experiment is executed with a bisimulation degree of k = 1, ..., 10, using the following procedure. We run the algorithms six times with the specific configuration. We use the first run as a warm-up and do not account it for our measurements. The next five runs are used to measure the variables.

5.3 Implementation

All algorithms, i.e., the native algorithms of Schätzle et al. and Kaushik et al. and their generic BRS-variants are implemented using the same underlying framework and paralellization approach, i.e., are implemented in Scala upon the Apache Spark Framework. This API offers flexible support for parallel computation and message passing, which enables implementation of Map-Reduce and Signal-Collect routines. We use an Ubuntu 20 system with 32 cores and 2 TB RAM. The Apache Spark contexts were given the full resources. Time and memory measurements were taken using the Apache Spark Monitoring API.

5.4 Measures

We evaluate the algorithms' running time and memory consumption. For every run of an experiment, we report the total run time, the run time of each of the k iterations, and the maximum JVM on-heap memory consumption.

6 Results

We present full results for each algorithm, iteratively calculating k-bisimulation for every value of k = 1, ..., 10 and every dataset, in Figure 2. Table 3 summarizes the average total run time (minutes) for each experiment, for the computation of 10-bisimulation. The BRS algorithm takes an additional initialization step (see Algorithm 3, lines 13–19). Table 4 reports the maximum JVM on-heap memory in GB for each experiment. The BRS-Schätzle algorithm computes the 10-bisimulation the fastest on all datasets, except for BSBM100M, where BRS-Kaushik is fastest. Native Schätzle consumes the least memory on all smaller datasets. Native Schätzle consumes slightly more memory on BSBM1B than BRS, whereas on BTC2B, the memory consumption is about the same.

Smaller Datasets (100M + Edges). Figure 2 shows the average run time (minutes) for each of the ten iterations on the smaller datasets. The native Kaushik experiments take much longer than the others (Figures 2a, 2c, and 2e), so we provide plots without native Kaushik (Figures 2b, 2d, and 2f), to allow easier comparison between the other algorithms.



Fig. 2: Average iteration times (minutes) on the smaller datasets in (a) to (f) and larger datasets in (g) and (h).

	Schätzle	e et al.	Kaus	shik et al.
	BRS	Native	BRS	Native
Laundromat100M	$5.56_{(0.50)}$	$7.72_{(0.07)}$	$5.60_{(0.15)}$	586.66 (21.09)
BTC150M	$4.08_{(1.55)}$	$6.14_{(0.38)}$	$4.54_{(1.34)}$	78.02 (3.71)
BTC2B	$61.96_{(11.38)}$	$83.74_{(1.96)}$	85.92 (13.6)	out of time
BSBM100M	$6.46_{(0.08)}$	$9.40_{(0.06)}$	$5.20_{(0.50)}$	77.84 (2.41)
BSBM1B	$54.44_{(4.35)}$	85.98 (3.83)	$64.00_{(3,22)}$	out of memory

Table 3: Total run time (in minutes) needed for computing the k = 1, ..., 10 bisimulation (average and standard deviation over 5 runs).

	Schätzl	e et al.	Ka	ushik et al.
	BRS	Native	BRS	Native
Laundromat100M	211.5	147.9	210.5	335.1
BTC150M	140.6	107.7	130.1	181.7
BTC2B	$1,\!249.1$	1,249.7	$1,\!249.3$	out of time
BSBM100M	248.6	113.1	172.0	327.0
BSBM1B	1,248.2	1,335.4	1,249.2	out of memory

Table 4: Maximal JVM on-heap memory (in GB) used for k = 1, ..., 10 bisimulation. We provide the maximal memory usage determined over five runs, instead of the average and standard deviation, in order to demonstrate what memory is needed to perform the bisimulations without running out of memory.

BRS-Schätzle and native Schätzle have relatively constant iteration time on all smaller datasets. For example, on Laundromat100M (Figure 2b), native Schätzle computes an iteration in about 0.7 to 0.95 minutes. BRS is slightly faster on all smaller datasets, but uses more memory (Table 4). BRS-Kaushik shows similar results to BRS-Schätzle. Again iteration time is relatively constant on all datasets. As before, it is fastest on BTC150M.

Native Kaushik shows a different behavior across the datasets. Iteration time varies on all datasets: the iteration time increases in early iterations until it reaches a maximum value, and then decreases. The only exception to this behavior occurs on BTC150M, where the iteration time decreases from iteration one to two before following the described pattern for the remaining iterations. For example, on Laundromat100M (Figure 2a) an iteration takes about 10 to 100 minutes. The maximum iteration time is reached in iteration six. Native Kaushik runs fastest on BSBM100M, taking about 77.84 minutes on average. BRS-Kaushik is much faster on all smaller datasets (Table 3) and uses slightly less memory (Table 4).

