

# Survey and cross-benchmark comparison of remaining time prediction methods in business process monitoring

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Predictive business process monitoring methods exploit historical process execution logs to generate predictions about running instances (called cases) of a business process, such as the prediction of the outcome, next activity or remaining cycle time of a given process case. These insights could be used to support operational managers in taking remedial actions as business processes unfold, e.g., shifting resources from one case onto another to ensure this latter is completed on time. A number of methods to tackle the remaining cycle time prediction problem have been proposed in the literature. However, due to differences in their experimental setup, choice of datasets, evaluation measures and baselines, the relative merits of each method remain unclear. This article presents a systematic literature review and taxonomy of methods for remaining time prediction in the context of business processes, as well as a cross-benchmark comparison of 16 such methods based on 17 real-life datasets originating from different industry domains.

CCS Concepts: • **Applied computing** → **Business process monitoring**;

Additional Key Words and Phrases: business process, predictive monitoring, process performance indicator, process mining, machine learning

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## 1 INTRODUCTION

A business process is a collection of events, activities, and decision points involving a number of actors and objects, which collectively lead to an outcome that is of value to a customer [15]. A typical example of a business process is an order-to-cash process: a process that starts when a purchase order is received and ends when the requested product(s) or service(s) have been delivered and the corresponding payment has been confirmed. An individual execution of a business process is called a *case*. In an order-to-cash process, each purchase order gives rise to a case.

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Business process monitoring is concerned with the analysis of events produced during the execution of a business process in order to assess the fulfillment of compliance requirements and performance objectives [15]. Process monitoring can take place offline, via periodically generated reports, or online, via dashboards displaying the performance of ongoing cases of a process in terms of performance indicators such as cycle time, resource utilization, and defect rate [16].

*Predictive business process monitoring* refers to a family of online process monitoring techniques that seek to predict the future state or properties of ongoing cases of a process based on models extracted from historical (completed) cases recorded in *event logs*. A recent survey of this field [28] highlighted the existence of a wide range of methods to predict compliance violations [6], undesirable outcomes [26, 27], the next activity or the remaining sequence of activities of a case [17, 58], or quantitative process performance indicators such as the remaining cycle time of a case [41, 42, 57, 59]. These predictions can be used to alert process workers to problematic cases or to support resource allocation decisions. For example, if the cycle time of a case is expected to exceed a given threshold stipulated by a service-level agreement (e.g., 10 working days), this prediction can be used to inform the customer of the expected delay in advance or to allocate a dedicated resource to this case in order to put it back on track towards timely completion.

Unfortunately, there is no unified approach to evaluate existing business process monitoring methods, as different authors have used different datasets, experimental settings, evaluation measures, and baselines to assess their proposed methods. This is the case in particular of methods for predicting the (remaining) cycle time of cases, where there is a wide range of disparate methods that have not been compared to each other in a uniform setting.

This paper aims to address this gap by: (i) performing a systematic literature review of predictive process monitoring methods for time-related properties; (ii) providing a taxonomy of existing methods; and (iii) performing a comparative experimental evaluation of 16 representative methods using a benchmark covering 17 real-world event logs. The contributions of this study are a categorized collection of outcome-oriented predictive process monitoring methods, a ranking of the relative accuracy of existing methods leading to recommendations for selecting a method in a given setting, and a benchmark designed to enable researchers to empirically compare new methods against existing ones in a unified setting. The benchmark is provided as an open-source framework that allows researchers to run the entire benchmark with minimal effort, and to configure and extend it with additional methods and datasets.

The above contributions significantly refine and extend those of a previous survey of predictive process monitoring methods [28]. This latter survey provides a broad taxonomy of predictive monitoring methods in terms of the target of the prediction (e.g., compliance violation, next event, remaining cycle time), the methods employed (generative models versus regression or classification models), and the datasets and evaluation measures employed. The present survey focuses on methods for predicting time-related properties (in particular, cycle time) but goes deeper in the taxonomy by considering differences in pre-processing and feature encoding approaches, and by providing an empirical comparison of the relative performance of existing methods in this field.

The rest of the paper is organized as follows. Section 2 summarizes some basic concepts in the area of predictive process monitoring. Section 3 describes the search and selection of relevant studies. Section 4 surveys the selected studies and provides a taxonomy to classify them, while Section 5 justifies the selection of methods to be evaluated in our benchmark. Section 6 reports on the benchmark of the selected studies while Section 7 identifies threats to validity. Finally, Section 8 summarizes the findings and identifies directions for future work.

## 2 BACKGROUND

Predictive process monitoring is a multi-disciplinary area that draws concepts from process mining on the one side, and machine learning on the other side. In this section, we introduce concepts from these two disciplines that we will use later in this paper.

### 2.1 Process mining

Business processes such as an order-to-cash process at a seller or a claims handling process at an insurance company are generally supported by information systems that record data about each individual execution of a process. For example, execution data of an order-to-cash process may be recorded in an enterprise resource planning (ERP) system, while execution data of a claims handling process would be recorded in a claims management system.

Process mining [55] is a research area across business process management and data science that is concerned with deriving useful insights from process execution data. Process mining techniques can support various phases of the business process management (BPM) lifecycle, such as process discovery, process analysis and process monitoring [55]. In particular, process monitoring is the last phase of the BPM lifecycle, and aims to support decision making at runtime, i.e., for the ongoing *cases* (e.g., orders, claims) of a business process. For example, in the context of a claims handling process, this means offering decision support to a claims manager in order to ensure that all open claims are completed on-time, within the service-level agreement established with the claimant. Decision support may be offered in the form of descriptive, predictive and prescriptive analytics on the performance of the running cases. For example, a descriptive analytic may be the duration of a case so far, a predictive analytic may be the predicted overall duration of the case until this completes, while a prescriptive analytic may be an intervention on a case to ensure this completes on-time. In this paper, we focus on *predictive* process monitoring.

The starting point of predictive process monitoring is the execution data of a given business process. This data may be available in the form of an *event log* recording historical data, or an *event stream*, recording live data. For example, an event log may record data related to all completed cases of a process (e.g., all claims completed in the last year), while an event stream may record data related to all open claims, as these cases unfold. In the following, we discuss the ingredients of event logs and event streams (here called *prefix logs*).

An event log consists of cases, each capturing a particular instance of the business process (e.g., a given order or claim). Each case consists of a number of *events* where each event represents the execution of a particular activity in the process. Each event has a range of attributes of which three are mandatory: i) the *case identifier* specifying which case generated this event, ii) the *event class* (or *activity name*) indicating which activity the event refers to, and iii) the *timestamp* indicating when the event occurred. An event may refer to the start or completion of an activity, and so would the event timestamp. In this paper, we refer to the *completion* timestamp and assume each event log has only completion timestamps, unless otherwise noted. Besides case identifier, event class and timestamp, an event may carry additional attributes in its payload, such as the name of the resource executing the activity, or the data used as input/output.

As a running example, let us consider a patient treatment process in a hospital, where each case refers to a particular patient receiving treatment. The patient's date of birth and gender can be considered case-level attributes, or simply *case attributes*. These attributes belong to the case and are therefore shared by all events relating to that case. In addition, each event may have its specific *event attributes* contained in the event payload. For instance, the name of the nurse responsible for taking the blood test may be recorded as an attribute of an event referring to activity "Perform blood test". In other words, case attributes are invariant, i.e., their values do not change throughout

the lifetime of a case, while event attributes are dynamic as they change from an event to the other. Table 1 provides an example event log for our patient treatment process.

Table 1. Extract of an event log of a patient treatment process

Case ID	Case attributes		Event attributes			
	D.O.B.	Sex	Activity	Completion Time	Resource	Cost
1	20/02/1982	M	Registration	1/1/2017 9:13:00	John	15
1	20/02/1982	M	Antibiotics	1/1/2017 9:14:20	Mark	25
1	20/02/1982	M	Triage	1/1/2017 9:16:00	Mary	10
1	20/02/1982	M	Release	1/1/2017 9:18:05	Kate	20
1	20/02/1982	M	Return	1/1/2017 9:18:50	John	20
1	20/02/1982	M	Blood test	1/1/2017 9:19:00	Kate	15
2	03/08/1967	F	Registration	2/1/2017 16:55:00	John	25
2	03/08/1967	F	Triage	2/1/2017 17:00:00	Mary	25
2	03/08/1967	F	Antibiotics	3/1/2017 9:00:00	Mark	10
2	03/08/1967	F	Release	3/1/2017 9:01:50	Kate	15

The activity name of the first event in case 1 is *Registration*, which occurred on 1/1/2017 at 9:13 AM. This is the time when the first patient was registered in the hospital information system this event log has been extracted from. The additional event attributes show that the cost of the activity was 15 units and the activity was performed by *John*. These latter two are event attributes. The events in each case also carry two case attributes: the patient's date of birth and gender.

Let us provide formal definitions for the above-introduced terms.

*Definition 2.1 (Event).* An *event* is a tuple  $\langle a; c; t; \langle d_1; \dots; d_m \rangle \rangle$  where  $a$  is the activity name,  $c$  is the case identifier,  $t$  is the timestamp and  $\langle d_1; \dots; d_m \rangle$  (where  $m \geq 0$ ) are event attributes names and the corresponding values assumed by them.

Let  $E$  be the event universe, i.e., the set of all possible event classes, and  $T$  the time domain. Then there is a function  $\tau: E \rightarrow T$  that assigns timestamps to events.

The sequence of events generated by a given case forms a *trace*. Formally,

*Definition 2.2 (Trace).* A *trace* is a non-empty sequence  $\langle e_1; \dots; e_n \rangle$  of events such that  $\forall i \in \{1; \dots; n\}; e_i \in E$  and  $\forall i; j \in \{1; \dots; n\}; e_i.c = e_j.c$ . In other words, all events in the trace refer to the same case.

