Modeling Dynamical Systems with Data Stream Mining

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Abstract. We address the task of modeling dynamical systems in discrete time using regression trees, model trees and option trees for on-line regression. Some challenges that modeling dynamical systems pose to data mining approaches are described; these motivate the use of methods for mining data streams. The algorithm FIMT-DD for mining data streams with regression or model trees is described, as well as the FIMT-DD based algorithm ORTO, which learns option trees for regression. These methods are then compared on several case studies, i.e., tasks of learning models of dynamical systems from observed data. The experimental setup, including the datasets, and the experimental results are presented in detail. These demonstrate that option trees for regression work best among the considered approaches for learning models of dynamical systems from streaming data.

Keywords: dynamical systems, data streams, data mining, regression and model trees, option trees

1. Introduction

The central question addressed by this paper is the applicability of the paradigm of mining data streams, and in particular tree-based approaches to on-line regression, to the task of system identification in discrete-time. In this paper, we propose the use of regression trees [1], model trees [15] and option trees [12] to address the this task. Namely, these approaches have been adapted to the on-line setting \cite{9,10} and are very suitable for the task of system identification.

First, regression and model trees allow us to address the general task of system identification, including structural and parameter identification, as they can approximate arbitrary non-linear functions with piece-wise linear functions. Second, they can handle well both missing and noisy data, which are likely to be encountered in practice. Third, they are very efficient and can handle large quantities of data: They scale well both in terms of the number of data points, i.e., time points in the case of system identification, and the number of independent variables (features, attributes), i.e., input and system variables in the case of system identification. Recently, tree-based approaches to classification and regression have been adapted to work in the on-line, streaming setting \cite{3}.

To address the central question of this paper, we perform a thorough experimental evaluation, where we apply several tree-based approaches to several case studies (benchmark problems) in discrete-time modeling of dynamical systems coming from the area of control engineering. Using an evaluation methodology and performance measures appropriate for the task at hand, we evaluate the performance of the tree-based approaches to on-line regression and answer several more specific experimental questions.
The remainder of the paper is structured in the following way. First, in Sec. 2, we present the background related to this work. Next, in Sec. 3, we present the task of modeling dynamical systems in discrete time using data stream mining and present the tree-based algorithms for regression on data streams that will be used to solve the task. Furthermore, in Sec. 4, we describe the case studies and in Sec. 5, we introduce the experimental setup, including the experimental questions and the evaluation methodology. Finally, in Sec. 6, we present and discuss the results of the experiments in Sec. 7, we conclude the paper and suggest several directions for further work.

2. Background

2.1. Modeling Dynamical Systems in Discrete Time

Dynamical systems are described in terms of system variables ($y$) and, in the case of controlled systems, control variables ($u$), depending on either discrete or continuous time ($t$). Modeling dynamical systems in continuous time is concerned with the formulation of differential equations for the time derivatives of the $\dot{y}(t)$ system variables in terms of the input variables $u(t)$ and the systems variables themselves $y(t)$. The model of a dynamical system should correspond to the observed behavior of the system.

If we discretize this problem, we sample the time space from an initial time $t_0$ at fixed intervals with a time step of $\Delta t$, at which we obtain observations. We describe the discrete time space with a variable $k$ that corresponds to $t$ with the relation $k \sim i \iff t = t_0 + i \Delta t$, for $i = 1, 2, 3, \ldots$

In discrete time, recurrence relations replace differential equations as the formalism for describing the dynamical system. Discrete-time modeling of dynamical systems is then concerned with the formulation of the recurrence relations which best describe the observed system behavior. These recurrence relations define the present values of the system variables $y(k)$ in terms of past values of the system variables and present and past values of the input variables, i.e., in terms of the vector

$$x(k) = [u(k-1), \ldots, u(k-n), y(k-1), \ldots, y(k-n)]^T,$$

where $n$ is the system lag.

2.2. System Identification via Regression

To model a dynamical system in discrete time we need to define a function $f$ so that

$$y(k) = f(x(k)),$$

where $x(k)$ is defined by Eq. (1). The task of finding such a function $f$ from an observed behavior of the system is called system identification. When the form of $f$ is known and only its constant parameter values are to be determined, the task is called parameter identification. When the form of $f$ is not known, the task at hand is much more difficult. We need to determine both the form and the constant parameter values of the function $f$. In this case, the task is called structural identification.

When we observe a dynamical system we take measurements of the input and output vector pairs $u, y$ at each time point $k$, so that we record the behavior of the system. An example of such a behavior, recorded for the gas-liquid separator case study presented in Sec. 4.1, see Fig. [3]. Through the external dynamics approach [14], we transform the
observations into a table of lagged input and output vector pairs. The columns of the table are as described by the vector \( x(k) \) from Eq. 1. Each row of the table corresponds to a point in time where an observation has been made (see Tab. 1).

The external dynamics approach allows us to formulate the task of system identification as a regression task, where we want to predict the values of the system variables from the lagged values of the system and input variables. The dependent (response) variable is \( y(k) \), while the independent variables are gathered in the vector \( x(k) \). The data used to learn the regression function \( y(k) = f(x(k)) \) are taken from the recorded behavior as described above.