Larger Datasets (1B+ Edges). BRS-Schätzle computes the k = 1, ..., 10 bisimulation on BTC2B and BSBM1B with a relatively constant iteration time. The total run time (Table 3), is lowest on BSBM1B: 54.44 minutes on average. Native Schätzle also has nearly constant running time across iterations. The average running time for all ten iterations is lowest on BTC2B: 83.74 minutes on average, compared to 85.98 minutes on BSBM1B. Comparing the run times of

the two implementations, BRS is 37% on the BSBM1B dataset and 26% faster on BTC2B. Both algorithms require about the same memory during execution on BTC2B. On BSBM1B, native Schätzle uses slightly more memory than BRS. Native Kaushik did not complete one iteration on BTC2B in 24 hours, so execution was canceled. The algorithm ran out of memory on BSBM1B (Table 4).

Finally, BRS-Kaushik also has similar iteration times. On BSBM1B (Figure 2h), computing a bisimulation iteration ranges from about 4 to 6 minutes. On BTC2B (Figure 2g), the execution times for iterations one to three range from 2 to 4 minutes. Iterations four to nine take about 8 minutes each. For the final iteration ten, the running time slightly increases to about 9.5 minutes.

7 Discussion

Main Results. Our results show that, on all datasets, the generic BRS algorithm outperforms the native bisimulation algorithms of Schätzle et al. and Kaushik et al. for k = 10 (see Table 3). Since the BRS algorithm has an initialization phase, which is not present in the two native algorithms, we examine the data more closely to see at which value of k BRS begins to outperform the native implementations. We provide the exact numbers per iteration together with the standard deviation and averaged over five executions in Appendix D. Our generic BRS-Kaushik outperforms the native Kaushik et al. for all k: this is unsurprising, as BRS parallelizes the original sequential algorithm. The time taken to compute k-bisimulation is the total time for iterations up to, and including, k. The comparison of BRS-Schätzle with the native Schätzle et al. algorithm shows that our generic algorithm begins to outperform the native one at k=3for the Laundromat100M, BSBM100M, and BSBM1B; at k = 4 for BTC150M; and at k = 5 for BTC2B. Again, we emphasize that BRS is a generic algorithm supporting all graph summary models definable in FLUID (see introduction), whereas native algorithms compute only one version of bisimulation each.

Note that, for each dataset and each algorithm, we iteratively compute in total ten bisimulation-based graph summaries using every value of k from 1 to 10. As can be seen in Figure 2, the execution times per bisimulation iteration are fairly constant over all iterations for all datasets for BRS and Schätzle et al. The native Kaushik et al. algorithm sequentially computes the refinement of the partitions, so execution time directly relates to the number of splits per k-bisimulation iteration. This can be seen by the curve in the plots of the smaller datasets (see Figures 2a, 2c, and 2e). In particular, Kaushik et al. detect that full bisimulation has been reached and do not perform further computations: this happens after computing the 7th iteration (7-bisimulation, see Figure 2e) on the BSBM100M dataset but not on the other datasets on which the execution of the Kaushik et al. algorithm completed. This kind of termination check could also be added to the BRS algorithm but this is nontrivial, as we discuss below.

This consistent per-k iteration runtime is explained by each iteration of the BRS algorithm processing every vertex, considering all its neighbors. Thus, the

running time of the iteration depends primarily on the size of the graph, and not on how much the k-bisimulation that is being computed differs from the (k-1)bisimulation that was computed at the previous iteration. The native Schätzle et al. algorithm behaves in the same way. There is some variation in per-iteration execution time but this may be because, as each vertex is processed, duplicate incoming messages must be removed. The time taken for this will depend on the distribution of the incoming messages, which will vary between iterations.

We consider experimenting with values between k = 1 and 10 to be appropriate. The native Kaushik et al. algorithm terminates when full bisimulation is reached, which shows that full bisimulation is reached at k = 7 on the BSBM100M dataset, but has not been reached up to k = 10 for Laundromat100 and BTC150M. The rapidly decreasing per-iteration running times for native Kaushik et al. for BTC150M (Figure 2c) suggests that full bisimulation will be reached in a few more iterations. A similar curve can be observed for Laundromat100 (Figure 2a), but the running time for the k = 10 iteration is higher, suggesting that full bisimulation will not be reached for several iterations. Thus, on these datasets, computing 10-bisimulations is a reasonable thing to do, as full bisimulation has not yet been reached. Note that, for the other algorithms that we consider (native Schätzle, and the two instances of BRS), per-iteration running time is largely independent of k and of whether full bisimulation as been reached (see the per-iteration measurements on BSBM100M in Table 9).

