

# **MATLAB Software for Recursive Identification and Scaling Using a Structured Nonlinear Black-box Model – Revision 6**

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## **Abstract**

This report is intended as a users manual for a package of MATLAB™ scripts and functions, developed for recursive prediction error identification of nonlinear state space systems and nonlinear static systems. The core of the package is an implementation of related output error identification and scaling algorithms. The algorithms are based on a continuous time, structured black box state space model of a nonlinear system. Furthermore, to initialize the algorithm an initiation scheme based on Kalman filter theory is included. The purpose of the initialization algorithm is to find initial parameters for the prediction error algorithm, and thus reducing the risk of convergence to local false minima. An RPEM algorithm for recursive identification of nonlinear static systems, that re-uses the parameterization of the nonlinear ODE model, is also included in the software package. In this version of the software a new discretization of the continuous time model based on the midpoint integration algorithm is added. The software can only be run off-line, i.e. no true real time operation is possible. The algorithms are however implemented so that true on-line operation can be obtained by extraction of the main algorithmic loop. The user must then provide the real time environment. The software package contains scripts and functions that allow the user to either input live measurements or to generate test data by simulation. The scripts and functions for the setup and execution of the identification algorithms are somewhat more general than what is described in the references. There is e.g. support for automatic re-initiation of the algorithms using the parameters obtained at the end of a previous identification run. This allows for multiple runs through a set of data, something that is useful for data sets that are too short to allow convergence in a single run. The re-initiation step also allows the user to modify the degrees of the polynomial model structure and to specify terms that are to be excluded from the model. This makes it possible to iteratively re-fine the estimated model using multiple runs. The functionality for display of results include scripts for plotting of data, parameters, prediction errors, eigenvalues and the condition number of the Hessian. The estimated model obtained at the end of a run can be simulated and the model output plotted, alone or together with the data used for identification. Model validation is supported by two methods apart from the display functionality.

First, a calculation of the RPEM loss function can be performed, using parameters obtained at the end of an identification run. Secondly, the accuracy as a function of the output signal amplitude can be assessed.

Keywords: Identification, Kalman filter, MATLAB™, Nonlinear systems, Ordinary differential equation, RPEM, Recursive algorithm, Sampling, Scaling, Software, State space model.

### **Prerequisites**

This report only describes the parts of [ 1 ], [ 2 ], [ 3 ], [ 4 ], [ 5 ], [ 6 ] and [12] that are required for the description of the software. Hence the user is assumed to have a working knowledge of the algorithm of these publications and of MATLAB™, see e.g. [ 7 ]. This, in turn, requires that the user has a working knowledge of system identification and in particular of recursive identification methods as described in e.g. [ 8 ]. The algorithm for identification of static systems is described in some detail in the report.

### **Revisions**

Revision 1: [ 10 ] describes revision 1.0 of the accompanying SW. The software of revision 1 has been tested with MATLAB 5.3, MATLAB 6.5, and MATLAB 7.0 running on PCs and UNIX workstations.

Revision 2: [ 11 ] includes functionality for recursive identification of static nonlinear systems. See sections 9 and 10 of the report. Furthermore, an error has been corrected in the RPEM algorithm. The timing error of one sample in the output equation of the RPEM affected identification results slightly in case an explicit dependence of input signals was used in the output equation.

Revision 3: This revision adds a recursive algorithm for initialization of the dynamic prediction error algorithm. The new algorithm is based on Kalman filter theory, for details see sections 11 and 12.

Revision 4: This revision corrects a bug that led to erroneous behaviour of the initialization algorithm, for model orders higher than 2. The bug was due to a missed scaling in the differentiation part of the code. The references [ 5 ] and [ 6 ] are updated accordingly.

Revision 5: This revision includes the use of midpoint algorithm as a discretization algorithm.

Revision 6: This revision corrects an error associated with scaling, introduced in the previous revision.

## **Installation**

The file SW.zip is copied to the selected directory and unzipped. MATLAB is opened and a path is set up to the selected directory using the path browser. The software is then ready for use.

Note: This report is written with respect to the software, as included in the SW.zip file. It may therefore be advantageous to store the originally supplied software for reference purposes.

## **Error reports**

When errors are found, these may be reported in an e-mail to:

[torbjorn.wigren@it.uu.se](mailto:torbjorn.wigren@it.uu.se).

## **1. Introduction**

Identification of nonlinear systems is an active field of research today. There are several reasons for this. First, many practical systems show strong nonlinear effects. This is e.g. true for high angle of attack flight dynamics, many chemical reactions and electromechanical machinery of many kinds, see e.g. [ 1 ], [ 2 ], [ 3 ], [ 4 ], [5], [6] and the references therein for further examples. Another important reason is perhaps that linear methods for system identification are quite well understood today, hence it is natural to move the focus to more challenging problems.

There are already a number of identification methods available for identification of nonlinear systems. These include grey-box differential equation methods, where numerical integration is combined with optimization in order to optimize the unknown physical parameters that appear in the differential equations. An alternative approach is to start with a discrete time black box model, and to apply existing methodology from the linear field to the solution of the nonlinear identification problem. This is the approach taken in the NARMAX method and its related algorithms. There, a least squares formulation can often be found, a fact that facilitates the solution. Other methods apply neural networks for modeling of nonlinear dynamic systems. See [ 1 ], [ 2 ] and the references therein for a more detailed survey.

This report focuses on software that implements new nonlinear recursive system identification methods. Contrary to the above methods, this black box method estimates continuous time parameters in a general state space model, with a known and possibly nonlinear measurement equation. The main identification methods belong to the class of recursive prediction error methods (RPEMs) and the methods are of output error type. Advantages include the fact that the stability of the estimated model is checked by a projection algorithm, at each iteration step. The least squares approaches above cannot guarantee a resulting stable model - this needs to be checked after the identification has been

completed. A further advantage is that the connection to the physical parameters can be retained to a greater extent than if a discrete time nonlinear model is used as the starting point. There are also disadvantages. A major disadvantage with output error methods is that they sometimes converge to a local sub-optimal minimum point of the criterion function, meaning that careful initialization is needed. The effect of local minimum points is reduced for the method described in [ 1 ] – [ 4 ], by a method that scales the states, the estimated parameters of the model and, most importantly, the Hessian of the criterion function. The scaling is implemented by a scaling of the sampling period used when running the identification algorithm, see [ 1 ] – [ 3 ] and [ 12 ] for details. One important aspect of this scaling method is that corresponding un-scaled parameter values can be calculated in a post-identification step.

The nonlinear identification algorithms are based on a continuous time black box state space model. This model is structured in that only one right hand side component of the ordinary differential equation (ODE) model is parameterized as an unknown function. As shown in [ 1 ] and [ 2 ] this avoids over-parameterization. The restriction imposed on the model structure may seem restrictive. However, it is motivated in [ 1 ] and [ 2 ] that the selected structure can always (locally in the states) model systems with more general right hand sides, a fact that extends the applicability of the method significantly. The selected parameterization of the right hand side function of the ODE is a linear-in-the-parameters multi-variate polynomial in the states and input signals. The approach taken allows for MIMO nonlinear system identification. The covariance matrix of the measurement disturbances is estimated on-line.

The second revision of the software package adds an RPEM algorithm for recursive identification of nonlinear static systems. The algorithm re-uses the parameterization of the nonlinear function used in the RPEM for identification of nonlinear dynamic systems described above. Most of the code of the SW package has been re-used, however a number of scripts have been modified and appear in two versions. This is marked by the inclusion of “Static” in one of the duplicated m-files. All other m-files can be used as described for the RPEM for identification of nonlinear dynamics. Note that scaling is not applicable in the static case. The static algorithm is described in sections 9 and 10.