### 2.3. Data Stream Mining

Data stream mining is a sub-field of data mining, which is concerned with the development of methods for modeling of data arriving in the form of a stream [3]. Mining a data stream is substantially different from classical data mining, mainly in the way data is reformulated according to the external dynamics approach.

#### Table 1

An excerpt from the recorded behavior of the GLS system depicted in Fig. 1, reformulated according to the external dynamics approach.

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#### Fig. 1

Recorded values of the input variables (valves 1 and 2, Liquid level) and system variable (pressure) of the gas-liquid separator system. Input variables are plotted with dotted lines, the system variable is plotted with a continuous line.
available. In the classical batch-learning data mining setting, a fixed and complete dataset is given as input to a learning algorithm, which can then use the data in an unrestricted fashion. The examples or data points are assumed to all come from the same probability distribution. In contrast to classical data mining, where all the data is available from the beginning of the process, in the on-line approach data stream mining the data instances arrive sequentially in time.

The key properties of data streams and on-line learning are:

1. the examples (data points) arrive sequentially,
2. the (data mining) algorithm has no control over the order of arrival of the examples,
3. there can potentially be arbitrarily many examples,
4. the distribution of examples need not be stationary,
5. there is a need for a response in real-time, and
6. after an example is processed it is discarded or archived – we can not access it directly, unless we explicitly store it into memory (which is comparatively small comparing to the number of examples).

3. Modeling Dynamical Systems in Discrete Time Using Data Stream Mining

The task of discrete-time modeling of nonlinear dynamical systems from measured data has been approached using different regression techniques. Approaches used for this purpose include the basis-function approaches of Artificial Neural Networks [14] and fuzzy modeling, as well as the nonparametric approaches of kernel methods [2] and Gaussian Process models [16].

The task of system identification, as defined in the previous subsection, matches the properties of learning models on data streams almost to the letter (see Sec. 2.3). This allows us to formulate the task of modeling dynamical systems in discrete time using data stream mining algorithms. If we observe a dynamical system, the individual data points arrive sequentially along the time dimension: The system identification (learning) algorithm has no control over the order of arrival of the examples/observations. Depending on whether we continue to observe the system as time goes by, there can potentially be infinitely many examples. Depending on how frequently we make observations, the amount of data collected can easily grow over the available storage capacity. Finally, the laws underlying the behavior of a given system can change over time, especially if external disturbances appear in its environment (e.g., a pollution introduced into an ecosystem). Such dynamical systems are called time-varying dynamical systems.

To address the task of system identification using data stream mining, we use algorithms for learning tree-like models for regression [9,10]. They can learn model trees, as well as option trees for regression, and can also detect changes in the streaming data. Here, we briefly describe the algorithms that we use for modeling of dynamical systems.

3.1. FIMT-DD: Learning Model Trees with Change Detection

The FIMT-DD algorithm learns regression and model trees from data streams [9]. A model tree consists of internal nodes, which contain tests on independent variables (called
Fig. 2. An example regression tree learned on data from the gas-liquid separator domain.

The pseudo code of the FIMT-DD algorithm can be found in Alg. 1. The key points of the algorithm are the selection of splits for the internal nodes, the learning of linear models in the leaf nodes and the detection of changes. Algorithms for learning trees on data streams use statistical tests to determine when enough examples have been accumulated to make a split decision.

Algorithm 1 The FIMT-DD algorithm. Pseudocode.

Input: $\delta$ – confidence level, $N_{\text{attr}}$ – number of attributes, $n_{\text{min}}$ – chunk size.
Output: $T$ – root node of the current model/regression tree.

for $\infty$ do
    $e \leftarrow \text{ReadNextExample}()$
    $\text{Leaf} \leftarrow \text{Traverse}(T, e)$
    $\text{Seen} \leftarrow \text{SeenAtLeaf}(\text{Leaf}) + 1$
    $\text{UpdateStatistics}(\text{Leaf}, e)$
    $\text{UpdateLeaf}(\text{Leaf}, e)$
    if $\text{Seen} \mod n_{\text{min}} = 0$ then
        for $i = 1 \rightarrow N_{\text{attr}}$ do
            $h_i \leftarrow \text{FindBestSplitForAttribute}(i)$
        end for
        $h_A \leftarrow \text{BestSplit}(h_1, \ldots, h_{N_{\text{attr}}})$
        $h_B \leftarrow \text{SecondBestSplit}(h_1, \ldots, h_{N_{\text{attr}}})$
        if $\frac{\text{VR}(h_A)}{\text{VR}(h_B)} < 1 - \sqrt{\frac{\ln(1/\delta)}{2 \cdot \text{Seen}}}$ then
            $\text{MakeSplit}(h_A, T)$
        end if
    end if
end for
Algorithms for learning decision trees on data streams start with a single leaf node. Leaves are then split when enough evidence has accumulated that the more complex tree resulting from the split will perform better than the leaf node. Deciding when to split a leaf is a critical step in tree construction. For this purpose, FIMT-DD utilizes the variance reduction heuristic along with the Hoeffding bound [6].