Conversely, our primary motivation is graph summarization. When two vertices in a graph are 10-bisimilar, this means they have equivalent neighborhoods out to distance 10. This means that they are already "largely similar" for that value of k. Thus, we feel that, having computed k-bisimulation for a relatively large value of k, there is little advantage in going to k+1. In other words, adding another iteration of k + 1 still leaves us with vertices that are "largely similar".

Scalability to Graph Size. From the execution time of computing kbisimulation for k = 1, ..., 10, we observe that BRS-Schätzle, BRS-Kaushik. and native Schätzle scale linearly. Here, we consider the scalability of the algorithms with respect to graph size. To this end, we fix on iterations $k = 1, \ldots, 10$ and compare the total time taken to compute these bisimulations for input graphs of different (large) sizes. BTC150M contains approximately 5M vertices and 145M edges. BTC2B has around 80M vertices and 2B edges, which is a factor of about 18 and 14 larger, respectively. BRS-Schätzle takes 4.08 minutes, BRS-Kaushik 4.54 minutes, and native Schätzle 6.14 minutes on BTC150M. On BTC2B, they take 15 times, 19 times, and 14 times longer. BSBM1B contains about 10 times as many vertices and edges as BSBM100M. The experiments on BSBM1B took about 10 times longer for BRS-Schätzle, 12 times for BRS-Kaushik, and 9 times for native Schätzle. Thus, the scaling factor of the execution times is approximately equal to that of the graph's size. Finally, the total runtimes of native Kaushik on the different graphs indicate that the algorithm does not scale linearly with the input graph's size. The initial partition P_0 has one block per label set and Laundromat100M has many different label sets (Definition 2). The algo-

rithm checks stability of every block against every other, leading to a run time that is quadratic in the number of blocks.

Generalization and Threats to Validity. We use synthetic and realworld graphs, which is important to analyze the practical application of an algorithm [5, 17]. Two GSMs were used for evaluation of our hash-based BRS algorithm. The GSM of Schätzle et al. computes a forward k-bisimulation, based on edge labels (Definition 1). The GSM of Kaushik et al. computes a backward kbisimulation based on vertex labels (Definition 2). Hence, the two GSMs consider different structural features for determining vertex equivalence. Regardless, for both GSMs, the BRS algorithm scales linearly with the number of bisimulation iterations and the input graph's size and computes the aggregated k = 1, ..., 10bisimulations the fastest on every dataset.

All algorithms are implemented in Scala, in the same framework. Each algorithm was executed using the same procedure on the same machine with exclusive access during the experiments. Each experimental configuration was run six times. The first run is discarded in the evaluations to address side effects.

8 Conclusion and Future Work

We focus on the performance (runtime and memory use) of our generic, parallel BRS algorithm for computing different bisimulation variants, and how this performance compares to specific algorithms for those bisimulations, due to Schätzle et al. and Kaushik et al. Our experiments comparing $k = 1, \ldots, 10$ bisimulations on large synthetic and real-world graphs show that our generic, hash-based BRS algorithm outperforms the respective native bisimulation algorithms on all datasets for all $k \geq 5$ and for smaller k in some cases. The experimental results indicate that the parallel BRS algorithm and native Schätzle et al. scale linearly with the number of bisimulation iterations and the input graph's size. Our experiments also show that our generic BRS-Kaushik algorithm effectively parallelizes the original sequential algorithm of Kaushik et al., showing runtime performance similar to BRS-Schätzle. Overall, we recommend using our generic BRS algorithm over implementations of specific algorithms of k-bisimulations. Due to the support of a formal language for defining graph summaries [6], the BRS approach is flexible and easily adaptable, e.g., to changes in the desired features and user requirements, without sacrificing performance.

Future work includes incorporating a check (similar to Kaushik et al.) for whether full bisimulation has been reached. This is nontrivial as the algorithm of Kaushik et al. is based on splitting partitions, whereas BRS is based on describing the equivalence class in which each vertex lives. It is easy to check that no classes have been split, but harder to check that every pair of vertices that had the same description at the previous iteration have the same description at the current iteration, since the description of every vertex changes at each iteration.

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Appendix

A Additional Dataset Statistics

Table 5 lists degree statistics of the datasets, where d(G) is the average degree of vertices in G, $d_G(v)$ the degree of a vertex $v \in V$, and $\Delta(G)$ the maximum degree in G.

Graph	$\mu(d_G(v))$	$\Delta(G)$	$\mu(d_G^-(v))$	$\Delta^{-}(G)$	$\mu(d_G^+(v))$	$\Delta^+(G)$
L'mat100M	5.89 ± 569	1,570,748	2.95 ± 559	1,570,748	2.95 ± 108	545,688
BSBM100M	$9.76 \pm 1,144$	2,273,014	5.04 ± 1144	2,273,014	5.04 ± 6	76
BTC150M	$58.43 \pm 5,655$	5,629,275	29.22 ± 504	283,686	$29.22 \pm 5,629$	5,628,254
BSBM1B	$10.58 \pm 3,862$	23,924,441	5.46 ± 3862	23,924,441	5.46 ± 6	85
BTC2B	$48.40 \pm 17,119$	65,879,409	24.20 ± 1556	3,856,778	$24.20 \pm 17,038$	65, 878, 298

Table 5: Degree Statistics of the Datasets.