The third revision adds a Kalman filter based algorithm to the software package. The new algorithm is based on the same model structure as the RPEM for identification of nonlinear dynamic systems, and is intended to provide initial parameters for the RPEM. Most parts of the scripts and functions of the previous versions of the SW package can be used in combination with the new algorithm.

The fourth revision corrects a problem in the previous revision, associated with the differentiation of the measurements.

The fifth and sixth revisions includes the use of the midpoint integration method as another discretization algorithm applied on the continuous time black box state space model. All of the

previous functions and scripts can be used with the new algorithm. The sixth edition corrects an error associated with the scaling that appeared in the fifth edition.

Recursive system identification is a software dependent technology. Hence, when publishing new methodology in this field, it is relevant to also provide useful software for application of the presented algorithms. This facilitates a quick practical exploitation of new ideas. The development of the present MATLAB™ software package is motivated by this fact.

The present software package is developed and tested using MATLAB 5.3, MATLAB 6.5, MATLAB 7.0 and MATLAB 7.9. The software package does not rely on any MATLAB toolboxes. It consists of a number of scripts and functions. Briefly, the software package consists of scripts for setup, scripts for generation or measurement of data, scripts for execution of the RPEM and scripts for generation and plotting of results. There is presently no GUI, the scripts must be run from the command window. Furthermore, input parameters need to be configured in one or several of the setup scripts, as well as when running the scripts. In case of data generation by simulation, the ODE that defines the data generating system must be specified in standard MATLAB style. The software can only be run off-line, i.e. there is no support for execution in a real time environment. The major parts of the algorithmic loop can however easily be extracted for such purposes.

The report is organized according to the flow of tasks a user encounters when applying the scripts of the package. A detailed description of the software is given for the nonlinear dynamic case, the static case is described more briefly in the end of this report. Before the software is described some basic facts about the ODE model and the scaling method are reviewed.

## 2. Model – ODE case

The nonlinear MIMO model to be defined here is used for estimation of an unknown parameter vector  $\theta$  from measured inputs  $\mathbf{u}(t)$  and outputs  $\mathbf{y}_m(t)$ , given by

$$\begin{aligned} \mathbf{u}(t) &= \left( u_1(t) \quad \dots \quad u_1^{(n_1)}(t) \quad \dots \quad u_k(t) \quad \dots \quad u_k^{(n_k)}(t) \right)^T \\ \mathbf{y}_m(t) &= \left( y_{m,1}(t) \quad \dots \quad y_{m,p}(t) \right)^T \end{aligned} \quad (1)$$

The superscript  $^{(k)}$  denotes differentiation  $k$  times. The starting point for the derivation of the model is the following  $n$ :th order state space ODE

$$\begin{pmatrix} \dot{x}_1^{(1)} \\ \vdots \\ \dot{x}_{n-1}^{(1)} \\ \dot{x}_n^{(1)} \end{pmatrix} = \begin{pmatrix} x_2 \\ \vdots \\ x_n \\ f(x_1, \dots, x_n, u_1, \dots, u_1^{(n_1)}, \dots, u_k, \dots, u_k^{(n_k)}, \theta) \end{pmatrix} \quad (2)$$

$$\begin{pmatrix} y_1 \\ \vdots \\ y_p \end{pmatrix} = \begin{pmatrix} c_{11} & \dots & c_{1n} \\ \vdots & \ddots & \vdots \\ c_{p1} & \dots & c_{pn} \end{pmatrix} \begin{pmatrix} x_1 \\ \vdots \\ x_n \end{pmatrix},$$

where  $\mathbf{x} = (x_1 \dots x_{n-1} x_n)^T$  is the state vector. The following polynomial parameterization of the right hand side function of (2) is used

$$\begin{aligned} & f(x_1, \dots, x_n, u_1, \dots, u_1^{(n_1)}, \dots, u_k, \dots, u_k^{(n_k)}, \boldsymbol{\theta}) \\ &= \sum_{i_{x_1}=0}^{I_{x_1}} \dots \sum_{i_{x_n}=0}^{I_{x_n}} \sum_{i_{u_1}=0}^{I_{u_1}} \dots \sum_{i_{u_k}=0}^{I_{u_k}} \sum_{i_{u_k}=0}^{I_{u_k}} \theta_{i_{x_1} \dots i_{x_n} i_{u_1} \dots i_{u_k} \dots i_{u_k}} (x_1)^{i_{x_1}} \dots (x_n)^{i_{x_n}} (u_1)^{i_{u_1}} \dots \\ & \dots (u_1^{(n_1)})^{i_{u_1}^{(n_1)}} \dots (u_k)^{i_{u_k}} \dots (u_k^{(n_k)})^{i_{u_k}^{(n_k)}} = \boldsymbol{\varphi}^T(\mathbf{x}, \mathbf{u}) \boldsymbol{\theta} \end{aligned} \quad (3)$$

Here

$$\begin{aligned} \boldsymbol{\theta} &= \left( \theta_{0 \dots 0} \quad \dots \quad \theta_{0 \dots I_{u_k^{(n_k)}}} \quad \theta_{0 \dots 010} \quad \dots \quad \theta_{0 \dots 01 I_{u_k^{(n_k)}}} \quad \dots \right. \\ & \quad \left. \theta_{0 \dots 011} \quad \dots \quad \theta_{0 \dots 01 I_{u_k^{(n_k)}}} \quad \dots \quad \theta_{0 \dots 011 I_{u_k^{(n_k)}}} \quad \dots \right)^T \\ \boldsymbol{\varphi} &= \left( 1 \quad \dots \quad \left( (u_k^{(n_k)})^{I_{u_k^{(n_k)}}} \right) \quad u_k^{(n_k-1)} \quad \dots \quad \left( (u_k^{(n_k-1)})^{I_{u_k^{(n_k)}}} \right) \dots \quad (u_k^{(n_k-1)})^{I_{u_k^{(n_k-1)}}} \quad \dots \right. \\ & \quad \left. \left( (u_k^{(n_k-1)})^{I_{u_k^{(n_k-1)}}} \right) \left( (u_k^{(n_k-1)})^{I_{u_k^{(n_k-1)}}} \right) \dots \left( (x_1)^{I_{x_1}} \dots (x_n)^{I_{x_n}} (u_1)^{I_{u_1}} \dots (u_k^{(n_k)})^{I_{u_k^{(n_k)}}} \right)^T \right) \end{aligned} \quad (4)$$

Please see example 5 below for a low order example of the above parameterization. In order to obtain a discrete time model that is suitable for scaling two different discretization algorithms have been applied to (2), the Euler method and the midpoint method.

### 2.1 Euler integration method

The main reason for using the Euler method is that the sampling appears explicitly and linearly in the right hand side of the resulting difference equation model (5). This is convenient when the scaling algorithm is introduced. The result of the discretization is

$$\begin{pmatrix} x_1(t + T_s, \theta) \\ \vdots \\ x_{n-1}(t + T_s, \theta) \\ x_n(t + T_s, \theta) \end{pmatrix} = \begin{pmatrix} x_1(t, \theta) \\ \vdots \\ x_{n-1}(t, \theta) \\ x_n(t, \theta) \end{pmatrix} + T_s \begin{pmatrix} x_1'(t, \theta) \\ \vdots \\ x_{n-1}'(t, \theta) \\ x_n'(t, \theta) \end{pmatrix} \quad (5)$$

$$\begin{pmatrix} y_1(t, \theta) \\ \vdots \\ y_p(t, \theta) \end{pmatrix} = \begin{pmatrix} c_{11} & \dots & c_{1n} \\ \vdots & \ddots & \vdots \\ c_{p1} & \dots & c_{pn} \end{pmatrix} \begin{pmatrix} x_1(t, \theta) \\ \vdots \\ x_n(t, \theta) \end{pmatrix} = \mathbf{C}\mathbf{x}(t, \theta) \quad (6)$$

It can be remarked that the Euler method may require fast sampling in order not to introduce significant discretization errors. This is fortunately a less important effect in system identification applications. The reason is that the minimization algorithm uses the parameters as instruments to fit the model output to the measured data, as expressed by the criterion function. Even if an additional bias would be introduced in the estimated parameters, the input-output properties of the identified model can be expected to describe the data well.