A set of *completed traces* (i.e., traces recording the execution of completed cases) comprises an *event log*.

*Definition 2.3 (Event log).* An event log  $L$  is a set of completed traces, i.e.,  $L = \{t_i : i \in \{1; \dots; K\}\}$ , where  $S$  is the universe of all possible traces and  $K$  is the number of traces in the event log.

Event and case attributes are typically divided into categorical (qualitative) and numeric (quantitative) data type [49]. With respect to our running example, categorical attributes are *Sex*, *Activity* and *Resource*, while the only numeric attribute is *Cost*. Additionally, date of birth and completion timestamp can be used to extract more meaningful categorical and numeric attributes, such as the age of a patient, day of the week, time elapsed since the previous activity and so on.

Importantly, each data type requires different preprocessing to be used in a predictive model. Numeric attributes can either be represented as such or transformed to a different scale, while for

categorical attributes, one-hot encoding is typically applied. One-hot encoding represents each value of the categorical attribute with a binary vector with the  $i$ -th component set to one, and the rest set to zero. In other words, it provides a 1-to- $N$  mapping where  $N$  is the number of possible distinct values (*levels*), or cardinality of an attribute [31].

In predictive process monitoring we aim to make predictions for traces of incomplete cases (as stored in an event stream), rather than for traces of completed cases (as stored in an event log). To capture such partial traces, we define a function that returns the first  $k$  events of a trace of a (completed) case.

*Definition 2.4 (Prefix function).* Given a trace  $\tau = \langle e_1; \dots; e_n \rangle$  and a positive integer  $k \leq n$ ,  $hd^{k, \tau} = \langle e_1; \dots; e_k \rangle$ .

For example, for a sequence  $\tau = \langle a; b; c; d; e \rangle$ ,  $hd^{2, \tau} = \langle a; b \rangle$ .

The application of a prefix function will result in a *prefix log*, where each possible prefix of an event log becomes a trace. A prefix log captures formally the notion of an event stream, i.e., a container of partial traces.

*Definition 2.5 (Prefix log).* Given an event log  $L$ , its prefix log  $L^{\#}$  is the event log that contains all prefixes of  $L$ , i.e.,  $L^{\#} = \{hd^{k, \tau} : \tau \in L, 1 \leq k \leq |\tau|\}$ .

For example, a complete trace consisting of three events would correspond to three partial traces in the prefix log – the partial trace after executing the first, the second, and the third event. For event logs where the case length may vary significantly from case to case, instead of including every prefix of a trace, it is common to include only prefixes with a specific length in the prefix log during the training procedure. This can be done by retaining all prefixes of up to a certain length [26, 52]. An alternative approach is to only include prefixes  $hd^{k, \tau}$  such that  $hd^{k, \tau} \equiv 1 \pmod{\gamma}$  where  $\gamma$  is a gap size. This approach has been explored in [14].

## 2.2 Machine learning

At the core of every predictive process monitoring system are prediction models built for specific prediction goals using an event log  $L$  of complete cases. At runtime, these prediction models are applied to prefixes  $hd^{k, \tau}$  of ongoing cases read from an event stream, to make predictions about the future performance of such cases. Prediction models are typically built using machine learning algorithms.

Machine learning is a research area of computer science concerned with the discovery of models, patterns, and other regularities in data [32]. A machine learning system is characterized by a learning algorithm used on training data. The algorithm defines a process of learning from information extracted, usually as *feature vectors*, from the training data. In this work, we will deal with *supervised* learning, meaning that training data is *labeled*. Labeled training data is represented in the following form:

$$D = \{ \langle \mathbf{x}_1; \dots; \mathbf{x}_n \rangle : n \in \mathbb{N} \} \quad (1)$$

where  $\mathbf{x}_i \in \mathbb{X}$  are  $m$ -dimensional feature vectors ( $m \in \mathbb{N}$ ) and  $y_i \in \mathbb{Y}$  are the corresponding labels, i.e., values of the target variable.

Feature vectors extracted from the labeled training data are used to fit a predictive model that would assign labels on new data given labeled training data while minimizing error and model complexity. In other words, a model generalizes the pattern identified in the training data, providing a mapping  $\mathbb{X} \rightarrow \mathbb{Y}$ . The labels can be either continuous, e.g., cycle time of an activity, or discrete,

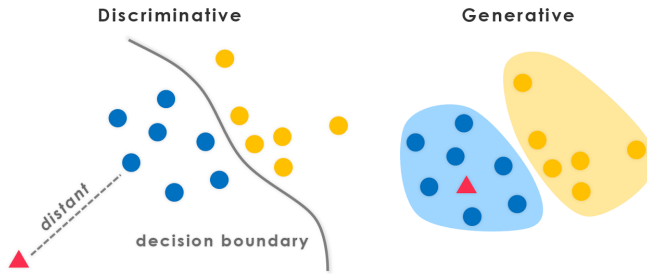


Fig. 1. Discriminative and generative models [34].

e.g., loan grade. In the former case, the model is referred to as a *regression* model; while in the latter case we are talking about a *classification* model.

From a probabilistic perspective, the machine learning objective is to infer a conditional distribution  $P^1Y|X^0$ . A standard approach to tackle this problem is to represent the conditional distribution with a parametric model, and then to obtain the parameters using a training set containing  $\{x_{n_i}, y_{n_i}\}$  pairs of input feature vectors with corresponding target output vectors. The resulting conditional distribution can be used to make predictions of  $y$  for new values of  $x$ . This is called a *discriminative* approach, since the conditional distribution discriminates between the different values of  $y$  [25].

Another approach is to calculate the joint distribution  $P^1X; Y^0$ , expressed as a parametric model, and then apply it to find the conditional distribution  $P^1Y|X^0$  to make predictions of  $y$  for new values of  $x$ . This is commonly known as a *generative* approach since by sampling from the joint distribution one can generate synthetic examples of the feature vector  $x$  [25].

To sum up, *discriminative* approaches try to define a (hard or soft) decision boundary that divides the feature space into areas containing feature vectors belonging to the same class (see Figure 1). In contrast, *generative* approaches first model the probability distributions for each class and then label a new instance as a member of a class whose model is most likely to have generated the instance [25].

In machine learning literature, various discriminative and generative algorithms have been proposed to address classification and regression tasks. Decision trees are a common choice in predictive business process monitoring, owing to their simplicity and interpretability [12, 21]. To increase the prediction accuracy, decision trees are often combined into *ensembles*. Ensemble methods train multiple predictors for the same task and combine them together for the final prediction. Figure 2 illustrates the general idea behind ensemble learning. Multiple base learners, or “weak” learners,  $f_k$  are fitted on the training data or its subset. Their predictions are then combined into one ensemble learner  $f_\Sigma$  by applying some aggregation function  $\sigma$ . A popular aggregation strategy is gradient boosting, where predictions from base learners are combined by following a gradient learning strategy [22].

### 2.3 Predictive process monitoring

Given an event log of complete cases of a business process, and a prefix case of this process as obtained from an event stream, we want to predict a performance measure of such prefix case in the future. For example, we may want to predict the time of this case until completion (or remaining time) or its outcome at completion. This task is sketched in Figure 3. A *prediction point* is the point in time where the prediction takes place. A *predicted point* is a point in the future where the performance measure has the predicted value. A prediction is thus based on the predictor’s knowledge of the history of the process until the prediction point as well as knowledge of the

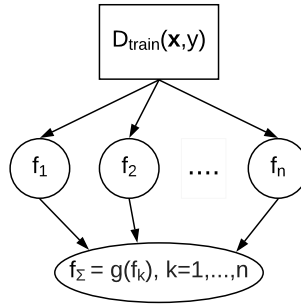


Fig. 2. Ensemble learning architecture.

future until the predicted point. The former is warranted by the predictor’s *memory* while the latter is based on the predictor’s *forecast*, i.e., predicting the future based on trend and seasonal pattern analysis. Finally, the prediction is performed based on a *prediction algorithm*.

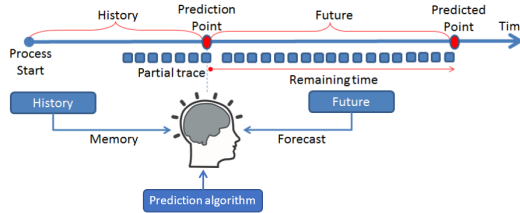


Fig. 3. Overview of predictive process monitoring.

Since in real-life business processes the amount of uncertainty increases over time (see “cone of uncertainty” [47]), the prediction task becomes more difficult and generally less accurate. As such, predictions are typically made up to a specific point of time in the future, i.e., the time horizon  $h$ . The choice of  $h$  depends on how fast the process evolves and on the prediction goal.

### 3 SEARCH METHODOLOGY

In order to retrieve and select studies for our survey and benchmark, we conducted a *Systematic Literature Review* (SLR) according to the approach described in [24]. We started by specifying the research questions. Next, guided by these goals, we developed relevant search strings for querying a database of academic papers. We applied inclusion and exclusion criteria to the retrieved studies in order to filter out irrelevant ones, and last, we divided all relevant studies into primary and subsumed ones based on their contribution.

#### 3.1 Research questions

The purpose of this survey is to define a taxonomy of methods for predictive monitoring of remaining time of business processes. The decision to focus on remaining time is to have a well-delimited and manageable scope, given the richness of the literature in the broader field of predictive process monitoring, and the fact that other predictive process monitoring tasks might rely on different techniques and evaluation measures.

In line with the selected scope, in this benchmark, we aim to answer the following research questions:

RQ1 What methods exist for predictive monitoring of remaining time of business processes?



RQ2 How to classify methods for predictive monitoring of remaining time?