The basic version of FIMT-DD learns regression trees, where the predictions in the leaves are constant. An extension of the FIMT-DD algorithm learns model trees instead of regression trees. In model trees, at a given leaf, instead of returning the average of target variable values for the accumulated examples, we use linear regression to predict the target value of incoming examples.

To this end, we place a single layer perceptron into each leaf. Given inputs \( x_1, \ldots, x_n \) (attribute values), the output is computed as
\[
o = \sum_{i=1}^{n} x_i w_i + b,
\]
where \( w_i \) are the weights and \( b \) is the threshold. Using the squared error function for an example \( e, E(x) = \frac{1}{2} (y - o)^2 \), where \( y \) is the actual target value and \( o \) is the predicted target value, we can continuously update the perceptron according to the delta or Widrow-Hoff additive rule. The complete perceptron learning scheme (procedure UpdateLeaf in Alg. [1]) is as follows

1. The weights \( w_1, \ldots, w_n \) and the threshold \( b \) are initialized to random values.
2. Consider a vector \( x = (x_1, \ldots, x_n) \) and calculate \( o \).
3. Update the weights using the delta rule \( w \leftarrow w + \eta (y - o)x \); (\( \eta \) is the learning rate).
4. Return to step 2.

In our case, the learning rate is not kept constant but is decreased inversely proportional to the number of instances \( \eta = \frac{\eta_0}{1 + N \eta_d} \), where \( \eta_0 \) is the initial learning rate, \( N \) is the number of examples processed and \( \eta_d \) is the learning rate decay parameter.

To detect changes, FIMT-DD makes use of the Page-Hinckley test (see Alg. [2]). The test is a mechanism for detecting changes in a signal \( x(k) \). At any time point \( k \), the test considers the following two variables: a cumulative sum \( m(k) \) and its minimal value \( M(k) = \min_{i=1,\ldots,k} m(i) \). The sum \( m(k) \) is defined as the cumulative difference between the signal \( x(k) \) and its average \( \overline{x}(k) \) corrected with an additional parameter \( \alpha \) (minimal absolute amplitude of change that we wish to detect), \( m(k) = \sum_{i=1}^{k} (x(k) - \overline{x}(k)) - \alpha \).

The Page-Hinckley test monitors the difference between \( m(k) \) and \( M(k) \), \( PH(k) = m(k) - M(k) \). When this difference passes a threshold \( \lambda \), an alarm is triggered which informs the algorithm of a possible change. To detect possible change in the data, we use the Page-Hinckley test on the absolute error signal \( |y - \hat{y}| \), where \( y \) is the true value of an example and \( \hat{y} \) is the predicted value. The error is back-propagated towards the root of the tree, as the Page-Hinckley test is monitored at each node in the tree.

When an alarm is triggered, the FIMT-DD algorithm responds by starting to grow an alternate subtree. The tree is grown at the node at the lowest level which has triggered the alarm. The alternate tree is grown in parallel with the original subtree, while the growth of the original subtree is stopped. At a time interval of \( T_{min} \), the viability of both the original \( O \) and the alternate tree \( A \) is compared using the faded \( Q_i \) statistic. If \( Q_i(O, A) \) is greater than 0, that means that the alternate subtree is better in terms of the predictive performance and the original subtree is replaced with the alternate. However, if after \( 10T_{min} \) examples, the alternate tree has not managed to produce better predictions, we discard the alternate tree and again allow the growth of the original tree.
Algorithm 2 Traverse procedure of the FIMT-DD algorithm, pseudocode.

Input: $T$ – node (root, internal or leaf), $e$ – training example
Output: $Leaf$ – leaf node where the example $e$ will be traversed

if $\text{IsLeaf}(T) \neq \text{True}$ then
    $Change \leftarrow \text{UpdatePHTest}(T, e)$
    if $Change = \text{True}$ then
        $\text{InitiateAlternateTree}(T, e)$
    end if
    $Child \leftarrow \text{ApplyNodeTestToExample}(T, e)$
    $Leaf \leftarrow \text{Traverse}(Child, e)$
else
    $Leaf \leftarrow T$
end if
return $Leaf$

3.2. ORTO: Learning Option Trees for Regression on Data Streams

The ORTO algorithm is an extension of the FIMT-DD algorithm which introduces option nodes into the learned regression (or model) trees [10]. When making a prediction for a new example with a regular regression (or model) tree, the example is sorted down exactly one branch of the tree, i.e., sorted to exactly one branch of each internal node it encounters. In contrast, when sorting an example through an option node, the example is multiplied and sorted to each of the children of an option node. This means that in each option node multiple predictions are produced, corresponding to the multiple branches out of the node followed by the example. These predictions are then aggregated to form a single prediction, e.g., by averaging. An example option tree for regression is presented in Fig. 3.