Laundromat100M has the smallest average and maximum values for both degree and in-degree. BTC150M has the highest average and maximum degrees and the highest maximum in-degree. BSBM100M has the largest maximum indegree and smallest maximum out-degree of our datasets.

We note that standard deviations in Table 5 are surprisingly large. This is due to some vertices having degrees orders of magnitude larger than the average, as shown by the maximum degrees.

We note that, in the real-world datasets, average total degree is average indegree plus average out-degree, to the precision quoted. This is expected, because every edge adds one to the total degree of each of its endpoints. This is not the case for the two versions of the synthetic BSBM dataset. This dataset contains relatively many self-loops and, by convention, a self-loop adds 1 to a vertex's total degree, while also adding 1 to its in- and out-degrees.

B Results of the BRS Algorithm

As the BRS algorithm takes an additional initialization step (see Algorithm 3, lines 13–19), Table 6 shows the breakdown of the running time of the algorithms in initialization and computing the actual k bisimulation iterations. It is separated into average run time for initialization and computation of all ten iterations.

On all smaller datasets, i. e., Laundromat100M, BSBM100M, and BTC150M and the initialization step of BRS-Schätzle takes about 1 minute on average (Table 6). Regarding BRS-Kaushik, the initialization also takes around 1 minute on all smaller datasets (Table 6). Regarding the larger dataset BSBM1B, the BRS-Schätzle algorithm takes 10.20 minutes for initialization and 44.24 minutes for the 10 iterations. On BTC2B the algorithm needs 16.22 minutes to initialize

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	BRS-	Schätzle et al.	BRS-Kaushik et a			
	Init.	Iterations	Init.	Iterations		
Laundromat100M	1.10	4.46	1.10	4.50		
BSBM100M	1.00	5.46	1.05	4.15		
BTC150M	1.47	2.61	1.44	3.10		
BSBM1B	10.20	44.24	10.43	53.57		
BTC2B	16.22	45.74	16.29	69.63		

Table 6: Initialization and bisimulation iteration (minutes) of average total run time for BRS.

the graph and 45.74 minutes to compute the 10 iterations. The BRS-Kaushik takes on BSBM1B 10.43 minutes for initialization and 53.57 minutes for the 10 iterations. On BTC2B, the BRS-Kaushik algorithm needs 16.29 minutes for initialization and 69.93 minutes for the 10 iterations.

Native Kaushik also has an initialization step, where the vertices are partitioned based on their label. This partitioning is done in parallel and took less than a second, so we do not account for it separately.

C Discussion of the Results of the Kaushik Algorithm

Experimental results for native Kaushik indicate that the algorithm does not scale linearly with the input graph's size. Its worst-case complexity is $\mathcal{O}(k \cdot m)$ [16], where m is the number of edges in the input graph.

A reason for the long running time of native Kaushik on Laundromat100M could be the size of the initial partition P_0 . It depends on the number of different label sets present in the graph's vertices (Definition 2). Laundromat100M contains 33, 431 different label sets, which is much higher than for BTC150M with 69 and BSBM100M with 1, 289 (Table 2). As a consequence, the algorithm has to perform stability checks and (potential) splits on blocks more often than on the other datasets. This does not contradict the similar run times on BTC150M and BSBM100M (69 label sets vs. 1, 289 label sets). First, BSBM100M reaches full bisimulation w.r.t. Definition 2 in iteration seven and hence execution is stopped early. Second, as can be seen in Figure 2e, the iteration time starts to decrease rapidly from iteration four to five on BSBM100M, going down from about 20 minutes to about 1 minute. This indicates that the partition hardly changes any further and therefore only a small number of stability checks and splits are performed in iterations five to seven.

Native Kaushik operates on the blocks of the graph's current partition. In each iteration *i*, the algorithm produces a partition P_i , which is stable w.r.t. all the blocks in P_{i-1} [16]. Consequently, if a block is split into two new blocks, these must also be checked for stability in that iteration (see also discussion in Section 5.3). Hence, the more blocks are split in an iteration, the more steps must be performed by the algorithm. In addition, bisimulation relationships between the vertices change far more often in early iterations [25]. This explains

the specific shape of the run time curve (Figures 2a, 2c and 2e) for the native Kaushik et al. algorithm. The exception is BTC150M (Figure 2c), where the iteration time first decreases from iteration one to two, indicating that the 1-and 2-bisimulations of this graph are very similar.