RQ3 What type of data has been used to evaluate these methods, and from which application domains?

RQ4 What tools are available to support these methods?

RQ5 What is the relative performance of these methods?

RQ1 is the core research question, which aims to identify existing methods to perform predictive monitoring of remaining time. With RQ2, we aim to identify a set of classification criteria on the basis of input data required (e.g., input log) and the underlying predictive algorithms. RQ3 explores what tool support the different methods have, while RQ4 investigates how the methods have been evaluated and in which application domains. Finally, with RQ5, we aim to cross-benchmark existing methods using a set of real-life logs.

### 3.2 Study retrieval

Existing literature in predictive business process monitoring was searched for using Google Scholar, a well-known electronic literature database, as it covers all relevant databases such as ACM Digital Library and IEEE Xplore, and also allows searching within the full text of a paper.

Our search methodology is based on the one proposed in [28], with few variations. Firstly, we collected publications using more specific search phrases, namely “predictive process monitoring”, “predictive business process monitoring”, “predict (the) remaining time”, “remaining time prediction” and “predict (the) remaining \* time”. The latter is included since some authors refer to the prediction of the remaining *processing* time, while others may call it remaining *execution* time and so on. We retrieved all studies that contained at least one of the above phrases in the title or in the full text of the paper. The search was conducted in March 2018 to include all papers published between 2005 and 2017.

The initial search returned **670** unique results which is about 3 times more than the ones found in [28], owing to the differences in search methodologies (Table 2). Figure 4 shows how the studies are distributed over time. We can see that the interest in the topic of predictive process monitoring grows over time with a sharp increase over the past few years.

Table 2. Comparison of our search methodology with [28]

	Method in [28]	Our method
Keywords	1. “predictive monitoring” AND “business process” 2. “business process” AND “prediction”	1. “predictive process” monitoring 2. “predictive business process monitoring” 3. “predict (the) remaining time” 4. “remaining time prediction” 5. “predict (the) remaining * time”
Search scope	Title, abstract, keywords	Title, full text
Min number of citations	5 (except 2016 papers)	5 (except 2017 papers)
Years covered	2010-2016	2005-2017
Papers found after filtering	41	53
Snowballing applied	No	Yes, one-hop

In order to consider only relevant studies, we designed a range of exclusion criteria to assess the relevance of the studies. First, we excluded those papers not related to the process mining



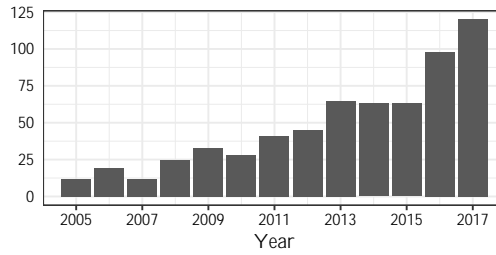


Fig. 4. Number of published predictive process monitoring studies over time.

field, written in languages other than English or papers with inaccessible full text. Additionally, to contain the reviewing effort, we have only included papers that have been cited at least five times. An exception has been made for papers published in 2017 – as many of them have not had a chance to accumulate the necessary number of citations, we required only one citation for them. Thus, after the first round of filtering, a total of **53** publications were considered for further evaluation.

Since different authors might use different terms, not captured by our search phrases, to refer to the prediction target in question, we decided to also include all studies that cite the previously discovered 53 publications (“snowballing”). Applying the same exclusion criteria, we ended up with **83** more studies. Due to the close-knit nature of the process mining community, there is a considerable overlap between these 83 studies and the 53 studies that had been retrieved during the first search stage. Accordingly, a total of **110** publications were finally considered on the scope of our review. All papers retrieved at each search step can be found at <https://goo.gl/kg6xZ1>.

The remaining 110 papers were further assessed with respect to exclusion criteria:

- EX1 The study does not actually propose a predictive process monitoring *method*. With this criterion, we excluded position papers, as well as studies that, after a more thorough examination, turned out to be focusing on some research question other than predictive process monitoring. Furthermore, here we excluded survey papers and implementation papers that employ existing predictive methods rather than propose new ones.
- EX2 The study does not concern remaining time predictions. Common examples of other prediction targets that are considered irrelevant to this study are failure and error prediction, as well as the next activity prediction. At the same time, prediction targets such as case completion time prediction and case duration prediction are inherently related to remaining time and therefore were also considered in our work. Additionally, this criterion does not eliminate studies that address the problem of predicting deadline violations in a boolean manner by setting a threshold on the predicted remaining time rather than by a direct classification.
- EX3 The study does not take an *event log* as input. In this way, we exclude methods that do not utilize at least the following essential parts of an event log: the case identifier, the timestamp and the event classes. For instance, we excluded methods that take as input numerical time series without considering the heterogeneity in the control flow. In particular, this is the case in manufacturing processes which are of linear nature (a process chain). The reason for excluding such studies is that the challenges when predicting for a set of cases of heterogeneous length are different from those when predicting for linear processes. While methods designed for heterogeneous processes are usually applicable to those of linear nature, it is not so vice versa. Moreover, the linear nature of a process makes it possible to apply other, more standard methods that may achieve better performance.

The application of the exclusion criteria resulted 25 relevant studies which are described in detail in the following section.

#### 4 ANALYSIS AND CLASSIFICATION OF METHODS

Driven by the research questions defined in Section 3.1, we identified the following dimensions to categorize and describe the relevant studies.

Type of input data RQ2

Awareness of the underlying business process RQ2

Family of algorithms RQ2

Type of evaluation data (real-life or artificial log) and application domain (e.g., insurance, banking, healthcare) RQ3.

Type of implementation (standalone or plug-in, and tool accessibility) RQ4

This information, summarized in Table 3, allows us to answer the first research question. In the remainder of this section, we provide a brief summary of each main study and then proceed with surveying the studies along the above classification dimensions.

##### 4.1 Overview

In the pioneering predictive monitoring approach [59], the authors aim to predict the remaining cycle time of a case by using non-parametric regression and leveraging activity duration and occurrence as well as other case-related data. Van der Aalst et al. [54] propose a set of approaches to build transition systems based on a given abstraction of the events in the event log and annotate them with time information extracted from the logs. In particular, information about elapsed, sojourn, and remaining time is reported for each state of the transition system. This information is then used to make predictions for the completion time of an ongoing case.

Folino et al. [19, 20] proposed an extension of the approach presented [57]. These studies combine an annotated transition system with a context-driven predictive clustering approach. Predictive clustering is based on the idea that different scenarios can be characterized by different predictors. Furthermore, contextual information is utilized to make more accurate predictions, in addition to control-flow [19] or control-flow and resources [20]. However, these transition system-based approaches require users to choose the log abstraction function. To address this limitation, Bevacqua et al. [3] and Cesario et al. [8] replace transition system predictors with standard regression algorithms. Additionally, in [6], the clustering component is further improved in order to address accuracy and scalability issues.

The approach presented in [57] is also refined by Polato et al. in [39] and [40] who apply machine learning techniques to annotate the transition systems in order to predict the remaining time of an ongoing case. Namely, in [39], the transition systems are annotated with machine learning models, such as Naïve Bayes and Support Vector Regression models. Further extension of this approach is presented in [40], where sojourn times are implicitly included in the Support Vector Regression model, and the evaluation is performed on both stationary and dynamic processes that might contain seasonal drifts.

Another technique based on the extraction of explicit models, in the form of sequence trees, is proposed by Ceci et al. in [7] who aim to predict the completion time and the next activity of an ongoing case. As with the predictive clustering approach, the sequence trees enable clustering of traces with similar sequences of activities and building a predictor for each node of the sequence tree by using control-flow and event attributes.

Rogge-Solti and Weske [1] leverage generally distributed transitions stochastic Petri nets (GDT-SPN) to predict the remaining time of a case. The approach takes as inputs (i) a stochastic process

Table 3. Overview of the 25 relevant studies resulting from the search (ordered by year and author).

Study	Year	Input data	Process-aware?	Algorithm	Domain	Implementation
van Dongen et al. [59]	2008	event log data	No	regression	public administration	ProM 5
van der Aalst et al. [57]	2011	event log data	Yes	transition system	public administration	ProM 5
Folino et al. [19]	2012	event log data	No	clustering	logistics	ProM
Pika et al. [37]	2012	contextual information event log data	No	stat analysis	nancial	ProM
van der Spoel et al. [58]	2012	event log data	Yes	process graph regression	healthcare	n/a
Bevacqua et al. [3]	2013	event log data	No	clustering regression	logistics	ProM
Bolt and Sepúlveda [4]	2013	event log data	Yes	stat analysis	telecom simulated	n/a
Folino et al. [20]	2013	event log data	No	clustering	customer service	n/a
Pika et al. [38]	2013	contextual information event log data	No	stat analysis	insurance	ProM
Rogge-Solti and Weske [41]	2013	event log data	Yes	stoch Petri net	logistics simulated	ProM
Ceci et al. [7]	2014	event log data	Yes	sequence trees regression	customer service work ow management	ProM
Folino et al. [21]	2014	event log data	No	clustering regression	logistics software development	n/a
de Leoni et al. [11]	2014	contextual information event log data	No	regression	no validation	ProM
Polato et al. [39]	2014	contextual information event log data	Yes	transition system regression	public administration	ProM
Senderovich et al. [44]	2014	event log data	Yes	classification queueing theory transition system	nancial	n/a
Metzger et al. [30]	2015	event log data	Yes	neural network constraint satisfaction	logistics	n/a
Rogge-Solti and Weske [42]	2015	process model event log data	Yes	QoS aggregation stoch Petri net	nancial logistics	ProM
Senderovich et al. [45]	2015	event log data	Yes	queueing theory transition system	nancial telecom	n/a
Cesario et al. [8]	2016	event log data	Yes	clustering regression	logistics	n/a
de Leoni et al. [12]	2016	event log data	No	regression	no validation	ProM
Polato et al. [40]	2018	contextual information event log data	Yes	transition system regression	public administration customer service	ProM
Senderovich et al. [43]	2017	event log data	No	classification regression	nancial healthcare manufacturing	standalone
Tax et al. [50]	2017	contextual information event log data	No	neural network	customer service public administration	standalone
Navarin et al. [33]	2017	event log data	No	neural network	nancial customer service	standalone
Verenich et al. [60]	2017	event log data process model	Yes	regression classification ow analysis	nancial customer service	standalone

model, which may be available in advance or derived from historical data, (ii) an ongoing trace, and (iii) the current time in order to make the predictions. The authors extended their approach in [42] to take into account the elapsed time since the last event in order to make more accurate time predictions and to estimate the probability of missing a deadline.