Fig. 3. An example option tree learned on data from the gas-liquid separator domain. The highlighted area represents the best tree.
The pseudocode of ORTO is presented in Alg. 3. The split decision process of ORTO is almost identical to that of FIMT-DD, but it includes an important alteration, which allows the introduction of option nodes. We use two aggregation approaches, averaging and choosing the best path. **Averaging** is the more straightforward of the two approaches. The final prediction of the option node is obtained by averaging the predictions obtained from the children of the option node. We refer to this variant of the ORTO algorithm as ORTO-A. The other aggregation approach used is to **choose the (currently) best path**. The cumulative errors for each example processed are kept for each of the children of an option node, and when combining the predictions, the prediction of the child with currently the

**Algorithm 3 The ORTO algorithm. pseudocode.**

**Input:** \( \delta \) – confidence level, \( N_{attr} \) – num. of attributes, \( n_{min} \) – chunk size, \( Agg \) – aggregation method, \( \beta \) – option decay factor, \( T_{max} \) – max. number of options.

**Output:** \( T \) – current option tree.

\[
T \leftarrow \text{InitiateOptionTree}()
\]

for \( \infty \) do

\[
e \leftarrow \text{ReadNextExample}()
\]

\[
\text{Leaf} \leftarrow \text{Traverse}(T, e)
\]

\[
\text{Seen} \leftarrow \text{SeenAtLeaf}(\text{Leaf}) + 1
\]

\[
\text{UpdateStatistics}(\text{Leaf}, e)
\]

\[
\text{UpdateLeaf}(\text{Leaf}, e)
\]

if \( \text{Seen} \mod n_{min} = 0 \) then

\[
\text{for } i = 1 \rightarrow N_{attr} \text{ do}
\]

\[
\text{FindBestSplitForAttribute}(i)
\]

end for

\[
h_A \leftarrow \text{BestSplit}(h_1, \ldots, h_{N_{attr}})
\]

\[
h_B \leftarrow \text{SecondBestSplit}(h_1, \ldots, h_{N_{attr}})
\]

if \( \frac{VR(h_A)}{VR(h_B)} < 1 - \sqrt{\frac{\ln(1/\delta)}{2 \cdot \text{Seen}}} \) then

\[
\text{MakeSplit}(h_A, T)
\]

else

\[
O \leftarrow \text{CreateOptionNode}(T, \text{Leaf})
\]

\[
\text{AddSplit}(h_A, O)
\]

\[
\text{Count} \leftarrow 0
\]

\[
k \leftarrow \text{CountSplitCandidates}(h_1, \ldots, h_{N_{attr}})
\]

\[
l \leftarrow \text{Level}(T, \text{Leaf})
\]

\[
\text{for } i = 1 \rightarrow N_{attr} \text{ do}
\]

\[
\text{if } i \neq 1 \text{ and Count } < k \cdot \beta^l \text{ and } \frac{VR(h_i)}{VR(h_A)} > 1 - \sqrt{\frac{\ln(1/\delta)}{2 \cdot \text{Seen}}} \text{ and NumberOfOptions}(T) < T_{max} \text{ then}
\]

\[
\text{AddSplit}(h_i, O)
\]

\[
\text{Count} \leftarrow \text{Count} + 1
\]

end if

end for

end if

end if

return \( \text{Predict}(T, Agg, e) \)

end for
lowest error is returned as the option node’s prediction. For example, the leftmost child of the option node of the tree in Fig. 3 has the lowest error and would be used when making predictions. This variant of ORTO is referred to as ORTO-BT.

4. Case Studies

In this paper, we demonstrate the applicability of tree-based approaches to on-line regression to the task of system identification in discrete-time, on three use cases coming from the area of control engineering. These include the following case studies: modeling the gas-liquid separator [11], modeling the process of pH neutralization [5], and modeling a synthetic dynamical system [13]. Below we present briefly each of the case studies.

4.1. Case Study: Gas-Liquid Separator

The gas-liquid separator is a unit in a semi-industrial process plant which is part of a larger pilot plant. The unit’s purpose is to capture flue gases under low-pressure from the effluent channels using a water flow, cool the gases down, and supply them with higher pressure to other parts of the plant. A schematic representation of the unit is given in Fig. 4.

The flue gases coming from the effluent channels are absorbed by the water into the water circulation pipe through the injector $I_1$. The water flow is produced by the water ring pump $P_1$, which operates at constant speed. The gas-water mixture is fed by the pump into the tank $T_1$, where the gas-liquid separation occurs. The accumulated gases form a pressurized gas ‘cushion’. This pressure forces the flue gases to be blown out from the tank into a neutralization unit. Similarly, the cushion pushes the water to circulate back into the reservoir. The amount of water in this circuit is kept constant. The entire process is controlled through a manipulation of the $V_1$ and $V_2$ valves.