## 2.2 Midpoint integration method

The purpose of using the midpoint integration rule is mainly the higher accuracy of the algorithm compared to the Euler method giving more accurate parameter estimates. The results of the discretization of (2) is:

$$\begin{pmatrix} x_1(t + T_s, \theta) \\ \vdots \\ x_{n-1}(t + T_s, \theta) \\ x_n(t + T_s, \theta) \end{pmatrix} = \begin{pmatrix} x_1(t, \theta) \\ \vdots \\ x_{n-1}(t, \theta) \\ x_n(t, \theta) \end{pmatrix} \quad (7)$$

$$+ T_s \begin{pmatrix} x_1'(t, \theta) + \frac{T_s}{2} x_1''(t, \theta) \\ \vdots \\ x_n'(t, \theta) + \frac{T_s}{2} x_n''(t, \theta) \end{pmatrix} = \begin{pmatrix} x_1(t, \theta) + \frac{T_s}{2} \varphi^T \left( x_1(t, \theta), \dots, x_n(t, \theta), u_1(t), \dots, u_{n_k}(t) \right) \theta \\ \vdots \\ x_n(t, \theta) + \frac{T_s}{2} \varphi^T \left( x_1(t, \theta), \dots, x_n(t, \theta), u_1\left(t + \frac{T_s}{2}\right), \dots, u_{n_k}\left(t + \frac{T_s}{2}\right) \right) \theta \end{pmatrix}$$

$$\begin{pmatrix} y_1(t, \theta) \\ \vdots \\ y_p(t, \theta) \end{pmatrix} = \begin{pmatrix} c_{11} & \dots & c_{1n} \\ \vdots & \ddots & \vdots \\ c_{p1} & \dots & c_{pn} \end{pmatrix} \begin{pmatrix} x_1(t, \theta) \\ \vdots \\ x_n(t, \theta) \end{pmatrix} = \mathbf{C}\mathbf{x}(t, \theta) \quad (8)$$

where

$$\bar{x}_i(t, \theta) = x_i(t, \theta) + \frac{T_s}{2} \varphi^T \left( x_1(t, \theta), \dots, x_n(t, \theta), u_1\left(t + \frac{T_s}{2}\right), \dots, u_{n_k}\left(t + \frac{T_s}{2}\right) \right) \theta, \quad (9)$$

is used in the right hand side of the model (7).

### 3. Scaling – ODE case

#### 3.1 Euler method

During development of the RPEM described in [ 1 ] - [ 4 ] , it was noticed that problems with convergence to false local minimum points of the criterion were often highly related to the selection of the sampling period. The sampling period of course needs to be short enough during measurement, in order to capture the essential dynamics of the identified system. Hence the measurement sampling period cannot be arbitrarily selected. However, since the sampling period appears *explicitly* in the model ( 5 ) and in the corresponding gradient difference equation, *it is straightforward to apply identification algorithms based on ( 5 ) with another, scaled value of the sampling period*. This idea affects the updating of the states, the gradient and any projection algorithm that is used to control the stability of the model. A scale factor  $\alpha$  appears before the multiplication with the sampling period  $T_s$  in those three quantities. To explain the details, the scale factor  $\alpha$  and the scaled sampling period  $T_s^{Scaled}$  are first defined as

$$T_s^{Scaled} = \alpha T_s. \quad (10)$$

The model ( 5 ), ( 6 ) , as applied in the identification algorithm is then transformed into

$$\begin{pmatrix} x_1^s(t + T_s, \theta^s) \\ \vdots \\ x_{n-1}^s(t + T_s, \theta^s) \\ x_n^s(t + T_s, \theta^s) \end{pmatrix} = \begin{pmatrix} x_1^s(t, \theta^s) \\ \vdots \\ x_{n-1}^s(t, \theta^s) \\ x_n^s(t, \theta^s) \end{pmatrix} + T_s^{Scaled} \begin{pmatrix} f_1(x_1^s(t, \theta^s), \dots, x_n^s(t, \theta^s), u_1(t), \dots, u_k(t), \theta^s) \\ \vdots \\ f_n(x_1^s(t, \theta^s), \dots, x_n^s(t, \theta^s), u_1(t), \dots, u_k(t), \theta^s) \end{pmatrix} \quad (11)$$

$$\begin{pmatrix} y_1^s(t, \theta^s) \\ \vdots \\ y_p^s(t, \theta^s) \end{pmatrix} = \begin{pmatrix} c_{11} & \dots & c_{1n} \\ \vdots & \ddots & \vdots \\ c_{p1} & \dots & c_{pn} \end{pmatrix} \begin{pmatrix} x_1^s(t, \theta^s) \\ \vdots \\ x_n^s(t, \theta^s) \end{pmatrix} = \mathbf{C} \mathbf{x}^s(t, \theta^s) \quad (12)$$

where the superscript  $^s$  denotes scaled quantities. Note that the original sampling period must be retained in all time arguments, so as to refer to the correct measurement times. The gradient follows by differentiation of ( 11 ) and ( 12 )

$$\frac{d\mathbf{x}^s(t + T_s, \theta^s)}{d\theta^s} = \frac{d\mathbf{x}^s(t, \theta^s)}{d\theta^s} \quad (13)$$



$$+ T_S^{Scaled} \begin{pmatrix} \frac{dx_2^s(t, \theta^s)}{d\theta^s} \\ \vdots \\ \frac{dx_n^s(t, \theta^s)}{d\theta^s} \\ \varphi^T(\mathbf{x}^s(t, \theta^s), \mathbf{u}(t)) \end{pmatrix} + T_S^{Scaled} \begin{pmatrix} 0 \\ \vdots \\ 0 \\ \boldsymbol{\theta}^T \left( \frac{d\boldsymbol{\varphi}(\mathbf{x}^s(t, \boldsymbol{\theta}^s), \mathbf{u}(t))}{d\mathbf{x}^s} \right) \left( \frac{d\mathbf{x}^s(t, \boldsymbol{\theta}^s)}{d\boldsymbol{\theta}^s} \right) \end{pmatrix}$$

$$(\boldsymbol{\psi}^s(t, \boldsymbol{\theta}^s))^T = \frac{d\mathbf{y}^s(t, \boldsymbol{\theta}^s)}{d\boldsymbol{\theta}^s} = \mathbf{C} \frac{d\mathbf{x}^s(t, \boldsymbol{\theta}^s)}{d\boldsymbol{\theta}^s}. \quad (14)$$

Note that the above change from  $\boldsymbol{\theta}$  to  $\boldsymbol{\theta}^s$  is *not* to be treated as a change of variables in the differentiation leading to (13) and (14). The originally derived gradient is applied, but with a scaled sampling period. The last affected quantity of the RPEM algorithm is the projection algorithm that is [1]

$$\mathbf{S}^s(\boldsymbol{\theta}^s) = \mathbf{I}_n + T_S^{Scaled} \begin{pmatrix} 0 & 1 & 0 & \cdots & 0 \\ 0 & 0 & 1 & \ddots & \vdots \\ \vdots & \vdots & \ddots & \ddots & 0 \\ 0 & 0 & \cdots & 0 & 1 \\ (\boldsymbol{\theta}^s)^T \frac{d\boldsymbol{\varphi}(\mathbf{x}^s(t, \boldsymbol{\theta}^s))}{d\mathbf{x}^s} \end{pmatrix} \quad (15)$$

$$D_M = \left\{ \boldsymbol{\theta}^s \mid \left| \text{eig}(\mathbf{S}^s(\boldsymbol{\theta}^s)) \right| < 1 - \delta \right\}, \delta > 0 \quad (16)$$

$$\left[ \hat{\boldsymbol{\theta}}^s(t) \right]_{\mathbf{b}_M} = \begin{cases} \hat{\boldsymbol{\theta}}^s(t) & \hat{\boldsymbol{\theta}}^s(t) \in D_M \\ \hat{\boldsymbol{\theta}}^s(t - T_S) & \hat{\boldsymbol{\theta}}^s(t) \notin D_M \end{cases} \quad (17)$$

In (15),  $\mathbf{S}^s(\boldsymbol{\theta}^s)$  denotes the linearized system matrix of the model,  $D_M$  denotes the model set, here defined as the asymptotically stable models with a margin  $\delta$  to the stability limit. The last equation stops the updating of the parameter vector in case the update would result in values outside the model set. Other details of an RPEM where the scaling algorithm is used can be found in [1] – [4].