Senderovich et al. [43] propose and evaluate different strategies for extracting inter-case features for predicting the completion time of an ongoing trace. Such inter-case features consider not only the information related to the ongoing case in question, but also the other, concurrent cases.

De Leoni et al. [1, 12] propose a general framework to predict various characteristics of running instances, including the remaining time, based on correlations with other characteristics and using decision and regression trees.

Tax et al. [50] propose deep learning models based on long short term memory (LSTM) neural networks to predict the next activity to be executed and its timestamp. By iteratively repeating this process until the case is predicted to be finished, the approach also allows for the remaining time prediction. An alternative to this incremental approach is proposed by Navarin et al. [31], who train an LSTM model to directly predict the remaining time as a single scalar value.

Metzger et al. [30] propose three approaches to predict time-based constraint violations, namely machine learning, constraint satisfaction and QoS aggregation. The authors show that all the approaches achieve satisfactory results and then propose to combine them. Results on a real-life case study show that combining these techniques improves the prediction accuracy [41] and [45]. Senderovich et al. apply queuing theory to predict possible delays in business process executions. The proposed approaches refine traditional techniques based on transition systems to take into account queuing effects. The problem of predicting deadline violations in business processes has also addressed by Pika et al. [17, 38] by identifying process risk indicators that cause the possibility of a delay.

## 4.2 Input data

As stipulated by EX3 criterion, all surveyed proposals take as input an event log. Such a log contains at least a case identifier, an activity and a timestamp. In addition, many techniques leverage case and event attributes to make more accurate predictions. For example, in the pioneering predictive monitoring approach described in [59], the authors predict the remaining processing time of a trace using activity durations, their frequencies and various case attributes, such as the case priority. Many other approaches, e.g., [14], [40], [43] make use not only of case attributes but also of event attributes, while applying one or more kinds of sequence encoding. Furthermore, some approaches, e.g., [19] and [43], exploit contextual information, such as workload indicators, to take into account inter-case dependencies due to resource contention and data sharing. Finally, a group of works, e.g., [30] and [60] also leverage a process model in order to replay ongoing process cases on it. Such works treat remaining time as a cumulative indicator composed of cycle times of elementary process components.

## 4.3 Process awareness

Existing techniques can be categorized according to their process-awareness, i.e., whether or not the methodology exploits an explicit representation of a process model to make predictions. As can be seen from Table 3, nearly a half of the techniques are process-aware. Most of them construct a transition system from an event log using set, bag (multiset) or sequence abstractions of observed events. State transition systems are based on the idea that the process is composed of a set of consistent states and the movement between them [50]. Thus, a process behavior can be predicted if we know its current and future states.

Bolt and Sepúlveda [1] exploit query catalogs to store the information about the process behavior. Such catalogs are groups of partial traces (annotated with additional information about each partial trace) that have occurred in an event log, and are then used to estimate the remaining time of new executions of the process.

Also queuing models can be used for prediction because if a process follows a queuing context and queuing measures (e.g., arrival rate, departure rate) can be accurately estimated and at the process actual execution, the movement of a queuing item can be reliably predicted. Queuing theory and regression-based techniques are combined for delay prediction in [44, 45].

Furthermore, some process-aware approaches rely on stochastic Petri nets [41, 42] and process models in BPMN notation [60].

#### 4.4 Family of algorithms

Non-process aware approaches typically rely on machine learning algorithms to make predictions. In particular, these algorithms take as input labeled training data in the form of feature vectors and the corresponding labels. In case of remaining time predictions, these labels are continuous. As such, various regression methods can be utilized, such as regression trees [2] or ensemble of trees, i.e., random forest [58] and XGBoost [43].

An emerging family of algorithms for predictive monitoring are artificial neural networks. They consist of one layer of input units, one layer of output units, and multiple layers in-between which are referred to as hidden units. While traditional machine learning methods heavily depend on the choice of features on which they are applied, neural networks are capable of translating the data into a compact intermediate representation to aid a hand-crafted feature engineering process. A feedforward network has been applied [30] to predict deadline violations. More sophisticated architectures based on recurrent neural networks were explored in [50] and [33].

Other approaches apply trace clustering to group similar traces to fit a predictive model for each cluster. Then for any new running process case, predictions are made by using the predictor of the cluster it is estimated to belong to. Such approach is employed e.g., in [19] and [21].

Another range of proposals utilizes statistical methods without training an explicit machine learning model. For example, Pika et al. [37, 38] make predictions about time-related process risks by identifying and leveraging process risk indicators (e.g., abnormal activity execution time or multiple activity repetition) by applying statistical methods to event logs. The indicators are then combined by means of a prediction function, which allows for highlighting the possibility of transgressing deadlines. Conversely, Bolt and Sepúlveda [4] calculate remaining time based on the average time in the catalog which the partial trace belongs to, without taking into account distributions and confidence intervals.

Rogge-Solti and Weske [11] mine a stochastic Petri net from the event log to predict the remaining time of a case using arbitrary firing delays. The remaining time is evaluated based on the fact that there is an initial time distribution for a case to be executed. As inputs, the method receives the Petri net, the ongoing trace of the process instance up to current time, the current time and the number of simulation iterations. The algorithm returns the average of simulated completion times of each iteration. This approach is extended [42] to exploit the elapsed time since the last observed event to make more accurate predictions.

Finally, Verenich et al. [60] propose a hybrid approach that relies on classification methods to predict routing probabilities for each decision point in a process model, regression methods to predict cycle times of future events, and flow analysis methods to calculate the total remaining time. A conceptually similar approach is proposed by Polato et al. [40] who build a transition system from an event log and enrich it with classification and regression models. Naive Bayes classifiers are used to estimate the probability of transition from one state to the other, while support vector regressors are used to predict the remaining time from the next state.

#### 4.5 Evaluation data and domains

As reported in Table 3, most of the surveyed methods have been validated on at least one real-life event log. Some studies were additionally validated on simulated (synthetic) logs.

Importantly, many of the real-life logs are publicly available from the TU Center for Research Data<sup>1</sup> Among the methods that employ real-life logs, we observed a growing trend to use publicly available logs, as opposed to private logs which hinder the reproducibility of the results due to not being accessible.

Concerning the application domains of the real-life logs, we noticed that most of them pertain to banking (8 studies), logistics and customer service (6 studies each) and public administration (5 studies) processes.

#### 4.6 Implementation

Providing publicly available implementation and experiment data greatly facilitates reproducibility of research experiments, as well as to enable future researchers to build on past work. To this end, we found that nearly a half of the methods provide implementation as a plug-in for the ProM framework<sup>2</sup> The reason behind the popularity of ProM can be explained by its open-source and portable framework, which allows researchers to easily develop and test new algorithms. Also, ProM provides an Operational Support (OS) environment<sup>3</sup> that allows it to interact with external workflow management systems at runtime. In this way, process mining can also be performed in an online setting. Another 4 methods have a standalone implementation in Python<sup>3</sup>. Finally, 8 methods do not provide a publicly available implementation.

#### 4.7 Predictive monitoring workflow

As indicated in Table 3, most predictive monitoring methods make use of machine learning algorithms based on regression, classification or neural networks. Such methods typically proceed in two phases: offline, to train a prediction model based on historical cases, and online, to make predictions on running process cases (Figure 5) [51].

In the offline phase, given an event log, case prefixes are extracted and filtered, e.g., to retain only prefixes up to a certain length, to create a prefix log (cf. Section 2). Next, similar prefixes are grouped into homogeneous buckets, e.g., based on process states or similarities among prefixes and prefixes from each bucket are encoded into feature vectors. Then feature vectors from each bucket are used to fit a machine learning model.

In the online phase, the actual predictions for running cases are made, by reusing the buckets, encoders and predictive models built in the offline phase. Specifically, given a running case and a set of buckets of historical prefixes, the correct bucket is first determined. Next, this information is used to encode the features of the running case. In the last step, a prediction is extracted from the encoded case using the pre-trained model corresponding to the determined bucket.

Similar observations can be made for non-machine learning-based methods. For example, in [51], first, a transition system is derived and annotated and then the actual predictions are calculated for running cases. In principle, this transition system akin to a predictive model can be mined in advance and used at runtime.

#### 4.8 Primary and subsumed (related) studies

Among the papers that successfully passed both the inclusion and exclusion criteria, we determined primary studies that constitute an original contribution for the purposes of our benchmark, and subsumed studies that are similar to one of the primary studies and do not provide a substantial contribution with respect to it.

<sup>1</sup>[https://data.4tu.nl/repository/collection:event\\_logs\\_real](https://data.4tu.nl/repository/collection:event_logs_real)

<sup>2</sup><http://promtools.org>

<sup>3</sup><https://www.python.org>

Fig. 5. Predictive process monitoring workflow.