This process is modeled with a set of differential equations [11], one of which is presented below: The equation describes the rate of change of the $p_1$ variable, which

![Fig. 4. A schematic representation of the system in the gas-liquid separator case study.](image-url)
corresponds to the relative air pressure in the tank $T_1$

$$\frac{dp_1}{dt} = f_a(h_1) \cdot \left[ \alpha_1 + \alpha_2 p_1 + \alpha_3 p_1^2 + f_b(u_1)\sqrt{p_1} + f_c(u_2)\sqrt{p_1 + \alpha_4 + \alpha_5 h_1} \right], \quad (2)$$

where $h_1$ is the liquid level in $T_1$, $u_1$ and $u_2$ are the command signals for the $V_1$ and $V_2$ valves, respectively, $\alpha_1, \ldots, \alpha_5$ are constants, while $f_a$, $f_b$, and $f_c$ are functions of the appropriate variables. In more detail, $f_a$ is a rational function of $h_1$, while $f_b$ and $f_c$ are valve characteristics, i.e., exponential functions of input signals $u_1$ and $u_2$, respectively.

In previous work, a complete non-linear dynamic model was derived and modeled in MATLAB by using Simulink [17]. The model consists of several parts: models of the valves $V_1$ and $V_2$, a model of the air pressure in the separator, a model of the water and air flow through the injector, and a model of the water levels in the separator and the reservoir (see Fig. 5). For the purpose of comparison, we replicated the simulation model using the same parameters (see Sec. 5).

The modeling task in this case study is to build a predictive model that can predict the value $p_1$ from its previous (lagged) values, as well as the current and previous values of the input variables. For the purpose of modeling, $h_1$ was also taken as an input variable, in addition to the control signals $u_1$ and $u_2$ for the valves $V_1$ and $V_2$. For this dataset a system lag of 3 was selected.

The dataset is composed of two runs of the gas-liquid separator, each sampled twice per second for about two hours, netting around 14500 examples each. For the purposes of learning a predictive model, we have concatenated these two runs into a single dataset, producing 29036 total examples.

**Fig. 5.** A model for the gas liquid separator in Simulink, consisting of five sub-models
4.2. Case Study: pH Neutralization

This case study is concerned with the problem of controlling alkalinity, a common problem in biotechnological and chemical processes. In particular, it seeks to identify the pH neutralization process, which exhibits severe nonlinear behavior.

The system is composed of several fluid streams: an acid stream $Q_1$, a buffer stream $Q_2$ and a base stream $Q_3$. The streams are mixed in a tank $T_1$. Prior to the mixing, the acid stream passes through another tank $T_2$. The observed variable is the effluent pH, which is controlled by manipulating the flow rate of the base stream $Q_3$, while both the acid and the buffer flows are kept constant. This process is schematically presented in Fig. 6.

The model of this dynamical system contains the following state, input (control) and output variables:

$$x = [W_{a4}, W_{b4}, h_1]$$

$$u = Q_3, y = \text{pH},$$

where $W_{a4}$ and $W_{b4}$ are the effluent reaction invariants and $h_1$ is the liquid level in the tank $T_1$. Following from here, it is assumed that a (previously designed) controller is being used to keep the fluid level in the tank $T_1$ constant at $h_1 = 14$ cm. This is achieved through the manipulation of the flow rate of the effluent stream $Q_4$.

The mathematical model obtained has the following form:

$$x = f(x) + g(x)u$$

$$c(x, y) = 0,$$

where $f(x)$ and $g(x)$ are non-linear functions of the state vector $x$, while $c(x, y)$ is a non-linear function that implicitly defines the value of the output variable $y$.

In total, a set of 31998 data instances was produced by simulating the above model. We apply the lagging procedure with a system lag of 2. This provides 4 features to be used in the modeling of the system behavior, i.e., $y(k - 1), y(k - 2), u(k - 1)$ and $u(k - 2)$.

**Fig. 6.** A schematic representation of the system in the pH neutralization case study.
4.3. Case Study: Narendra

To evaluate how the FIMT-DD algorithm adapts to change, we consider the synthetic Narendra dataset [13]. This dynamical system is composed of an input variable \( u \) and a system variable \( y \), connected as follows

\[
y(k) = \frac{1}{1 + y(k-1)^2} + u(k-1)^3.
\]  

(4)

The value of the input variable \( u \) is selected uniformly at random from the interval \([-1, 1]\) after every 20 examples (time points) and kept constant for the next 20 examples. The initial system state is set as \( y(0) = 0 \). This system is modeled with a system lag of 1, which gives us two features \( y(k-1) \) and \( u(k-1) \) to model the system behavior.

Specifically to test the change detection and adaptation of FIMT-DD, we change the dynamics of the system. There are two types of changes that we introduce: change of the function and change of a parameter. In the former, the recurrence relation is replaced by

\[
\hat{y}(k) = \frac{1}{1 + \hat{y}(k-1)^2} + u(k-1),
\]  

(5)

replacing the exponent 3 in the second term with a value of 1, while in the latter, the relation changes into

\[
\hat{y}(k) = \frac{1}{1 + \hat{y}(k-1)^2} + 2u(k-1)^3,
\]  

(6)

replacing the constant 1 in front of the second term with the constant 2.

For this case study we generated datasets Narendra-F and Narendra-P, where the change is introduced in the function and in the parameter, respectively. Both types of changes are introduced half-way through the dataset. In total, 900000 instances were generated and in both cases the change was introduced at the 450000th time point.

When considering these two datasets, we compare two variants of the FIMT-DD algorithm, with an enabled and disabled change adaptation mechanism, respectively. In both cases, we use FIMT-DD to learn model trees.