D Detailed Results of Experiments

We provide the detailed experimental results. The results are split into results for the smaller datasets (Laundromat100M, BTC150M, BSBM100M) shown in Tables 7 to 9. The results for the larger datasets (BTC2B, BSBM1B) are shown in Tables 10 and 11. The tables for BRS include an additional iteration 0, which corresponds to the initialization routine of the algorithm.

Run			Aggregates											
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	1.1	0.4	0.5	0.5	0.4	0.4	0.6	0.6	0.5	0.5	0.5	6.0	0.49	0.07
2	1.1	0.3	0.4	0.3	0.3	0.3	0.6	0.4	0.3	0.3	0.3	4.6	0.35	0.09
3	1.1	0.4	0.5	0.4	0.7	0.6	0.5	0.4	0.4	0.4	0.4	5.8	0.47	0.10
4	1.1	0.4	0.5	0.4	0.4	0.7	0.5	0.5	0.4	0.4	0.5	5.8	0.47	0.09
5	1.1	0.4	0.4	0.4	0.4	0.8	0.4	0.4	0.4	0.4	0.5	5.6	0.45	0.12
μ	1.10	0.38	0.46	0.40	0.44	0.56	0.52	0.46	0.40	0.40	0.44	5.56		
σ	0.00	0.04	0.05	0.06	0.14	0.19	0.07	0.08	0.06	0.06	0.08	0.50		

(a) BRS algorithm executed with GSM Schätzle

Run					Aggregates								
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	1.1	0.8	0.7	0.8	0.8	0.7	0.7	0.7	0.7	0.8	7.8	0.78	0.12
2	1.1	0.8	0.7	0.8	0.7	0.7	0.7	0.7	0.7	0.7	7.6	0.76	0.12
3	0.7	0.8	1.1	0.8	0.7	0.8	0.8	0.7	0.7	0.7	7.8	0.78	0.12
4	0.7	0.8	1.1	0.8	0.7	0.8	0.7	0.7	0.7	0.7	7.7	0.77	0.12
5	0.7	0.7	1.1	0.8	0.7	0.8	0.8	0.7	0.7	0.7	7.7	0.77	0.12
μ	0.86	0.78	0.94	0.80	0.72	0.76	0.74	0.70	0.70	0.72	7.72		
σ	0.00	0.00	0.00	0.08	0.04	0.07	0.00	0.00	0.05	0.04	0.07		

Run		Iteration Ag												gregates	
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ	
1	1.1	0.3	0.4	0.4	0.4	0.4	0.4	0.6	0.7	0.5	0.5	5.7	0.46	0.11	
2	1.1	0.3	0.4	0.3	0.3	0.4	0.5	0.7	0.4	0.4	0.5	5.3	0.42	0.12	
3	1.1	0.3	0.4	0.4	0.5	0.7	0.5	0.5	0.4	0.4	0.5	5.7	0.46	0.10	
4	1.1	0.3	0.4	0.4	0.4	0.4	0.7	0.6	0.5	0.4	0.5	5.7	0.46	0.11	
5	1.1	0.3	0.4	0.4	0.4	0.7	0.4	0.5	0.4	0.4	0.6	5.6	0.45	0.11	
μ	1.10	0.30	0.40	0.38	0.40	0.52	0.50	0.58	0.48	0.42	0.52	5.60			
σ	0.00	0.00	0.00	0.04	0.06	0.15	0.11	0.07	0.12	0.04	0.04	0.15			

(b) Schätzle algorithm

(c) BRS algorithm executed with GSM Kaushik

Run					Iter	ation					Agg	regat	es
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	11.4	21.2	35.1	60.0	85.5	106.7	98.8	81.8	59.3	44.1	603.9	60.39	30.9
2	13.6	22.7	35.8	66.8	76.8	91.9	85.6	69.6	52.1	35.0	549.9	54.99	25.81
3	10.9	19.3	31.9	53.7	81.3	100.1	98.7	79.9	62.5	42.5	580.8	58.08	30.21
4	11.2	21.1	33.3	58.2	78.6	100.5	98.1	85.1	61.6	41.2	588.9	58.89	30.04
5	11.8	21.3	34.1	55.1	81.8	107.7	97.0	91.9	69.0	40.1	609.8	60.98	31.81
μ	11.78	21.12	34.04	58.76	80.80	101.38	95.64	81.66	60.90	40.58	586.66		
σ	0.96	1.08	1.37	4.59	2.97	5.67	5.06	7.28	5.45	3.09	21.09		

(d) Kaushik algorithm

Table 7: Detailed Results (minutes) on Laundromat100M for 10-bisimulation.