Specifically, a study is considered subsumed if:

- there exists a more recent and/or more extensive version of the study from the same authors (e.g., a conference paper is subsumed by an extended journal version), or
- it does not propose a substantial improvement/modification over a method that is documented in an earlier paper by other authors, or
- the main contribution of the paper is a case study or a tool implementation, rather than the predictive process monitoring method itself, and the method is described and/or evaluated more extensively in a more recent study by other authors.

This procedure resulted in 10 primary and 14 subsumed studies, as reported in Table 4. Some studies are subsumed by several primary studies. For instance, Metzger *et al.* [34] propose a feedforward neural network which is subsumed by a more sophisticated recurrent neural network architecture explored in [33]. At the same time, [30] describes a QoS technique which is subsumed by the flow analysis technique presented in [60].

## 5 METHODOLOGICAL FRAMEWORK

Assessing all the methods that resulted from the search would be infeasible due to the heterogeneous nature of the inputs required and the outputs produced. As such, we decided to abstract the details that are not inherent to the methods and focus on their core differences.

### 5.1 Prefix bucketing

While some machine learning-based predictive process monitoring approaches train a single predictor on the whole event log, others employ a multiple predictor approach by dividing the prefix traces in the historical log into several buckets and training a separate predictor for each bucket.



Table 4. Primary and subsumed studies

Primary study	Subsumed studies
van der Aalst et al. [57]	van Dongen et al. [59], Bolt and Sepúlveda [4]
Folino et al. [19]	Folino et al. [20, 21]
Rogge-Solti and Weske [42]	Rogge-Solti and Weske [41]
Senderovich et al. [45]	Senderovich et al. [44]
Cesario et al. [8]	Bevacqua et al. [3]
de Leoni et al. [12]	Pika et al. [37, 38], de Leoni et al. [11]
Polato et al. [40]	van der Spoel et al. [58], Polato et al. [39], Ceci et al. [7]
Senderovich et al. [43]	van der Spoel et al. [58]
Tax et al. [50]	Metzger et al. [30]
Navarin et al. [33]	Verenich et al. [60], Metzger et al. [30]

To this end, Teinmaa et al [51] surveyed several bucketing methods out of which three have been utilized in the primary methods:

**Zero bucketing.** All pre x traces are considered to be in the same bucket. As such, a single predictor is t for all pre xes in the pre x log. This approach has been used in [2], [43] and [50].

**Pre x length bucketing.** Each bucket contains the pre xes of a speci c length. For example, the n-th bucket contains pre xes where at least n events have been performed. One classi er is built for each possible pre x length. This approach has been used in [60].

**Cluster bucketing.** Here, each bucket represents a cluster that results from applying a clustering algorithm on the encoded pre xes. One classi er is trained for each resulting cluster, considering only the historical pre xes that fall into that particular cluster. At runtime, the cluster of the running case is determined based on its similarity to each of the existing clusters and the corresponding classi er is applied. This approach has been used in [19] and [8]

**State bucketing.** It is used in process-aware approaches where some kind of process representation, e.g., in the form of a transition system, is derived and a predictor is trained for each state, or decision point. At runtime, the current state of the running case is determined, and the respective predictor is used to make a prediction for the running case. This approach has been used in [40].

## 5.2 Prefix encoding

In order to train a machine learning model, all pre xes in a given bucket need to be represented, or encoded as fixed-size feature vectors. Case attributes are static, and their number is fixed for a given process. Conversely, with each executed event, more information about the case becomes available. As such, the number of event attributes will increase over time. To address this issue, various sequence encoding techniques were proposed in related work summarized in [60] and refined in [51]. In the primary studies, the following encoding techniques can be found:

**Last state encoding.** In this encoding method, only event attributes of the last events are considered. Therefore, the size of the feature vector is proportional to the number of event attributes and is fixed throughout the execution of a case.  $n = 1$  is the most common choice used, e.g., in [40], although in principle higher values can also be used.

**Aggregation encoding.** In contrast to the last-state encoding, aggregation encoding considers all events in the pre x of a case, rather than considering only the last events. Each attribute

is encoded into one or multiple features using different aggregation functions depending on the datatype of the attribute. If an attribute is of numerical type (e.g., amount), we map this attribute into one feature by means of a numerical aggregation function such as sum, average, minimum, or maximum. On the other hand, if an attribute is categorical (e.g., level), we map this attribute into one feature for every value in the attribute's domain by applying the count aggregation function. For example, the possible values of the attribute risk level are  $\{A; B; C; D\}$ . We map this attribute to four features: risk level-A, risk level-B, risk level-C, risk level-D. For a given prefix of a case, risk level-A is the number of times that the attribute risk level has had a value equal to A among the events in the prefix, and similarly for risk level-B, risk level-C, etc. In other words, the resulting features correspond to the absolute frequency of each possible value in the attribute's domain within the current prefix of a case. Note that the activity attribute (i.e., the attribute corresponding to the type of activity referenced by an event) is a categorical attribute and hence it is encoded using the count function. This leads to one feature for each type of activity (e.g., one feature for Submit loan application, another for Check credit history, etc.). The value of each feature is the frequency of the type of activity in question, within a given prefix. This encoding is suggested for example in [12].

**Index-based encoding.** Here for each position in a prefix, we concatenate the event occurring in that position and the value of each event attribute in that position:  $v_1^k \dots v_n^k$ , where  $k$  is the total number of attributes of an event. This type of encoding is lossless, i.e., it is possible to recover the original prefix based on its feature vector. On the other hand, with longer prefixes, it significantly increases the dimensionality of the feature vectors and hinders the model training process. This approach has been used in [60].

**Tensor encoding.** A tensor is a generalization of vectors and matrices to potentially higher dimensions [48]. Unlike conventional machine learning algorithms, tensor-based models do not require input to be encoded in a two-dimensional form, where  $n$  is the number of training instances and  $d$  is the number of features. Conversely, they can take as input a three-dimensional tensor of shape  $t \times p \times d$ , where  $t$  is the number of events and  $d$  is the number of event attributes, or features derived from each event. In other words, each prefix is represented as a matrix where rows correspond to events and columns to features for a given event. The data for each event is encoded as-is. Case attributes are encoded as event attributes having the same value throughout the prefix. Hence, the encoding for LSTM is similar to the index-based encoding except for two differences: (i) case attributes are duplicated for each event, (ii) the feature vector is reshaped into a matrix.

To aid the explanation of the encoding types, Tables 5–7 provide examples of feature vectors derived from the event log in Table 1. Note that for the index-based encoding, each trace in the event log produces only one training sample per bucket, while the other encodings produce as many samples as many prefixes can be derived from the original trace, i.e.,  $|p|$  to

These three canonical encodings can serve as a basis for various modifications thereof. For example, de Leoni et al. [14] proposed the possibility of combining last state and aggregation encodings.

While the encoding techniques stipulate how to incorporate event attributes in a feature vector, the inclusion of case attributes and inter-case metrics, such as the number of currently open cases, is rather straightforward, as their number is fixed throughout the case lifetime.

While last state and aggregation encodings can be combined with any of the bucketing methods described in Section 5.1, index-based encoding is commonly used with prefix-length bucketing,

Table 5. Feature vectors created from the log in Table 1 using last state encoding.

Sex	Age	Activity_last	Time_last	Resource_last	Cost_last
M	37	Registration	0	John	15
M	37	Antibiotics	80	Mark	25
M	37	Triage	180	Mary	10
M	37	Release	305	Kate	20
M	37	Return	350	John	20
M	37	Blood test	360	Kate	15
F	52	Registration	0	John	25
F	52	Triage	300	Mary	25
F	52	Antibiotics	57900	Mark	10
F	52	Release	58010	Kate	15

Table 6. Feature vectors created from the log in Table 1 using aggregated encoding.

Sex	Age	Regist.	Antib.	Triage	Release	Return	Blood test	John	Mark	Mary	Kate	sum_Time	sum_Cost
M	37	1	0	0	0	0	0	1	0	0	0	0	15
M	37	1	1	0	0	0	0	1	1	0	0	80	40
M	37	1	1	1	0	0	0	1	1	1	0	180	50
M	37	1	1	1	1	0	0	1	1	1	1	305	70
M	37	1	1	1	1	1	0	2	1	1	1	350	90
M	37	1	1	1	1	1	1	2	1	1	2	360	105
F	52	1	0	0	0	0	0	1	0	0	0	0	25
F	52	1	0	1	0	0	0	1	0	1	0	300	50
F	52	1	1	1	0	0	0	1	1	1	0	57900	60
F	52	1	1	1	1	0	0	1	1	1	1	58010	75

Table 7. Feature vectors created from the log in Table 1 using index-based encoding, buckets of length

Sex	Age	Activ_1	Time_1	Res_1	Cost_1	Activ_2	Time_2	Res_2	Cost_2	Activ_3	Time_3	Res_3	Cost_3
M	37	Regist.	0	John	15	Antib.	80	Mark	25	Triage	180	Mary	10
F	52	Regist.	0	John	25	Triage	300	Mary	25	Antib.	57900	Mark	10

as the feature vector size depends on the trace length. Nevertheless, two options have been proposed in related work to combine index-based encoding with other bucketing types:

Fix the maximum pre x length and, for shorter pre xes, impute missing event attribute values with zeros or their historical averages. This approach is often referred to as **padding** in machine learning [46] and has been used in the context of predictive process monitoring in [52] and [33].

Use the sliding window method to encode only recent (up-to window size) history of the pre x. This approach has been proposed in [43].

### 5.3 Predictive algorithms

Machine learning-based predictive process monitoring methods have employed a variety of classification and regression algorithms, with the most popular choice being decision trees (e.g., [1]). Although quite simple, decision trees have an advantage in terms of computation performance and interpretability of the results. Other choices include random forests [51, 60], support vector regression [40] and extreme gradient boosting [43].