### 3.2 Midpoint method

The scaling of the sampling parallels the previous section, where the sampling  $T_S$  is scaled according to (7). Applying the scaled sampling to the discretized model (7) – (8) generates the following result:

$$\begin{pmatrix} x_1^s(t+T_s, \theta^s) \\ \vdots \\ x_{n-1}^s(t+T_s, \theta^s) \\ x_n^s(t+T_s, \theta^s) \end{pmatrix} = \begin{pmatrix} x_1^s(t, \theta^s) \\ \vdots \\ x_{n-1}^s(t, \theta^s) \\ x_n^s(t, \theta^s) \end{pmatrix} \quad (18)$$

$$+T_s^{-1} \begin{pmatrix} x_1^s(t, \theta^s) - T_s^{-1} x_1^s(t, \theta^s) \\ \vdots \\ x_{n-1}^s(t, \theta^s) - T_s^{-1} x_{n-1}^s(t, \theta^s) \\ \psi(x_1^s(t, \theta^s), \dots, x_n^s(t, \theta^s), u_1, \dots, u_k, t) \theta^s \\ \psi(x_1^s(t, \theta^s), \dots, x_n^s(t, \theta^s), u_1(t + \frac{T_s^{-1}}{2}), \dots, u_k(t + \frac{T_s^{-1}}{2})) \theta^s \end{pmatrix}$$

$$\begin{pmatrix} y_1^s(t, \theta^s) \\ \vdots \\ y_p^s(t, \theta^s) \end{pmatrix} = \begin{pmatrix} c_{11} & \dots & c_{1n} \\ \vdots & \ddots & \vdots \\ c_{p1} & \dots & c_{pn} \end{pmatrix} \begin{pmatrix} x_1^s(t, \theta^s) \\ \vdots \\ x_n^s(t, \theta^s) \end{pmatrix} = \mathbf{C} \mathbf{x}^s(t, \theta^s) \quad (19)$$

where the superscript  $^s$  denotes scaled quantities, and  $\bar{x}_1^s$  is the scaled version of (9). The gradient for the new algorithm is then:

$$\frac{d\mathbf{x}^s(t+T_s, \theta^s)}{d\theta^s} = \frac{d\mathbf{x}^s(t, \theta^s)}{d\theta^s} + T_s^{-1} \begin{pmatrix} \frac{dx_1^s(t, \theta^s)}{d\theta^s} - T_s^{-1} \frac{dx_1^s(t, \theta^s)}{d\theta^s} \\ \vdots \\ \frac{dx_{n-1}^s(t, \theta^s)}{d\theta^s} + T_s^{-1} \frac{df(x_1^s, \dots, x_n^s, u_1, \dots, u_k, t, \theta^s)}{d\theta^s} \\ \frac{df(x_1^s, \dots, x_n^s, u_1(t + \frac{T_s^{-1}}{2}), \dots, u_k(t + \frac{T_s^{-1}}{2}), \theta^s)}{d\theta^s} \end{pmatrix} \quad (20)$$

where

$$\frac{df(\bar{x}_1^s, \dots, \bar{x}_n^s, u_1, \dots, u_k^{(n_k)}, \theta)}{d\theta} = \frac{\partial f}{\partial \bar{\mathbf{x}}} \bigg|_{t+\frac{T_s^{\text{Scaled}}}{2}} \frac{d\bar{\mathbf{x}}^s}{d\theta^s} + \frac{\partial f}{\partial \theta^s} \bigg|_{t+\frac{T_s^{\text{Scaled}}}{2}} \quad (21)$$

$$\frac{d\bar{\mathbf{x}}^s}{d\boldsymbol{\theta}^s} = \frac{d\mathbf{x}^s(t, \boldsymbol{\theta}^s)}{d\boldsymbol{\theta}^s} + \boldsymbol{\varphi}(\mathbf{x}^s(t, \boldsymbol{\theta}^s), u(t)) + \frac{T_s^{Scaled}}{2} \begin{pmatrix} 0 & 1 & 0 & \dots & 0 \\ 0 & 0 & 1 & \ddots & \vdots \\ \vdots & \vdots & \ddots & \ddots & 0 \\ 0 & 0 & \dots & 0 & 1 \end{pmatrix} \quad (22)$$

$$\frac{\partial f}{\partial \boldsymbol{\theta}^s} = \boldsymbol{\varphi}(\mathbf{x}^s(t, \boldsymbol{\theta}^s), \mathbf{u}(t)) \quad (23)$$

$$\frac{\partial f}{\partial \mathbf{x}^s} = (\boldsymbol{\theta}^s)^T \frac{d\boldsymbol{\varphi}(\mathbf{x}^s(t, \boldsymbol{\theta}^s))}{d\mathbf{x}^s(t, \boldsymbol{\theta}^s)} \frac{d\mathbf{x}^s(t, \boldsymbol{\theta}^s)}{d\boldsymbol{\theta}^s} \quad (24)$$

When a scaled value of the sampling period is applied, the algorithm still attempts to minimize the criterion, thereby obtaining other minimizing parameter values than when the true sampling period is used. When testing the scaling algorithm experimentally, dramatic improvements were sometimes observed in the algorithmic behavior. Convergence speeds could be improved and initial values that lead to divergence and instability could be made to work well.

The application of the scaling algorithm results in other estimated parameter values than what would be obtained without scaling. Fortunately, as shown in [ 2 ] for the algorithm based on Euler discretization, the original parameters can be calculated from the estimated ones. The transformation is given by a diagonal transformation matrix that is a function of the applied scale factor. The results for the algorithm based on midpoint discretization algorithm appear in [ 12 ]. The analysis of the effect of the scaling is continued in [ 3, where the effect of the conditioning of the Hessian of the criterion function is analyzed in detail. This shows that the effect of the scaling is quite dramatic. Changes of the condition number by several orders of magnitude were obtained there, for a simple simulated second order example.

The scaling can be applied also to the initialization algorithm. There the scaling needs to be applied in the linearized system matrix, and in the differentiation step in case the model order is higher than 2.

#### 4. Software package overview – RPEM ODE case

The software package is command driven, i.e. no GUI is available. It consists of a number of MATLAB scripts and functions. These are described in the next subsection.