Run					Ite	erati	on					Agg	grega	ates
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	1.4	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	2.4	0.10	0.00
2	1.4	0.4	0.4	0.4	0.8	0.5	0.4	0.4	0.4	0.5	0.4	6.0	0.46	0.12
3	1.7	0.2	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.6	3.3	0.16	0.15
4	1.4	0.2	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.4	2.8	0.14	0.09
5	1.7	0.4	0.4	0.4	0.6	0.4	0.4	0.4	0.4	0.4	0.4	5.9	0.42	0.06
μ	1.47	0.26	0.22	0.22	0.34	0.24	0.22	0.22	0.22	0.24	0.38	4.08		
σ	0.14	0.12	0.15	0.15	0.3	0.17	0.15	0.15	0.15	0.17	0.16	1.55		

Run					Itera	ation	L				Agg	grega	tes
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	0.6	0.6	0.7	0.8	0.7	0.6	0.6	0.6	0.6	0.6	6.4	0.64	0.07
2	0.5	0.6	0.7	0.8	0.6	0.6	0.6	0.6	0.6	0.6	6.2	0.62	0.07
3	0.6	0.6	0.8	0.8	0.6	0.6	0.6	0.6	0.6	0.6	6.4	0.64	0.08
4	0.5	0.6	0.8	0.8	0.6	0.6	0.6	0.6	0.6	0.6	6.3	0.63	0.09
5	0.5	0.5	0.6	0.7	0.6	0.5	0.5	0.5	0.5	0.5	5.4	0.54	0.07
μ	0.54	0.58	0.72	0.78	0.62	0.58	0.58	0.58	0.58	0.58	6.14		
σ	0.05	0.04	0.07	0.04	0.04	0.04	0.04	0.04	0.04	0.04	0.38		

Run					Ite	erati	on					Agg	grega	tes
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	1.4	0.4	0.4	0.3	0.4	0.6	0.4	0.4	0.4	0.4	0.4	5.5	0.41	0.07
2	1.4	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.7	3.0	0.16	0.18
3	1.4	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.5	2.8	0.14	0.12
4	1.4	0.4	0.5	0.5	0.4	0.4	0.4	0.4	0.4	0.4	0.4	5.6	0.42	0.04
5	1.4	0.4	0.6	0.5	0.4	0.4	0.4	0.4	0.4	0.4	0.5	5.8	0.44	0.07
μ	1.44	0.28	0.34	0.30	0.28	0.32	0.28	0.28	0.28	0.28	0.50	4.54		
σ	0.14	0.15	0.21	0.18	0.15	0.19	0.15	0.15	0.15	0.15	0.11	1.34		

(b) Schätzle algorithm

(c) BRS algorithm executed with GSM Kaushik

Run					Itera	ation					Agg	rega	tes
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	3.9	0.9	3.7	12.2	17.0	17.8	12.7	6.6	2.8	1.0	78.6	7.86	6.17
2	3.8	0.9	3.7	11.2	16.8	17.0	12.9	6.0	2.9	1.5	76.7	7.67	5.92
3	4.1	0.9	3.9	11.7	17.6	17.6	12.6	6.1	3.1	1.1	78.7	7.87	6.14
4	3.9	0.9	4.0	14.1	20.5	18.5	12.2	6.0	2.5	1.2	83.8	8.38	6.95
5	3.6	0.9	3.5	10.8	15.5	16.0	12.0	6.3	2.7	1.0	72.3	7.23	5.55
μ	3.86	0.90	3.76	12.00	17.48	17.38	12.48	6.20	2.80	1.16	78.02		
σ	0.16	0.00	0.17	1.15	1.66	0.84	0.33	0.23	0.20	0.19	3.71		

(d) Kaushik algorithm

Table 8: Detailed Results (minutes) on BTC150M for 10-bisimulation.

Run					Ite	erati	on					Agg	grega	\mathbf{tes}
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	1.0	0.5	0.5	0.5	0.6	0.8	0.5	0.5	0.5	0.5	0.4	6.3	0.53	0.10
2	1.0	0.5	0.6	0.5	0.5	0.9	0.5	0.5	0.5	0.5	0.5	6.5	0.55	0.12
3	1.0	0.5	0.6	0.5	0.5	0.9	0.5	0.5	0.5	0.5	0.5	6.5	0.55	0.12
4	1.0	0.5	0.5	0.5	0.6	0.9	0.5	0.5	0.5	0.5	0.5	6.5	0.55	0.12
5	1.0	0.5	0.5	0.5	0.9	0.5	0.5	0.5	0.5	0.6	0.5	6.5	0.55	0.12
μ	1.00	0.50	0.54	0.50	0.62	0.80	0.50	0.50	0.50	0.52	0.48	6.46		
σ	0.00	0.00	0.05	0.00	0.15	0.15	0.00	0.00	0.00	0.04	0.04	0.08		