A recent cross-benchmark of 13 state-of-the-art commonly used machine learning algorithms on a set of 165 publicly available classification problems [5] found that gradient tree boosting generally achieves better accuracy than random forest, while random forest is more accurate than decision trees. Gradient boosting has also been shown to outperform other algorithms in the context of business process data [51]. As a result, predictive process monitoring methods using predictive algorithms that are inherently more accurate will perform better.

In order to eliminate the bias associated with the usage of different predictors in related work, we decided to fix extreme gradient boosting (XGBoost) as the main predictor across all the compared techniques. XGBoost is based on the theory of boosting, wherein the predictions of several weak learners (models whose predictions are slightly better than random guessing), are combined to produce a strong learner [2]. These weak learners are combined by following a gradient learning strategy. At the beginning of the calibration process, a weak learner is fit to the whole space of data, and then, a second learner is fit to the residuals of the first one. This process of fitting a model to the residuals of the previous one goes on until some stopping criterion is reached. Finally, the output of XGBoost is a weighted mean of the individual predictions of each weak learner. Regression trees are typically selected as weak learners [54]. It should be noted that the choice of a machine learning algorithm on one side and the choice of encoding and bucketing techniques on the other side are orthogonal choices—any algorithm can be combined with any feasible bucketing-encoding pair.

Nevertheless, we set aside recurrent neural networks (RNN) as another predictor applied in the primary method [50]. RNNs, and specially their long short term memory (LSTM) variant, have been shown to deliver consistently high accuracy in sequence modeling domains exhibiting temporal dynamics, including business [29, 33] and manufacturing [61, 63] processes. In this benchmark, we will evaluate LSTM models with zero bucketing, i.e., train a single LSTM model for all prefixes, as this is the only bucketing type that has been used with LSTMs in the literature. As unlike other predictors, LSTMs do not require attending the input data, we will naturally use tensor encoding with them. Similarly to the other predictors, to overcome the problem of feature vectors increasing with the prefix size, we fix the maximum number of events in the prefix (cf. Section 6.2.2) and for shorter prefixes, pad the data for missing events with zeros as in [33, 52].

## 5.4 Discussion

Summarizing the above observations, we devised a taxonomy of predictive monitoring techniques to be evaluated in our cross-benchmark (Figure 6). The taxonomy is framed upon a general classification of machine learning approaches into generative and discriminative ones (cf. Section 2.2).

The former correspond to process-aware predictive monitoring techniques, meaning that there is an assumption that an observed sequence is generated by some process that needs to be uncovered via probabilistic reasoning. The process can be represented via a state transition system [38], a Petri net [42] or a queueing model [45]. In our benchmark, we use implementations provided in [42, 57] as such, only making changes in the way the predictions are evaluated, in order to bring it in line with the other methods (cf. Section 6.2.2). The other primary method [60] had to be excluded due to the absence of a publicly available implementation.

Conversely, discriminative approaches are non-process-aware techniques that learn a direct mapping from inputs to the output via regression, without providing a model of how input sequences are actually generated. Having analyzed the discriminative studies, we have observed that they tend to mix different encoding types [2] or different bucketing types [19], while some combinations thereof have not been explored. As such, in our benchmark, we decided to evaluate all feasible

Fig. 6. Taxonomy of methods for predictive monitoring of remaining time.

combinations of canonical encoding and bucketing types, including those that have not been used in any existing approach in the literature. In this way, we will be able to assess the benefits of each combination, while abstracting implementation nuances of each individual primary method.

Finally, a range of newer studies propose hybrid methods that combine generative and discriminative approaches [40, 60]. Such methods can generally be approximated with state bucketing that, for every process state, fits a model to predict the remaining time starting from that state. Alternatively, the flow analysis technique [60] provides a higher degree of granularity by splitting the remaining time into its integral components – cycle times of individual activities to be executed according to the process model.

## 6 BENCHMARK

Based on the methods identified in the previous subsection, we conducted an extensive benchmark to identify relative advantages and trade-offs in order to answer RQ5. In this section, we describe the datasets, the evaluation setup and metrics, and present the results of the benchmark.

### 6.1 Datasets

We conducted the experiments using 17 real-life event datasets. To ensure the reproducibility of the experiments, the logs we used are publicly available at the Center for Research Data of March 2018, except for one log, which we obtained from the demonstration version of a software tool.

We excluded from the evaluation those logs that do not pertain to business processes (e.g., JUnit 4.12 Software Event Log and NASA Crew Exploration Vehicle). Such logs are usually not case-based or they contain only a few cases. Furthermore, we discarded the log that comes from the Environmental permit application process, as it is an earlier version of the BPIC 2015 event log from the same collection.

<sup>4</sup>In order to combine index-based encoding with bucketings other than prefix length-based, we use zero padding.

<sup>5</sup>[https://data.4tu.nl/repository/collection:event\\_logs\\_real](https://data.4tu.nl/repository/collection:event_logs_real)

All the logs have been preprocessed beforehand to ensure the maximum achievable prediction accuracy. Firstly, since the training and validation procedures require having a complete history of each case, we remove incomplete cases, as well as cases that have been recorded not from their beginning. For example, in the Road tra c nes log<sup>6</sup> we consider traces where the last recorded event isSend Fin to be pending and therefore incomplete. Secondly, we perform some basic feature engineering. For instance, using event timestamps, we extract weekday, hour and duration since the previous event in the given case and since the beginning of the case (elapsed time) from each log. Additionally, for categorical variables with many possible values, if some values appear very rarely (in less than 10 cases), these rare values are marked as *other*. Finally, we check if there are any data attributes that are constant across all cases and events, or cross-correlated with other attributes and discard them.

Table 8 summarizes the basic characteristics of each resulting dataset, namely the number of complete cases, the ratio of distinct (unique) traces (DTR), the number of event classes, the average number of distinct events per trace (DER), average case length, i.e., the average number of events per case and its coefficient of variation (CV), average case duration (in days) and its CV, and the number of case and event attributes. The datasets possess a very diverse range of characteristics and originate from a variety of domains.

Table 8. Statistics of the datasets used in the experiments.

log	# cases	DTR	event classes	DER	mean case length	CV case length	mean case duration	CV case duration	# attributes (case+event)	Domain
bpic2011	1140	0.858	251	0.505	131.342	1.542	387.283	0.875	6+10	HC
bpic2012a	12007	0.001	10	1	4.493	0.404	7.437	1.563	1+4	Fin
bpic2012o	3487	0.002	7	1	4.562	0.126	15.048	0.606	1+4	Fin
bpic2012w	9650	0.235	6	0.532	7.501	0.97	11.401	1.115	1+5	Fin
bpic2015_1	696	0.976	189	0.967	41.343	0.416	96.176	1.298	17+8	PA
bpic2015_2	753	0.999	213	0.969	54.717	0.348	159.812	0.941	17+8	PA
bpic2015_3	1328	0.968	231	0.975	43.289	0.355	62.631	1.555	18+8	PA
bpic2015_4	577	0.998	179	0.97	42	0.346	110.835	0.87	15+8	PA
bpic2015_5	1051	0.997	217	0.972	51.914	0.291	101.102	1.06	18+8	PA
bpic2017	31509	0.207	26	0.878	17.826	0.32	21.851	0.593	3+15	Fin
credit	10035	0	8	1	8	0	0.948	0.899	0+7	Fin
helpdesk	3218	0.002	5	0.957	3.293	0.2	7.284	1.366	7+4	CS
hospital	59228	0	8	0.995	5.588	0.123	165.48	0.671	1+20	HC
invoice	5123	0.002	17	0.979	12.247	0.182	2.159	1.623	5+10	FI
production	225		28	0.454	20.191	1.034	20.467	1.03	2+13	M
sepsis	1035	0.076	6	0.995	5.001	0.288	0.029	1.966	23+10	HC
tra c_ nes	150370	0.002	11	0.991	3.734	0.439	341.676	1.016	4+10	PA

AcronymsDTR distinct trace ratio, DER distinct event ratio, CV coefficient of variation

DomainsHC healthcare, Fin nancial, PA public administration, CS customer service, M manufacturing

## 6.2 Experimental Setup

In this section, we describe our approach to split the event logs into training and test sets along the temporal dimension. Next, we provide a description of our evaluation criteria and the baselines to

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compare against. Finally, we discuss the hyperparameter optimization procedure employed in our benchmark.

**6.2.1 Data split** In order to simulate a real-life situation where prediction models are trained using historical data and applied to running cases, we employ a so-called temporal split to divide the event log into train and test cases. Specifically, all cases in the log are ordered according to their start time and the first 80% are used to fit the models, while the remaining 20% are used to evaluate the prediction accuracy. In other words, the classifier is trained with all cases that started before a given date  $T_1$  which would represent a current point in time in a real-life scenario, and the testing is done only on cases that start afterwards. Technically, cases that start before and are still running at  $T_1$  should not be included in either set. However, to prevent the exclusion of a significant number of cases, in our experiments, we allow the two sets not to be completely temporally disjoint.

Fig. 7. Temporal split of the training and test sets.

**6.2.2 Evaluation metrics** Two measures commonly employed to assess a predictive process monitoring technique are accuracy and earliness [26, 53]. Indeed, in order to be useful, a prediction should be accurate and should be made early on to allow enough time to act upon.