##### 4.1 Scripts, functions and command flow

Roughly, the scripts and functions can be divided into five groups:

- *Live data measurements.* The two scripts of this group set up and perform clocked live data measurements. The scripts are **SetupLive.m** and **MeasurementMain.m**.
- *Simulated data generation.* The four scripts of this group define a dynamic system, that is then used for generation of simulated data. The scripts and functions are **SetupData.m**, **f.m**, **h.m** and **GenerateData.m**.
- *Recursive identification.* The five scripts of this group perform the actual identification tasks, supporting user interaction. The scripts and functions are **SetupRPEM.m**, **RPEM.m**, **h\_m.m**, **dhd\_x\_m.m** and **ReInitiate.m**.
- *Supporting functions - not called by the user.* The four functions of this group are called by scripts of the previous group. They are all related to the implementation of the RHS model of the identified ODE. The scripts are **GenerateIndices.m**, **f\_m.m**, **dfdx\_m.m** and **dfdtheta\_m.m**.
- *Preparation and display of results.* There are eleven scripts in this group. They all prepare, compute and display results of the identification process. The scripts are **PlotData.m**, **SimulateModelOutput.m**, **PlotParameters.m**, **PlotPredictionErrors.m**, **PlotEigenvalues.m**, **PlotCondition.m**, **ComputeRPEMLossFunction.m**, **PlotModelOutput.m**, **PlotSystemAndModelOutput.m**, **PlotResidualErrors.m** and **MeanResidualAnalysis.m**.

These groups of scripts and functions need to be operated in a particular order to make sense. This order of execution between scripts and functions is displayed with arrows in Figure 1. A single directional arrow indicates that the script/function pointed at may be executed only after the execution of the pointing script/function. See Figure 1 for details.

There are three major ways to exploit the five groups of scripts and functions.

1. In case the user has input and output signals available, the first step is to define and run the script **SetupLive.m**. This sets basic parameters like the sampling period. The user can then proceed directly to use the groups *Recursive Identification* and *Preparation and display of results*. The data, which can be simulated or live, should be stored in the (row) matrices  $u$  and  $y$ .
2. In case the user is to perform live measurements, all the steps of the *Live data measurement* group should be executed first. The user can then proceed directly to use the groups *Recursive Identification* and *Preparation and display of results*.
3. In case the user intends to use simulated data, this data can be generated by execution of the scripts and functions of the group *Simulated data generation*. The user can then proceed directly to use the groups *Recursive Identification* and *Preparation and display of results*.

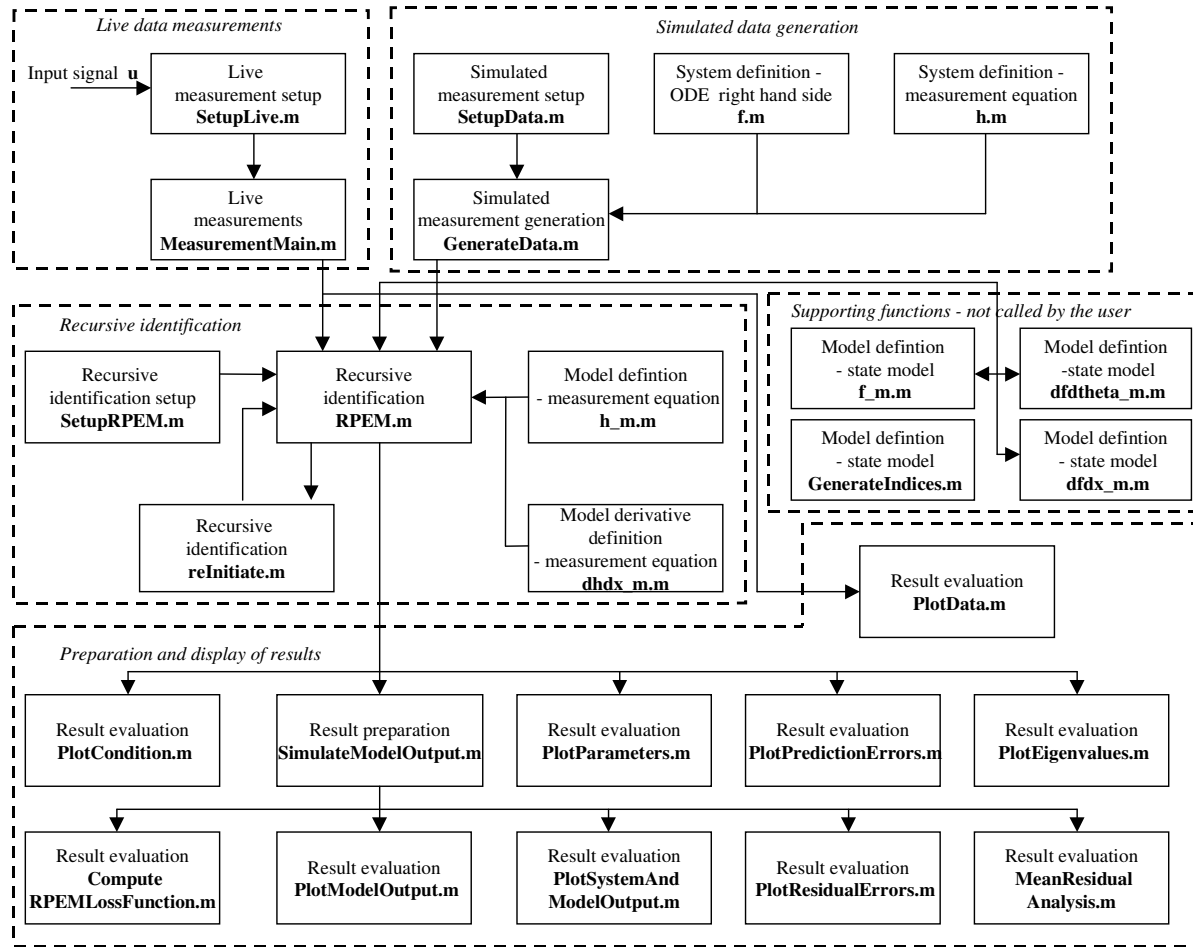


Figure 1: MATLAB scripts, functions and their relations in terms of execution order.

## 4.2 Restrictions

The main restrictions of the software are

- The software is command line driven - no GUI support is implemented.
- The software does not support true real time operation - there is no real time OS support implemented.
- The software has been tested and run using MATLAB 5.3, MATLAB 6.5, MATLAB 7.0 and MATLAB 7.9..

## 5. Data input – RPEM ODE case

The generation of data begins the section where the actual software is described. Since the user has access to all source files, the descriptions below do not describe code related issues and internal variables. Only the parts that are required for the use of the software package are covered. When m-files are reproduced, only the relevant parts are included, the reader should be aware that more information can be found in the source code. Note that the setup files are to be treated as templates, the user is hence required to modify right hand sides only - no addition or deletion of code should be used in the normal use of the package.

### 5.1 Simulated data

The generation of simulated data requires that the user

1. Modifies the underlying ODE model, as given by **f.m** and **h.m**. The function **f.m** implements the RHS of the ODE, using a conventional MATLAB function call. Note that the built in ODE solvers of MATLAB are not used. Instead an Euler algorithm is implemented. The reason is that this allows the generation of simulated data that can be exactly described by the applied model, should this be desired. The function **h.m** implements the (possibly nonlinear) measurement equation. The functions allow for addition of systems noise and measurement noise.
2. Provides further input data in the script **SetupData.m**. The parameters that define the data generation are directly written into this script. These parameters define the sampling period, the data length, the dimensions of the system, the type and parameters of the input signal, the type and parameters of the disturbances, as well as the initial value of the ODE.
3. Executes **SetupData.m**. This loads the necessary parameters into the MATLAB workspace.
4. Generates data by execution of **GenerateData.m**. After the execution of this script, variables with sampling instances, input signals and output signals are available in the MATLAB workspace.