Run					Itera	ation	L				Agg	grega	tes
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	0.9	1.3	0.9	0.9	0.9	0.9	0.9	0.9	0.9	0.9	9.4	0.94	0.12
2	0.9	1.0	1.2	0.9	1.0	0.9	0.9	0.9	0.9	0.9	9.5	0.95	0.09
3	0.9	0.9	1.2	0.9	0.9	0.9	0.9	0.9	0.9	0.9	9.3	0.93	0.09
4	0.9	1.0	1.2	0.9	0.9	0.9	0.9	0.9	0.9	0.9	9.4	0.94	0.09
5	0.9	1.0	1.2	0.9	0.9	0.9	0.9	0.9	0.9	0.9	9.4	0.94	0.09
μ	0.90	1.04	1.14	0.90	0.92	0.90	0.90	0.90	0.90	0.90	9.40		
σ	0.00	0.14	0.12	0.00	0.04	0.00	0.00	0.00	0.00	0.00	0.06		

(b) Schätzle algorithm

Run					Ite	erati	on					Agg	grega	tes
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	1.0	0.4	0.5	0.4	0.4	0.6	0.7	0.4	0.4	0.4	0.4	5.6	0.46	0.10
2	1.0	0.3	0.4	0.3	0.3	0.3	0.3	0.5	0.5	0.3	0.3	4.5	0.35	0.08
3	1.1	0.4	0.4	0.4	0.5	0.7	0.5	0.4	0.4	0.4	0.5	5.7	0.46	0.09
4	1.0	0.4	0.5	0.4	0.4	0.6	0.6	0.4	0.4	0.4	0.4	5.5	0.45	0.08
5	1.1	0.3	0.4	0.3	0.3	0.3	0.3	0.5	0.5	0.3	0.4	4.7	0.36	0.08
μ	1.05	0.36	0.44	0.36	0.38	0.50	0.48	0.44	0.44	0.36	0.40	5.20		
σ	0.05	0.05	0.05	0.05	0.07	0.17	0.16	0.05	0.05	0.05	0.06	0.50		

(c) BRS algorithm executed with GSM Kaushik

Run				It	erat	ion					Agg	rega	tes
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	8.0	16.5	33.9	19.0	0.8	0.6	0.6	0.0	0.0	0.0	79.4	7.94	11.03
2	8.8	17.2	29.3	22.1	0.9	0.6	0.6	0.0	0.0	0.0	79.5	7.95	10.44
3	8.2	16.2	30.7	22.6	0.8	0.7	0.7	0.0	0.0	0.0	79.9	7.99	10.71
4	8.0	19.0	26.4	21.5	0.8	0.6	0.6	0.0	0.0	0.0	76.9	7.69	9.97
5	9.2	15.9	25.8	20.6	0.8	0.6	0.6	0.0	0.0	0.0	73.5	7.35	9.43
μ	8.44	16.96	29.22	21.16	0.82	0.62	0.62	0.00	0.00	0.00	77.84		
σ	0.48	1.11	2.96	1.27	0.04	0.04	0.04	0.00	0.00	0.00	2.41		

(d) Kaushik algorithm

Table 9: Detailed Results (minutes) on BSBM100M for 10-bisimulation.

Run					Ite	ratio	on					Agg	rega	\mathbf{tes}
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	16.0	4.7	5.2	4.9	5.0	5.6	5.1	5.1	5.1	5.1	5.9	67.7	5.17	0.33
2	16.0	4.5	5.1	4.9	5.0	5.9	5.0	5.0	5.8	5.1	5.3	67.6	5.16	0.40
3	16.4	1.6	1.8	1.7	1.9	2.2	1.9	1.9	1.9	1.9	6.0	39.2	2.28	1.25
4	16.0	4.5	5.1	4.9	5.2	5.5	5.1	5.0	5.8	5.2	5.4	67.7	5.17	0.33
5	16.4	4.6	4.9	4.8	6.2	5.7	5.0	5.0	4.9	5.0	5.1	67.6	5.12	0.45
μ	16.22	3.98	4.42	4.24	4.66	4.98	4.42	4.40	4.70	4.46	5.54	61.96		
σ	0.16	1.19	1.31	1.27	1.45	1.4	1.26	1.25	1.45	1.28	0.35	11.38		

Run					Itera	ation	L				Agg	rega	\mathbf{tes}
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	8.3	8.7	8.7	9.2	8.7	8.7	8.7	8.7	8.7	8.9	87.3	8.73	0.21
2	8.1	8.3	8.3	8.8	8.3	8.3	8.3	8.3	8.3	8.3	83.3	8.33	0.17
3	7.9	8.2	8.2	8.2	8.1	8.2	8.1	8.6	8.1	8.2	81.8	8.18	0.17
4	7.9	8.3	8.3	8.4	8.4	8.7	8.8	8.4	8.4	8.5	84.1	8.41	0.23
5	7.9	8.2	8.1	8.2	8.3	8.3	8.3	8.3	8.3	8.3	82.2	8.22	0.12
μ	8.02	8.34	8.32	8.56	8.36	8.44	8.44	8.46	8.36	8.44	83.74		
σ	0.16	0.19	0.20	0.39	0.20	0.22	0.27	0.16	0.20	0.25	1.96		