**Accuracy.** To assess the accuracy of the prediction of continuous variables, well-known error metrics are Mean Absolute Error (MAE), Root Mean Square Error (RMSE) and Mean Percentage Error (MAPE) [23], where MAE is defined as the arithmetic mean of the prediction errors, RMSE as the square root of the average of the squared prediction errors, while MAPE measures error as the average of the unsigned percentage error. We observe that the value of remaining time tends to be highly varying across cases of the same process, sometimes with values on different orders of magnitude. RMSE would be very sensitive to such outliers. Furthermore, the remaining time can be very close to zero, especially near the end of the case, thus MAPE would be skewed in such situations. Hence, we use MAE to measure the error in predicting the remaining time. Formally,

$$MAE = \frac{1}{n} \sum_{i=1}^n |y_i - \hat{y}_i| \quad (2)$$

where  $y_i \in Y = R$  is the actual value of a function in a given point and  $\hat{y}_i \in Y = R$  is the predicted value.

**Earliness.** A common approach to measure the earliness of the predictions is to evaluate the accuracy of the models after each arrived event or at fixed time intervals. Naturally, uncertainty decreases as a case progresses towards its completion. Thus, the earlier we reach the desired level of accuracy, the better the technique is in terms of its earliness.

To measure earliness, we make predictions for pre- $\text{fixes}^{k-1}$  of traces in the test set starting from  $k = 1$ . However, using all possible values of  $k$  is coupled with several issues. Firstly, a large number of pre- $\text{fixes}$  considerably increases the training time of the prediction models. Secondly, for a single model approach, the longer cases tend to produce much more pre- $\text{fixes}$  than shorter ones and, therefore, the prediction model is biased towards the longer cases. Finally, for a



multiple model approach, if the distribution of case lengths has a long tail, for very long prefixes, there are not enough traces with that length, and the error measurements become unreliable. Consequently, we use prefixes of up to 20 events only in both training and test phase. If a case contains less than 20 events, we use all prefixes, except the last one, as predictions do not make sense when the case has completed. In other words, the input for our experiments is a filtered prefix  $L = \{hd^{k^1} : 2 \leq L; 1 \leq k \leq \min\{j, j-1\}; 20\}$ .

Inherently, there is often a trade-off between accuracy and earliness. As more events are executed, due to more information becoming available, the prediction accuracy tends to increase, while the earliness declines [3, 53]. As a result, we measure the performance of the models w.r.t. each dimension separately.

**6.2.3 Hyperparameter optimization** Each prediction technique is characterized by model parameters and by hyperparameters [3]. While model parameters are learned during the training phase so as to fit the data, hyperparameters are set outside the training procedure and used for controlling how flexible the model is in fitting the data. For instance, the number of clusters in the k-means clustering procedure is a hyperparameter of the clustering technique. The impact of hyperparameter values on the accuracy of the predictions can be extremely high. Optimizing their value is therefore important, but, at the same time, optimal values depend on the specific dataset under examination [2].

A traditional way of performing hyperparameter optimization is grid search, which is an exhaustive search through a manually specified subset of the hyperparameter space of a learning algorithm. A grid search algorithm must be guided by some performance metric, typically measured by cross-validation on the training set or evaluation on a held-out validation set [2]. In our benchmark, to find the best set of hyperparameters, we perform grid search using five-fold cross-validation. For each combination of hyperparameters, we train a model based on the 80% of the original training set and evaluate its performance on the remaining validation set. The procedure is repeated five times, while measuring the validation performance over each split using the same metrics as for the test set. Then we select one combination of the parameters that achieves the best validation performance and retrain a model with these parameters, now using the whole training set.

For machine learning-based techniques, we use the implementation of XGBoost from the scikit-learn library [36] for Python which allows for a wide range of learning parameters as described in the work by Tianqi and Carlos [3]. Table 9 lists the most sensitive learning parameters that were optimized during the grid search. To achieve the optimal predictive power, these parameters are tuned for each combination of dataset, bucketing method, and sequence encoding method. For approaches that involve clustering, we use the k-means clustering algorithm, which is one of the most widely used clustering methods in the literature. K-means requires one to specify in advance the desired number of clusters. We selected  $k = 10$  as the highest value because larger values resulted in some clusters with too few samples to train a classifier. We also chose  $k = 2$  as a representative of a low value of  $k$  and  $k = 5$  as an intermediate value. Thus, we searched for the optimal value in the set  $\{2, 5, 10\}$ . In the case of the index-based bucketing method, an optimal configuration was chosen for each prefix length separately.

For LSTMs, we used the recurrent neural network library [1], with Tensorflow backend. As with XGBoost, we performed tuning using grid search using the parameters specified in Table 9. These parameters and their values were chosen based on the results reported in [51]. Other hyperparameters were left to their defaults.

A similar procedure is performed for methods that do not train a machine learning model. For the method in [57], we vary the type of abstraction set, bag of sequence to create a transition system. For the method in [2], we vary the stochastic Petri net properties: (i) the kind of transition

Table 9. Hyperparameters tuned via grid search.

Parameter	Explanation	Search space
XGBoost		
n_estimators	Number of decision trees ( weak learners) in the ensemble	f 250; 500g
learning_rate	Shrinks the contribution of each successive decision tree in the ensemble	f 0.02; 0.04; 0.06g
subsample	Fraction of observations to be randomly sampled for each tree.	f 0.5; 0.8g
colsample_bytree	Fraction of columns (features) to be randomly sampled for each tree.	f 0.5; 0.8g
max_depth	Maximum tree depth for base learners	f 3; 6g
LSTM		
units	Number of neurons per hidden layer	f 100; 200g
n_layers	Number of hidden layers	f 1; 2; 3g
batch	Number of samples to be propagated	f 8; 32g
activation	Activation function to use	f ReLUg
optimizer	Weight optimizer	f RMSprop; Nadamg

distribution normal, gaussian kernel or histogram and (ii) memory semantics global preselection or race with memory. We select the parameters that yield the best performance on the validation set and use them for the test set.

6.2.4 Feature encoding parameters Section 5.2, we noted that the aggregation encoding requires us to specify an aggregation function for each event attribute. As such, for activities and resource attributes, we use the count(frequency) aggregation function that returns the number of times a given activity has been executed, or the number of activities a given resource has executed. The same function is applied to other categorical event attributes. For each numeric event attribute, we include two features in the feature vector: the mean value of the attribute and its standard deviation across the pre x.

### 6.3 Evaluation results

In this section, we present the evaluation results of all benchmarked methods and discuss their relative merits and trade-offs in order to answer RQ5. To keep the names of the methods short, in the following tables and figures, we refer to them as bucketingType\_encodingType where bucketingType can be one of the four options described in Section 5.0: Bucket for zero bucketing, pre x for pre x length bucketing, cluster for cluster bucketing and state for state bucketing. Analogously, we denote aggregation encoding agg, last state encoding last and index-based encoding by index. Tensor encoding is only used with the LSTM-based method, and is not denoted.

Table 10 reports the prediction accuracy, averaged across all evaluated pre x lengths, together with its standard deviation. The averages are weighted by the relative frequency of pre xes with that pre x (i.e., longer pre xes get lower weights, since not all traces reach that length). In our experiences, we set an execution cap of 6 hours for each training configuration, so if some method did not finish within that time, the corresponding cell in the Table is empty. Furthermore, since the flow-analysis approach [40] cannot readily deal with unstructured process models, we only provide its results for the logs from which we were able to derive structured models.

Rows 5 and onward in Table 10 each refer to a combination of a bucketing and a feature encoding technique. In these rows, we report the results obtained using an XGBoost regressor (hyperparameter-optimized as discussed above). The TS and SPN rows (transition systems and stochastic Petri net) do not use a regressor but rather estimate the remaining time directly from the generative model they create. The FA row uses optimized XGBoost regressors to estimate the cycle time of each remaining activity.

Overall, we can see that in 14 out of 17 datasets, LSTM-based networks achieve the best accuracy, while low-analysis and index-based encoding with no bucketing and pre x-length bucketing achieve the best results in one dataset each. At the same time, we observe a trade-off between accuracy and explainability of the predictions, among the surveyed methods. Indeed, transition systems and stochastic Petri nets that provide more explainable predictions usually achieve lower prediction accuracy than black-box models such as LSTMs. This trade-off should be considered by process analysts when choosing a suitable prediction method, and it reflects the general machine learning observation that more accurate models are more complex and are harder for users to interpret [5].

Figure 8 presents the prediction accuracy in terms of MAE, evaluated over different pre x lengths <sup>7</sup>. Each evaluation point includes pre xes of exactly the given length. In other words, traces that are shorter than the required pre x are left out of the calculation. Therefore, the number of cases used for evaluation is monotonically decreasing when increasing the pre x length.

In most of the datasets, we see that the MAE decreases as cases progress. It is natural that the prediction task becomes trivial when cases are close to completion. However, for some datasets, the predictions become less accurate as the pre x length increases. This phenomenon is caused by the fact that these datasets contain some short traces for which it appears to be easy to predict the outcome. These short traces are not included in the later evaluation points, as they have already finished by that time. Therefore, we are left with longer traces only, which appear to be more challenging for the predictor, hence decreasing the total accuracy on larger pre x lengths. However, different techniques behave differently wrt. earliness. For example, LSTMs generally provide the most accurate predictions early on, but as the case progresses, other techniques may outperform LSTMs.

As a simple bulk measure to compare the performance of the benchmarked techniques, we plot their mean rankings achieved across all datasets in Figure 9. Ties were resolved by assigning every tied element to the lowest rank. The rankings illustrate that LSTMs consistently outperform other machine-learning baselines in terms of accuracy (measured by MAE), while stochastic Petri nets and transition systems are usually the least accurate methods.