*Example 1:* This and the following examples illustrates the use of the software package for identification of the system

$$\begin{aligned}\ddot{x}(t) + x(t) + (2 + u(t))\dot{x} &= u(t) \\ y(t) &= x(t) + e(t)\end{aligned}\quad (25)$$

This system is also used in [ 3 ] and [ 12 ], to asses effects of scaling. This system can be written in state space form as

$$\begin{aligned}\begin{pmatrix} \dot{x}_1(t) \\ \dot{x}_2(t) \end{pmatrix} &= \begin{pmatrix} x_2(t)(2 + u(t)) - u(t) \\ -x_1(t) - x_2(t) \end{pmatrix} \\ y(t) &= x_2(t) + e(t).\end{aligned}\quad (26)$$

Note that the ordering of states is not exactly as defined in the model ( 2 ). This is intentional since such situations are common in practical situations. It can be seen that the system is oscillatory with an input amplitude depending resonance frequency and damping.

The relevant parts of the files **f.m** and **h.m** become

**f.m**

```
function [f]=f(t,x,u,w)
    f(1,1)=x(2,1)*(2+u(1,1))-u(1,1);
    f(2,1)=-x(1,1)-x(2,1);
end
```

**h.m**

```
function [h]=h(t,x,u,e);
    h=x(2,1)+e(1,1);
end
```

Data is to be generated by simulation using a sampling period of  $T_s = 0.10s$ . 10000 input-output samples are to be generated. The input signal is to be selected with a uniform distribution in amplitude, with a mean of 0, a range  $[-1,1]$  and a clock period 3.0s. The measurement disturbance is to be white, zero mean with a standard deviation of 0.1.

The setup script that performs this task is **SetupData.m**

```
% dimensions...

nu=1; % Input signal dimension
nx_0=2; % State dimension
ny=1; % Output dimension, normally 1
```

```

% Input signal related...Type may be selected among:
%
% InputType=[
%     'PRBS      ';
%     'Gaussian  ';
%     'UniformPRBS';
%     'SineWave  ';
%     'Custom    '];

InputType=[
    'UniformPRBS'];
uAmplitude=[
    1.0];
uMean=[
    0];
uFrequency=[
    0.1];
ClockPeriod=[
    30];% Clock period vector in terms of sampling time

% System disturbance related...Type may be selected among:
%
% DisturbanceTypeSystem=[
%     'WGN      ';
%     'SineWave';
%     'Custom   '];

DisturbanceTypeSystem=[
    'WGN      ';
    'WGN      '];
wSigma=[
    0.0;
    0.0]; % Gaussian system noise standard deviation (discrete time)
wMean=[
    0;
    0];
wSineAmplitude=[

```



```

0;
0];
wSineFrequency=[
0;
0];

% Measurement disturbance related... Type may be selected among:
%
% DisturbanceTypeMeasuremet=[
%   'WGN      ';
%   'SineWave';
%   'Custom   '];

DisturbanceTypeMeasurement=[
    'WGN      '];
eSigma=0.1; % Gaussian measurement noise standard deviation
eMean=[
0;
0];
eSineAmplitude=[
0];
eSineFrequency=[
0];

% sampling time and data length

Ts=0.1; % Sampling time in seconds
N=10000; % Number of data points
SamplingInstances=(Ts:Ts:N*Ts);

% ODE related...

x0=[0.5 -1.0]'; % Initial values

```

The final step of the data generation is to execute the files **SetupData.m** and **GenerateData.m**. This is done in the MATLAB command window as follows

```

» SetupData
» GenerateData
...
percentReady =
    100
»

```

## 5.2 Live data

The generation of simulated data requires that the user

1. Is connected to the system via MATLAB. The connection must be such that commands to control DA-converters that generate input signals can be issued from within MATLAB. Similarly, commands that read AD-converters that sample output signals must be available from within MATLAB. The script **MeasurementMain.m** probably needs modification in a few parts in order to interface correctly to the AD- and DA-converters of the system of the user.
2. Generates an input signal, that is stored in the matrix (row vector in the one-dimensional input signal case) **u**.
3. Provides further data in the script **SetupLive.m**. The parameters that define the data generation are directly written into this script. These parameters defines the sampling period, the data length and the dimensions of the system.
4. Executes **SetupLive.m**. This loads the necessary parameters into the MATLAB workspace.
5. Generates data by execution of **MeasurementMain.m**. After the execution of this file, variables with input signals and output signals are available in the MATLAB workspace. This script operates as a loop that continuously polls the MATLAB real time clock, waiting for the next sampling instance. This means that it may not be possible to use the computer for other tasks during the data collection session. The reason for this solution is that it avoids the need for a real time OS connection. Note also that the calls to AD- and DA-converters may be different on other systems. This script is hence likely to require some modification.

*Example 2:* The setup script file **SetupLive.m** becomes (empty since simulated data is used here)

```

% dimensions...Note that nx is not really relevant,
% it is however required in the RPEM setup so it is set in this file

nu=[]; % Input signal dimension
nx=[]; % State dimension
ny=[]; % Output dimension, normally 1

% sampling time and data length

```

```
Ts=[]; % Sampling time in seconds
N=[]; % Number of data points
SamplingInstances=(Ts:Ts:N*Ts);
```

The measurement process is started by typing

```
» SetupData
» GenerateData
...
```

in the MATLAB command window. During the measurement session, the script continuously displays the time, the inputs as well as the measured outputs, as commanded to DA-converters and read by AD-converters. After termination all data that is needed for identification is available in the MATLAB workspace.

### 5.3 Display of data

After execution of either one of the chain of actions of section 5.1 or section 5.2, data can be plotted.

1. The **PlotData.m** script that is executed in the MATLAB command window. This script makes use of the dimensions of the system in order to divide the plot into several sub-windows, and in order to provide the axis text.

*Example 3:* The MATLAB command window command is

```
» PlotData
»
```

The following plot is generated

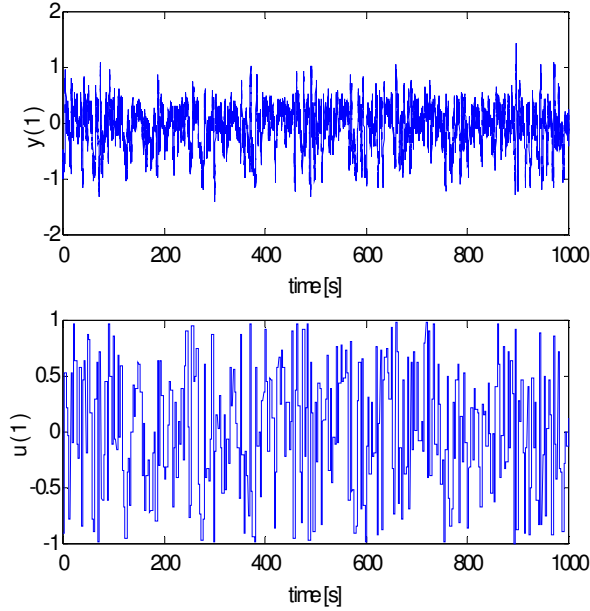


Figure 2: The result of a PlotData command.

## 6. Recursive Identification – RPEM ODE case

At this point everything is in place for a first identification run.

### 6.1 RPEM setup

The preparation for the identification run requires that the user

1. Modifies the output equation and the corresponding derivative of underlying ODE model, as given by **h\_m.m** and **dhd\_x\_m.m**. The function **h\_m.m** implements the output equation of the model. Note that this function is allowed to be a nonlinear function of the state and input. The function is not allowed to be dependent on the estimated parameters, it must be known a priori. Note also that the derivative of the function, with respect to the estimated state, needs to be supplied in the function **dhd\_x\_m.m**.
2. Provides further input data in the script **SetupRPEM.m**. The parameters that define the data generation are directly written into this script. These parameters define the dimension of the system, the initial value used in the ODE model, the gain sequence  $\mu(t)/t$ , the size of the initial value of the **R**-recursion, the initial value of the measurement covariance matrix  $\mathcal{A}(t)$ , the stability limit applied by the projection algorithm, the scale factor, as well as the down-sampling period used to avoid too large logs during long runs with high degree models. The reader is referred to [ 1 ] - [ 4 ] for details on these parameters, as well as on their use.