5. Experimental Design

In this section, we first explicitly state the experimental questions that we set out to answer. We then list the on-line regression algorithms that we evaluate and compare, along with their parameter settings. The section concludes with the evaluation methodology.

5.1. Experimental Questions

In our comparison, we address four specific experimental questions. First, we assess whether the data stream mining approaches are appropriate for the identification of dynamical systems in discrete time. Additionally, we are interested in determining which of the compared algorithms performs the best. Afterwards, we compare the model produced by the best performed algorithm with an expert model in terms of simulation. Finally,
we wish to evaluate the effectiveness of the change detection mechanism of the on-line tree learning approach, which would be necessary for the identification of time-varying dynamical systems.

To answer the question of whether the considered approaches perform satisfactorily on the task of identification of dynamical systems, we need to verify that they make small, preferably ever-decreasing, errors. Low values of the error are desired that should be lower than those of a pre-selected baseline method. This means accurate predictions of the evolution of the state of the system, i.e., its state at the next point in time.

To evaluate which of the algorithms performs best, we compare them on two benchmark control engineering problems. The first one consists of measured data (gas-liquid separator case study in Sec. 4.1), while the second is synthetic (pH neutralization case study in Sec. 4.2). Our task is primarily to examine the predictive performance of the algorithms. In addition, we also inspect their time and memory consumption.

For the gas-liquid separator case study, we compare the simulation of the system dynamics obtained by using the models learned by data stream mining with the simulation produced by the expert model described in Sec. 4.1. This allows us to determine whether the learned model, which required almost zero effort to create, captures the dynamics of the system and how it compares to the expert model, which required a lot of effort to produce.

Specifically, we use the best performing algorithm identified above (ORTO-A), which learns on only the first run of the gas-liquid separator. Similarly, the parameter tuning of the expert model is performed on the first run. The second run of the gas-liquid separator is used for the simulation. For the learned model, the lagged output values are replaced by its earlier predictions, i.e., aside from the initial system state, the learner is agnostic of the recorded state of the system during the second run.

Finally, an important aspect of learning from data streams is the ability to detect changes in the data. In the context of system identification, this corresponds to the successful identification of time-varying dynamical systems. We investigate the performance of on-line learning of model trees with and without change detection on an artificial task where two types of changes are introduced to a known dynamical system (see Sec. 4.3).

5.2. Evaluated Algorithms

We compare and evaluate the on-line regression algorithms described in Sec. 3. In particular, we compare FIMT-DD using regression and model trees as well as the ORTO-A and ORTO-BT algorithms for learning option trees, without change detection, on the pH neutralization and gas-liquid separator case studies. For the Narendra case study, we consider only FIMT-DD with and without change detection.

We use a single linear regression model learned on-line as a baseline, i.e., a single perceptron (see Sec. 3.1). It can also be interpreted as a model tree with only leaf at the root, which is never allowed to grow. The parameter settings chosen for the evaluated algorithms are listed in Tab. 2.

5.3. Evaluation on Data Streams

The evaluation of accuracy of algorithms on data streams is not straightforward. In this paper, we use the predictive sequential or prequential approach, which introduces a
fading factor into the calculation of error metrics \( \text{3} \). Instead of reporting the average error \( E_N = \frac{1}{N} \sum_{i=1}^{N} L(y_i, \hat{y}_i) \), where \( L \) is some loss function, \( y_i \) is the real value of the \( i \)-th example and \( \hat{y}_i \) is the algorithm’s prediction on the \( i \)-th example, we introduce the faded sum of errors \( S_N \) and the faded number of examples \( B_N \) defined as follows:
\[
S_N = L(y_N, \hat{y}_N) + \alpha S_{N-1} = \sum_{i=1}^{N} \alpha^{N-i} L(y_i, \hat{y}_i), \\
B_N = 1 + \alpha B_{N-1} = \sum_{i=1}^{N} \alpha^{N-i},
\]
to compute the faded error \( E_N = \frac{S_N}{B_N} \). We use the parameter value \( \alpha = 0.975 \) [3].

If we use the square loss function \( L(y, \hat{y}) = (y - \hat{y})^2 \), this allows us to compute the faded mean squared error. Additionally, we use the relative root mean squared error (RRMSE), where we take the square root of the ratio of the mean squared error of the algorithm with the mean squared error of the linear regression model, which is used as a baseline:
\[
RRMSE = \sqrt{\frac{\text{MSE}_{\text{alg}}}{\text{MSE}_{\text{lr}}}}.
\]

6. Results and Discussion

In this section, we present the experimental results organized according to the experimental questions, posed in the previous section. We first discuss the question of the satisfactory performance of data stream mining approaches for modeling dynamical systems. Next, we discuss the relative performance of the different approaches to on-line regression. Afterwards, we focus on the comparison of two simulations obtained by a learned and a human constructed model. Finally, we discuss how the on-line learning of regression trees (with and without change detection) deals with time varying dynamical systems.