To complement the above observations, we also compare aggregated error values. These values need to be normalized, e.g., the by mean case duration, so that they are on a comparable scale. In order to do that, for each log, we divide the average MAE values and their standard deviations across all pre xes reported in Table 10 by the mean case duration for that log reported in Table 8. The results for each technique are illustrated in the boxplots in Figure 10, where each point represents the results for one of the 17 datasets. We can see that LSTM-based techniques have an average error of 40% of the mean case duration across all datasets. In contrast, transition systems on average incur a 59% error. Importantly, for LSTMs the accuracy varies between 0.07 and 0.56, while index-based encoding with pre x length bucketing, the method that on average achieves the second most accurate results, is more volatile and varies between 0.08 and 0.90 of the mean case duration.

Another important observation is related to the temporal stability of the predictions. In general, methods that provide higher accuracy also have lower volatility of error across case lifetime (Figure 10b). In other words, the difference between successive predictions obtained from these methods is lower, as evidenced by lower standard deviation of the error metrics.

In order to assess the statistical significance of the observed differences in methods' performance across all datasets, we use the non-parametric Friedman test. The complete set of experiments indicate statistically significant differences according to this test ( $p \leq 4.642 \cdot 10^{-8}$ ). Following the

<sup>7</sup>To keep the plots readable, we removed 4 methods that generally achieve lower prediction accuracy across most datasets, according to Table 10: zero bucketing and state bucketing with last state and aggregation encoding.



Fig. 8. Prediction accuracy (measured in terms of MAE) across different prefix lengths

Fig. 9. Average ranking of the evaluated methods over all datasets.

(a) (b)

Fig. 10. Average normalized MAE values (a) and their standard deviation (b) across case lifetime.

procedure suggested in the recent work on evaluating machine learning algorithms in order to find which methods in particular differ from each other, we use the Nemenyi post-hoc test that compares all methods to each other.

Table 11. Post-hoc Nemenyi test of methods' rankings across all datasets.

	cluster_agg	cluster_last	LSTM	noBucket_agg	noBucket_last	pre_x_agg	pre_x_index	pre_x_last	state_last
cluster_last	0:996								
LSTM	0:098	0421							
noBucket_agg	0:998	0894	0:005						
noBucket_last	0:995	0847	0:003	1					
pre_x_agg	0:999	0931	0:007	1	1				
pre_x_index	0:999	1	0515	0833	0773	0883			
pre_x_last	1	0958	0:011	1	1	1	0923		
state_last	0:958	0631	0:001	1	1	1	0535	1	
TS	0:169	0:025	0	0:740	0804	0669	0:016	0:593	0946

Table 11 lists p-values of a pairwise post-hoc analysis. Since the test requires complete information for all pairwise comparisons, we included only 10 methods for which we have results on all 17 datasets. For most pairs, the null hypothesis that their performance is similar can not be rejected. However, the test underlines the impressive performance of LSTM, which significantly outperforms most of the other methods at the  $p < 0.05$  level.

While on average most combinations of bucketing and encoding methods provide more or less similar levels of accuracy, we can observe differences for individual datasets. For example, in the hospital dataset, it is clear that clustering with aggregation encoding is better than with clustering with last state encoding. Arguably, aggregating knowledge from all events performed so far provides much more signal than using raw attributes of the latest event.

In order to explain differences in performance of various bucketing and encoding combinations, we try to correlate the characteristics of event logs (Table 8) with the type of bucketing/encoding that achieves the best accuracy on that log. One can notice that if cases in the log are very heterogeneous in terms of case length, i.e., the coefficient of variation of case length is high enough, it is more beneficial to assign all traces to the same bucket. This can be observed in `bpic2011`, `bpic2012w` and `production` event logs, where standard deviation of case length is close to or exceeds mean case length. Furthermore, if cases in the log are short (e.g., `helpdesk`, `tra_c_nes`, `bpic2012a`, `bpic2012b`) or very distinctive from each other (e.g., `bpic2015_2`) last state encoding tends to capture the most signal. Notably, in the aforementioned logs, the index-based encoding, although lossless, is not optimal. This suggests that in these datasets, combining the knowledge from all events performed so far provides more signal for remaining time prediction than the order of events. However, standard classifiers like XGBoost are not able to learn such higher-level features, unlike LSTMs, which is why in some situations even the simple aggregations outperform the index-based encoding.

**Execution Times.** The machine learning models took between 15 minutes to 5 hours to train on conventional hardware depending on the size of the dataset. The choice of bucketing and feature encoding method does not significantly influence execution times, except for the fact that index-based encoding takes about three times more execution time than last state encoding and aggregation encoding due to the fact that it leads to larger feature sets. Similarly, the FA and SPN methods take between a few minutes to 3 hours. Note that model training is an offline operation. A few hours of model training time is acceptable in practical settings since this operation can be performed as an overnight batch job.

The LSTM models require between 15 and 90 seconds per training iteration depending on the dataset. In the benchmark, we applied 50 iterations and thus the training times per model ranged from 13 to 80 minutes on a single NVIDIA Tesla k80 Graphics Processing Unit (GPU). Training times on conventional CPU processors would be an order of magnitude higher.

The execution time to make a prediction for a given prefix (i.e., for each new event to be handled) is in the order of milliseconds for LSTM and machine learning methods and close to a second for the FA method.

## 7 THREATS TO VALIDITY

One of the threats to the validity of this study relates to the potential selection bias in the literature review. To minimize this, we described our systematic literature review procedure on a level of detail that is sufficient to replicate the search. However, in time the search and ranking algorithms of the used academic database (Google Scholar) might be updated and return different results. Another potential source of bias is the subjectivity when applying inclusion and exclusion criteria,

<sup>8</sup>These measurements were made on a laptop with a 2.4GHz Intel Core i7 processor and 16GB of RAM.



as well as when determining the primary and subsumed studies. In order to alleviate this issue, all the included papers were collected in a publicly available spreadsheet, together with decisions and reasons about excluding them from the study. Moreover, each paper was independently assessed against the inclusion and exclusion criteria by two authors, and inconsistencies were resolved with the mediation of a third author.

Another threat to validity is related to the comprehensiveness of the conducted experiments. In particular, only one representative setting of each technique was used. Namely, only one machine learning algorithm (XGBoost) and one clustering method (k-means) were tested over all relevant methods. It is possible that there exists, for example, a combination of an untested clustering technique and a predictor that outperforms the settings used in this study. Furthermore, the generalizability of the findings is to some extent limited by the fact that the experiments were performed only on 17 event logs. Although these are all real-life event logs from different application fields that exhibit different characteristics, it may be possible that the results would be different using other datasets or different log preprocessing techniques for the same datasets. In order to mitigate these threats, we built an open-source software framework which allows the full replication of the experiments, and made this tool publicly available. Moreover, additional datasets, as well as new sequence classification and encoding methods can be plugged in, so that the framework can be used for future experiments.

## 8 CONCLUSION

This study provided a survey, a taxonomy and an empirical evaluation of methods to predict the remaining cycle time of business process executions. The relevant existing studies were identified through a systematic literature review, which retrieved 25 studies dealing with the problem of remaining time prediction in business process executions. Out of these 25 studies, 10 of them propose distinct methods (primary studies). Through further analysis of the primary studies, a taxonomy was proposed based on three main aspects: (i) the type of input data required; (ii) the process-awareness of the methods; (iii) and the algorithms employed to derive a predictive model from the data.

It was found that 7 out of the 10 primary studies employ discriminative machine learning algorithms (specifically regression) to train predictive models, while three used generative or hybrid generative-discriminative methods. The discriminative and hybrid methods are further broken down into sub-categories depending on how they divide the input traces into homogeneous buckets (trace bucketing) and how the traces in the event log are encoded as feature vectors (feature encoding). Based on this taxonomy, we identified 16 distinct methods, some of which have been explicitly studied in the literature, while others are derived by combining features of previously studied methods.

The 16 identified methods were then empirically evaluated based on a benchmark consisting of a unified experimental setup (including hyperparameter optimization where applicable) executed over 17 real-life, publicly available event logs. To ensure reproducibility of the empirical results, all the selected methods were implemented as an open-source framework.

The results of the empirical evaluation show that the most accurate results are obtained using LSTM neural networks, possibly owing to their ability to automatically learn relevant features from trace prefixes. On the other end of the spectrum, generative methods based on transition systems and stochastic Petri nets achieve the lowest accuracy. Regression methods fill the middle ground between these two ends. We found that there is no statistically significant difference between different canonical combinations of bucketing and feature encoding approaches. In light of the above, the overall recommendation resulting from this study is to use LSTM methods when the necessary computational resources are available for model training (e.g., availability of GPUs).

Otherwise, simpler methods such as last-state feature encoding without bucketing should be preferred given that they are the simplest approach. The mean ranking of the various methods over the 17 datasets suggest that clustering-based methods should be preferred over state-based methods, but this observation was not confirmed by the statistical significance tests. If the computational resources allow so, it is recommended to try out multiple methods over a given dataset prior to selecting one.

The results suggest that there is room for new methods that would consistently outperform existing techniques based on machine learning and that would be less computationally demanding (for model training) than deep learning techniques such as LSTMs. We also note that most existing methods are based on black-box models. Generative and hybrid techniques such as flow analysis (FA) and Stochastic Petri nets (SPN) are the only ones that provide interpretable models, but the accuracy of SPNs is very low (and they turn out to be difficult to interpret for non-expert users), while FA does not have a consistently high accuracy across datasets. Designing interpretable and consistently accurate approaches to remaining cycle time prediction remains an open challenge. Finally, we note that existing techniques are designed with accuracy in mind, and they neglect alternative quality dimensions such as stability of the sequential predictions made for a given case. While the problem of stability in predictive process monitoring has been studied in the context of prediction of case outcomes [5], this question has not yet been studied in the context of remaining cycle time prediction. This is another avenue for future work in the field.

**Reproducibility** All the source code required to reproduce the reported evaluation is available at <https://github.com/verenich/time-prediction-benchmark>. All the datasets employed are publicly available via the links provided in Section 6.1.

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