3. Executes **SetupRPEM.m**. This loads the necessary parameters into the MATLAB workspace.

*Example 4:* The system is to be identified with a second order model. The projection algorithm is to use a stability radius of 0.975 and the scale factor is selected equal to 2. The initial value of the measurement covariance matrix is selected equal to 0.1. The initial value of the **R** - recursion ( its inverse affecting the initial algorithmic gain) is selected equal to 100. The selection of the gain sequence  $\mu(t)/t$  is a little more complicated, see [ 1 ] for details.

The functions **h\_m.m** and **dhd\_x\_m.m** become

**h\_m.m**

```
function [h_m]=h_m(x_m,u);  
    h_m=x_m(1,1);  
end
```

**dhd\_x\_m.m**

```
function [dhd_x_m]=dhd_x_m(x,u);  
    dhd_x_m=[1 0];  
end
```

The setup script **SetupRPEM.m** becomes

```
nx=2;  
x_m_0=[0.5 -1]';  
  
%  
% Remaining initial values  
%  
  
muFactor=300; % To stabilize Gamma and to reduce the gain  
if exist('theta_0_new')  
    muFactor=1000;  
end  
mu_0=5;  
mu0=0.9995;  
y_m_0=0;  
initialNoiseVariance=0.1; % Initial value for the prediction error  
variance  
scaleFactorR=100; % the size of the initial diagonal approximation  
of the Hessian
```

```

%
% Parameters
%

stabilityLimit=0.975; % The linearized pole radius used for
stability checking and projection
downSampling=10; % The downsampling factor used when data from the
run is saved
scalingTs=2; % The scaling factor with which the sampling period is
multiplied during identification

```

Finally the user executes **SetupRPEM.m** in the MATLAB command window

```

» setupRPEM
»

```

## 6.2 RPEM command window control and estimated parameters

In order to perform an identification run the user is required to execute and provide input to the script **RPEM.m**. The execution of this script makes use of four additional functions, implementing the polynomial model applied for modeling of the RHS of the ODE. These functions are **f\_m.m**, **dfdx\_m.m**, **dfdtheta\_m.m** and **GenerateIndices.m**. The latter function generates the exponents of all factors of all terms of the polynomial expansion. The generation of these indices involves nested loops. They are therefore calculated in advance and used in repetitive calls in the form of a table.

To identify the system, the user is required to

1. Execute the script **RPEM.m**
2. Provide the degrees of the polynomial model (*polynomialOrders*) when prompted. The *polynomialOrders* variable is a column vector with the first element corresponding to the maximal degree of  $x_1$ , the second element corresponding to the maximal degree of  $x_2$  and so on. The last element corresponds to the maximum degree of the derivative of highest degree of the last input signal component. In the present example, *polynomialOrders* = [1 2 3]' would mean that the highest degree term of the polynomial expansion is  $\theta_{123}x_1x_2^2u^3$ .
3. Provide a list of indices that are not to be used (*notUsedIndices*) by the algorithm. The indices exclude terms in the polynomial expansions. Providing an empty matrix ([]) indicates that no terms shall be excluded. The list of not used indices are to be provided as rows in a matrix, where the number of rows equals the number of terms that are to be excluded from the model. In the present

example *notUsedIndices* = [0 0 0; 1 1 1] would mean that the terms  $\theta_{000}$  and  $\theta_{111}x_1x_2u$  are to be excluded from the model.

4. Specify the algorithm used as discretization method (*method*) as a text string. The user can provide either the text string 'Euler' or 'midpoint'.
5. Provide the initial parameter vector (*theta\_0*). Note that this parameter vector needs to correspond to a linearized system with all poles within the stability radius indicated by the script **SetupRPEM.m**. If the initial parameter vector does not meet this criterion the user is prompted for *theta\_0* again. Observe that the scale factor of the sampling period needs to be accounted for - it is a part of the linearized model, cf. [ 1 ].

Note: A good strategy is to initialize the algorithm with a model that has time constants and a static gain that are similar to those of the system.

*Example 5:* The algorithm is in this example initialized with

$$\hat{\theta}(0) = (0.0000 \quad 1.0000 \quad -1.0000 \quad 0.0000 \quad -0.2500 \quad 0.0000 \quad 0.0000 \quad 0.0000)^T. \quad (27)$$

This corresponds to the model

$$\varphi(\mathbf{x}, u) = (1 \quad u \quad x_2 \quad x_2u \quad x_1 \quad x_1u \quad x_1x_2 \quad x_1x_2u)^T. \quad (28)$$

In this example comments and explanations have been added. To distinguish these from the actual commands the comments are in italics. The command sequence applied in the MATLAB command window is

» RPEM

ans =

Input polynomialOrders and notUsedIndices - *The script asks for the max degrees of states and inputs*

K» polynomialOrders=[1 1 1]

polynomialOrders =

1

1

1

K» notUsedIndices=[] - *The script asks for terms of the polynomial that are to be excluded*

notUsedIndices =

[]

K» return

allIndices = - *The script returns the degrees of all included terms, input degrees to the right*

0 0 0

0 0 1

0 1 0

0 1 1

```

1  0  0
1  0  1
1  1  0
1  1  1
ans =
Input method      - The script asks for a discretization method, possible choices so far 'Euler' and
                    'midpoint'
K» method = 'Euler'
method =
    'Euler'
K» return
ans =
Input theta_0      - The script asks for an initial parameter vector
K» theta_0=[0 1 -1 0 -0.25 0 0 0]'
theta_0 =
    0
    1.0000
   -1.0000
    0
   -0.2500
    0
    0
    0
K» return
LinearizedPoleRadii = - The script returns the polr-radii of the linearized, initial model
    0.9000
    0.9000
...
percentReady = - The script displays the fraction of the processing that is completed.
    100
ans = - The script displays the identified parameters – scaled parameters to the left
    0.0003    0.0013
    0.2508    1.0031
   -0.4946   -0.9893
   -0.0018   -0.0035
   -0.4992   -1.9968
   -0.2512   -1.0050

```



```
0.0163  0.0326
-0.0174 -0.0347
```

»

The estimated parameters of the left column correspond to the ones obtained directly from the RPEM, i.e. these are scaled parameters. The right column contain parameters that are recomputed to correspond to the original sampling period. Note that the exact result depends on the generated input signal. This may differ between systems and execution occasions since the seed for the random number generator may differ. Hence, slight variations of the estimated parameters are normal.

### 6.3 Re-initiation, multiple runs and iterative refinement

The script **RPEM.m** produces a result for a certain choice of degrees of the right hand polynomial of the ODE (if the stability check is not triggered so that the algorithm gets stuck close to the stability limit). In case the result is not deemed sufficient, then a higher degree right hand side may be needed. The opposite may also be true, i.e. the result is sufficient but the number of parameters used may be unnecessarily high. So there is a need to

- Modify the degrees of the polynomial model of the ODE.
- Remove specific terms of the polynomial model of the ODE.
- Rerun the RPEM from the previous end results, with a redefined right hand side polynomial.

Exactly this is supported by the script **reInitiate.m**.

Note: Support for stepping also of the model order would be preferred. Such stepping does however have complicated (nonlinear) stability impacts. For this reason the development of such functionality is postponed to later versions of the software package.

In order to perform a new RPEM run with modified degrees, then user is required to

1. Run the script **reInitiate.m**. That script prompts the user for *polynomialOrders* and *notUsedIndices*. The parameter vector at the end of the run, together with the previous and new degrees, are then used to re-initiate all relevant quantities of the RPEM.
2. Rerun **RPEM.m**. Note that the RPEM does not need to prompt the user for any further information this time.

*Example 6:* In this example the degree of the input signal is increased to 2.