6.1. Comparison to the Baseline

In Tab. [3], we present the faded mean squared errors for the baseline method of linear regression and each of the four tree-based approaches for the gas-liquid separator domain. The error is recorded after each 5000 examples. The performance figures for the pH domain are given in Tab. [4].

The results clearly show that the tree-based methods for regression perform very well in terms of predictive error. For the case of the gas-liquid separator, the mean squared error rapidly approaches zero for all four tree-based approaches, as well as for the baseline. However, the error of the baseline is about an order of magnitude larger. For the pH
neutralization case, the error is getting close to zero for the tree-based approaches, but stays very high for the linear regression baseline approach.

The relative root mean square error essentially compares the error of the tree-based approaches to the error the baseline, the latter having the root relative root mean square error of 1. For the gas-liquid separator domain, the relative root mean squared error is clearly lower than 1 for all tree-based approaches (see Fig. 7a). For the pH neutralization case it is even lower (see Fig. 7b). To sum up, we can conclude that all tree-based approaches perform successfully at the task of system identification in discrete-time.

### 6.2. Relative Performance of the Tree-Based Approaches

In this section, we present the performance of the tree-based methods in terms of the (faded) root relative mean squared error with respect to the baseline (see Fig. 7), memory and time consumption (see Fig. 8 and 9), both for the gas-liquid separator case (marked with (a) in the figures) and the pH neutralization case (marked with (b) in the figures).

For the case of the FIMT-DD algorithm, we can see that FIMT-DD clearly performs better when learning model trees than when learning regression trees (see Fig. 7). Furthermore, the usage of model trees makes learning faster (in terms of achieving lower error with the same number of examples). In addition, model trees also achieve lower error overall than regression trees (for any given number of examples).

Regarding the option trees, the ORTO-A algorithm clearly outperforms the ORTO-BT algorithm. While the option trees learned for both ORTO-A and ORTO-BT are the same, the aggregation by averaging yields more accurate predictions. In addition, ORTO-A is also less prone to fluctuations in the error.

### Table 3. The faded mean squared errors of the different modeling approaches on the gas-liquid separator dataset, recorded at intervals of 5000 examples.

<table>
<thead>
<tr>
<th># of examples</th>
<th>5000</th>
<th>10000</th>
<th>15000</th>
<th>20000</th>
<th>25000</th>
<th>30000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>0.0084</td>
<td>0.0051</td>
<td>0.0253</td>
<td>0.0117</td>
<td>0.0136</td>
<td></td>
</tr>
<tr>
<td>FIMT-DD Reg. trees</td>
<td>0.0064</td>
<td>0.0018</td>
<td>0.0050</td>
<td>0.0030</td>
<td>0.0011</td>
<td></td>
</tr>
<tr>
<td>FIMT-DD Model trees</td>
<td>0.0040</td>
<td>0.0007</td>
<td>0.0006</td>
<td>0.0016</td>
<td>0.0003</td>
<td></td>
</tr>
<tr>
<td>ORTO-A</td>
<td>0.0008</td>
<td>0.0002</td>
<td>0.0021</td>
<td>0.0002</td>
<td>0.0005</td>
<td></td>
</tr>
<tr>
<td>ORTO-BT</td>
<td>0.0022</td>
<td>0.0002</td>
<td>0.0018</td>
<td>0.0011</td>
<td>0.0018</td>
<td></td>
</tr>
</tbody>
</table>

### Table 4. The faded mean squared errors of the different modeling approaches on the pH dataset, recorded at intervals of 5000 examples.

<table>
<thead>
<tr>
<th># of examples</th>
<th>5000</th>
<th>10000</th>
<th>15000</th>
<th>20000</th>
<th>25000</th>
<th>30000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>3.0473</td>
<td>3.9209</td>
<td>4.1793</td>
<td>3.1796</td>
<td>3.8867</td>
<td>3.3447</td>
</tr>
<tr>
<td>FIMT-DD Reg. trees</td>
<td>0.9071</td>
<td>0.0606</td>
<td>0.0438</td>
<td>0.0343</td>
<td>0.0281</td>
<td>0.0195</td>
</tr>
<tr>
<td>FIMT-DD Model trees</td>
<td>0.4405</td>
<td>0.0745</td>
<td>0.0390</td>
<td>0.0313</td>
<td>0.0375</td>
<td>0.0198</td>
</tr>
<tr>
<td>ORTO-A</td>
<td>0.0733</td>
<td>0.0251</td>
<td>0.0252</td>
<td>0.0215</td>
<td>0.0150</td>
<td>0.0167</td>
</tr>
<tr>
<td>ORTO-BT</td>
<td>0.3576</td>
<td>0.0343</td>
<td>0.0402</td>
<td>0.0396</td>
<td>0.0219</td>
<td>0.0151</td>
</tr>
</tbody>
</table>
If we analyze the overall performance, the ORTO-A algorithm is the best performing algorithm. It consistently outperforms all of the other algorithms in terms of the error. In addition, it is the fastest among the algorithms to achieve low errors. These results hold both for the gas-liquid separator case, as well as the pH neutralization case. This is to be expected, as the ORTO algorithm was designed to overcome the problem of initial delay in learning in a streaming setting and to increase the predictive accuracy faster [10].