Run					Ite	ratio	on					Agg	rega	tes
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	16.1	1.9	2.5	3.8	7.0	7.0	7.5	7.2	7.4	7.2	7.3	74.9	5.88	2.11
2	16.1	2.1	2.7	4.2	7.0	8.3	7.5	8.0	7.6	8.0	11.8	83.3	6.72	2.77
3	16.6	2.4	2.8	4.3	7.9	7.8	8.0	7.8	7.8	7.8	10.9	84.1	6.75	2.55
4	16.1	1.9	2.6	3.9	7.1	7.3	7.7	7.2	7.3	7.1	7.1	75.3	5.92	2.10
5	16.6	5.2	5.9	7.4	11.2	11.1	11.2	10.7	10.6	10.8	11.3	112.0	9.54	2.28
μ	16.29	2.70	3.30	4.72	8.04	8.30	8.38	8.18	8.14	8.18	9.68	85.92		
σ	0.20	1.26	1.30	1.35	1.62	1.47	1.42	1.30	1.24	1.35	2.05	13.60		

(b) Schätzle algorithm

(c) BRS algorithm executed with GSM Kaushik

Table 10: Detailed Results (minutes) on BTC2B for 10-bisimulation.

Run	Iteration												Aggregates			
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ		
1	10.1	3.9	4.7	4.3	4.3	4.5	5.3	4.7	5.1	4.7	5.7	57.3	4.72	0.50		
2	10.1	2.9	3.7	3.3	3.4	3.3	4.1	4.1	4.1	3.8	3.1	45.9	3.58	0.42		
3	10.0	3.9	4.8	4.3	4.4	4.5	5.2	4.7	4.9	4.5	4.2	55.4	4.54	0.36		
4	10.1	3.9	4.8	4.4	4.4	4.5	5.2	4.9	5.2	4.5	5.8	57.7	4.76	0.51		
5	10.0	3.9	4.8	4.4	4.4	4.6	5.4	4.8	5.0	4.4	4.2	55.9	4.59	0.41		
μ	10.20	3.70	4.56	4.14	4.18	4.28	5.04	4.64	4.86	4.38	4.60	54.44				
σ	0.62	0.40	0.43	0.42	0.39	0.49	0.48	0.28	0.39	0.31	1.02	4.35				

Run	Iteration											Aggregates			
	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ		
1	9.0	8.1	8.1	8.2	8.3	8.2	8.2	8.5	8.2	8.2	83.0	8.30	0.26		
2	10.0	8.3	8.2	8.3	8.4	8.5	8.4	8.4	8.3	8.3	85.1	8.51	0.50		
3	8.8	8.9	8.9	9.0	9.0	12.5	9.3	9.0	9.0	9.0	93.4	9.34	1.06		
4	8.1	8.2	8.4	8.2	8.3	8.6	8.7	8.2	8.2	8.2	83.1	8.31	0.19		
5	8.5	8.4	8.3	8.4	8.5	9.2	8.5	8.5	8.5	8.5	85.3	8.53	0.23		
μ	8.88	8.38	8.38	8.42	8.50	9.40	8.62	8.52	8.44	8.44	85.98				
σ	0.64	0.28	0.28	0.30	0.26	1.58	0.38	0.26	0.30	0.30	3.83				

Run	Iteration											Aggregates		
	0	1	2	3	4	5	6	7	8	9	10	Σ	μ	σ
1	10.7	4.0	5.6	5.8	5.9	6.0	5.8	5.8	5.8	6.3	5.0	66.7	5.60	0.62
2	10.7	3.2	5.2	5.5	5.4	5.5	5.5	5.5	5.4	6.3	5.6	63.8	5.31	0.75
3	10.3	3.1	5.0	5.2	5.2	5.1	5.1	5.2	5.2	6.0	4.0	59.4	4.91	0.76
4	10.7	3.2	5.1	5.2	5.5	5.4	5.3	5.3	5.3	5.7	5.1	61.8	5.11	0.66
5	10.3	4.1	5.8	6.1	6.1	6.1	6.1	6.1	6.1	6.6	4.9	68.3	5.80	0.70
μ	10.43	3.52	5.34	5.56	5.62	5.62	5.56	5.58	5.56	6.18	4.92	64.00		
σ	0.27	0.44	0.31	0.35	0.33	0.38	0.36	0.33	0.34	0.31	0.52	3.22		

(b) Schätzle algorithm

(c) BRS algorithm executed with GSM Kaushik

Table 11: Detailed Results (minutes) on BSBM1B for 10-bisimulation.