If we analyze the performance in terms of time and memory used, we can observe that the ORTO-A and the ORTO-BT algorithms are the most resource intensive (see Fig. 8 and 9). However, the ORTO-BT algorithm uses a fractionally smaller part of the tree for prediction than the ORTO-A algorithm. Using model trees rather than regression trees, for the FIMT-DD algorithm, increases the time consumption slightly, while the memory consumption barely changes.
6.3. Comparison to the Simulated Model

Here we use ORTO-A, which we identified as the best performing method. Here, we compare its predictions to the recorded values, as well as the values obtained by simulating the expert-derived non-linear dynamic model (see Sec. 4.1). The simulation was performed in Simulink using the ODE3 (Bogacki-Shampine) fixed step solver.

The results of the comparison are presented in Fig. 10. It is clear that the model learned by ORTO-A captures the dynamics of the system. In the first half of the simulation, the pressure simulated by the learned model is often off of the real pressure. This may be due to the differences between the two runs of the gas-liquid separator.

While the simulation of the expert-derived model better captures the dynamics of the system, it should be noted that the construction of this model required a lot of human resources. Our method, on the other hand, was used out of the box, without any parameter tuning, requiring minimal additional resources, both in terms of human effort and computation time.

6.4. Change Detection

In Fig. 11, we show the performance of the FIMT-DD algorithm in terms of the faded mean squared error on the Narendra-F and Narendra-P datasets, from the Narendra case. In the former, a functional form of the dynamics is changed, while in the latter a parameter of the dynamics changes, to simulate a time-varying dynamical system (see Sec. 4.3).

Analyzing the performance of the Narendra-F dataset, we notice that the FIMT-DD algorithm with change detection does detect the change, builds an alternate subtree and replaces the original in about 25000 examples, as the change is introduced at the 450000-th example and the predictive accuracy increases at about the 475000th example (see Fig. 11a). However, the difference in the error between the FIMT-DD with and without change is small. The increase in error due to the abrupt change in the dynamics, which is in this case also relatively small, peaks at about 0.08.

The change introduced in the Narendra-P dataset, causes a larger increase in error, peaking at about 0.15 (see Fig. 11b). This is expected, as in this case the change doubled
Fig. 10. The recorded values of the system variable (pressure) and its values simulated by the model learned by ORTO-A and the constructed expert model of the pressure in the gas-liquid separator case study.

the range of the second term to $[-2, 2]$, while the change introduced in the Narendra-F dataset only changed its distribution on the $[-1, 1]$ interval. As before, the change is detected and the tree is adapted in about 25000 examples. Because the introduced change is more radical than before, regrowing of the affected subtree(s) produces better results.

7. Conclusions and Further Work

The central question addressed by this paper concerns the applicability of the paradigm of mining data streams, and in particular the tree-based approaches to on-line regression, to the task of system identification in discrete-time. To address this question, we perform a thorough experimental evaluation, where we applied several tree-based approaches for on-line regression to several benchmark problems in discrete-time modeling of dynamical

Fig. 11. The performance (faded MSE) of FIMT-DD without and with change detection.
systems. Using an evaluation methodology and performance measures appropriate for the task at hand, we answer several experimental questions, as summarized below.

Our conclusions are as follows. Overall, tree-based approaches to on-line regression are appropriate for solving the task of identification of dynamical systems in discrete-time. Among the considered approaches, the ORTO-A algorithm clearly stands out as the best-performing one. ORTO-A learns option trees for regression, where an example may be sorted down multiple branches of the tree in option nodes, and averages the predictions obtained from each of the branches in the option tree that an example follows.

When comparing the learned model to an expert constructed model, we noted that the expert model produces a simulation closer to the recorded data. However, the learned model still captures the dynamics of the system quite well. Furthermore, the learned model required minimal additional use of resources, human and machine, while the construction of the expert model and its parameter tuning is a labor intensive and error-prone process.

We also investigate the use of tree-based regression algorithms on data streams for the identification of time-varying systems. The FIMT-DD algorithm for learning model trees is applied to synthetic data, which include a change of system dynamics at a given point in time. The change detection mechanism and the adaptation method in the FIMT-DD algorithm successfully detect the introduced change and adapt the tree to achieve better performance (as compared to the FIMT-DD algorithm without change detection).

Many avenues remain for further work. Specifically, the change detection and adaptation mechanisms of tree-based on-line learning approaches should be tested on other real-world datasets, where change is either known or suspected to appear. Furthermore, the mechanism for detecting change, i.e., the Page-Hinckley test, should be automatically parameterized with regard to the relative error instead of the absolute error, as this would allow to detect changes of any size without the need to select the parameters manually.

More generally, the experimental questions posed in this paper should be answered for a larger set of real-world problem domains and corresponding datasets. Of particular interest is the identification of dynamical systems which have more than one system variable: These are called MIMO (multiple input multiple output systems). Additional on-line learning methods should also be considered for the task at hand, such as tree-based on-line ensemble approaches, including on-line methods for learning multi-target regression and model trees on data streams [8].

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References


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