The MATLAB command window commands become

```
» reInitiate
```

```
ans =
```

```
Input polynomialOrders and notUsedIndices
```

```
K» polynomialOrders=[1 1 2]'
```

```
polynomialOrders =
```

```
1
```

```
1
```

2

K» return

» RPEM

...

The identification is now finalized and everything is set for display of the results.

## 7. Display of results – RPEM ODE case

The display of results is straightforward. By a study of the source code, users should be able to tailor available scripts and also write own ones when needed.

### 7.1 Parameters

In order to plot the parameters the user is required to

1. Execute the script **PlotParameters.m**. The components of the parameter vector are then plotted as a function of time. Note that the time scale is assumed to be seconds. In case another time scale is required, the figure needs to be edited after plotting, or the script needs modification.

*Example 7:* The command in the MATLAB command window is

» PlotParameters

»

The following plot is then generated

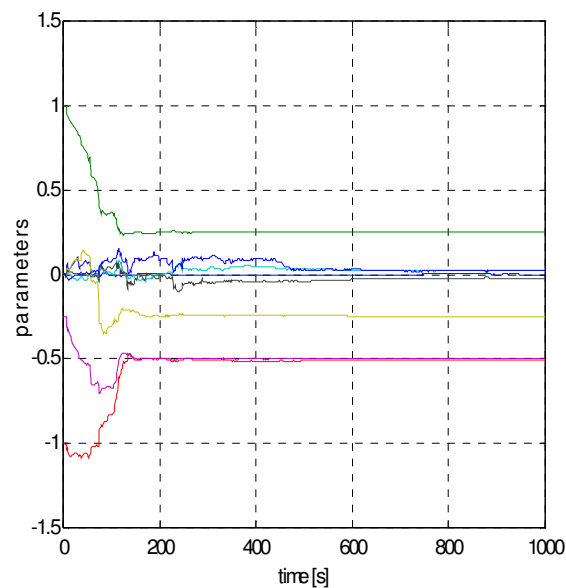


Figure 3: The result of a PlotParameters command.

## 7.2 Prediction errors

In order to plot the parameters the user is required to

1. Execute the script **PlotPredictionErrors.m**. The prediction errors are then plotted as a function of time. Note that the time scale is assumed to be seconds. In case another time scale is required, the figure needs to be edited after plotting, or the script needs modification.

*Example 8:* The MATLAB command window command is

```
» PlotPredictionErrors
```

```
»
```

The following plot is generated

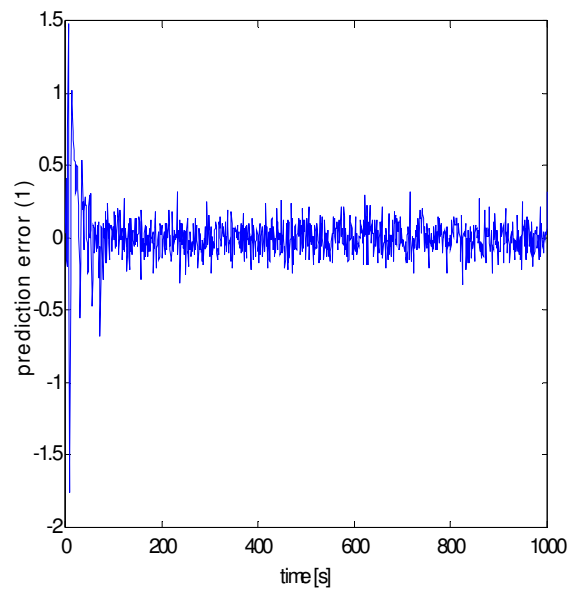


Figure 4: The result of a PlotPredictionErrors command.

## 7.3 Eigenvalues

In order to plot the convergence of the eigenvalues over time, the user is required to

1. Execute the script **PlotEigenvalues.m**. The eigenvalues of the Hessian are then plotted as a function of time. Note that the time scale is assumed to be seconds. In case another time scale is required, the figure needs to be edited after plotting, or the script needs modification.

*Example 9:* The MATLAB command window command is

```
» PlotEigenvalues
```

```
»
```

The following plot is generated

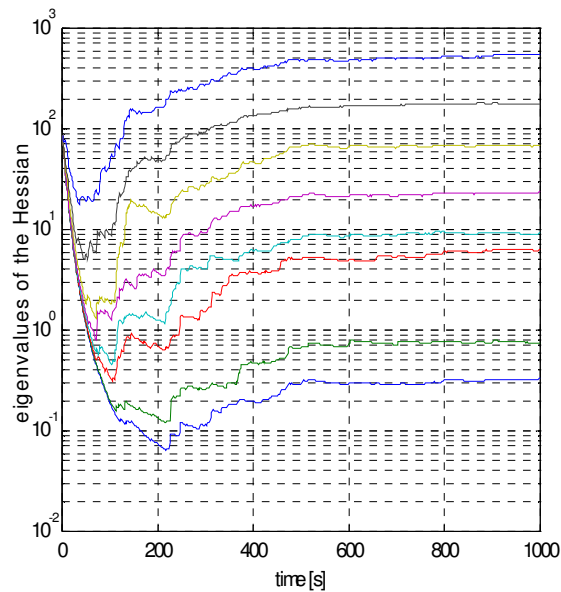


Figure 5: The result of a PlotEigenvalues command.

#### 7.4 Condition number

In order to plot the convergence of the condition number over time, the user is required to

1. Execute the script **PlotCondition.m**. The condition number is then plotted as a function of time.

Note that the time scale is assumed to be seconds. In case another time scale is required, the figure needs to be edited after plotting, or the script needs modification.

*Example 10:* The MATLAB command window command is

```
» PlotCondition
»
```

The following plot is generated

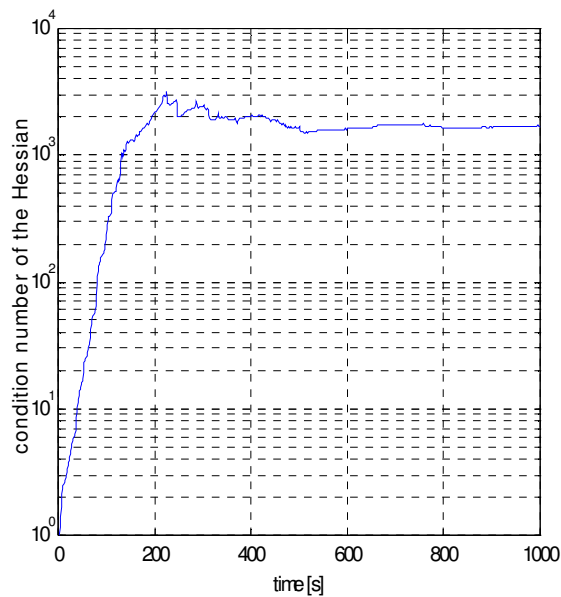


Figure 6: The result of a PlotCondition command.

### 7.5 End of run model simulation

The above plot commands make use of logged signals, covering the transient part of the identification process. This is not always desired. To compare the identified mode to the measured data it is e.g. more appropriate to simulate the model, using the parameters obtained at the end of the identification run.

Hence, to prepare for the remaining plot commands the user is required to

1. Execute the script **SimulateModelOutput.m**.

*Example 11:* The MATLAB command window command is

```
» SimulateModelOutput
```

```
...
```

```
percentReady =
```

```
100
```

```
»
```

The remaining plot commands and model validation commands can now be executed.

### 7.6 Simulated model output

In order to plot the simulated model output signal over time, the user is required to

1. Execute the script **PlotModelOutput.m**. The output signals of the model are then plotted as a function of time. Note that the time scale is assumed to be seconds. In case another time scale is required, the figure needs to be edited after plotting, or the script needs modification.

*Example 12:* The MATLAB command window command is

```
» PlotModelOutput
```

```
»
```

The following plot is generated

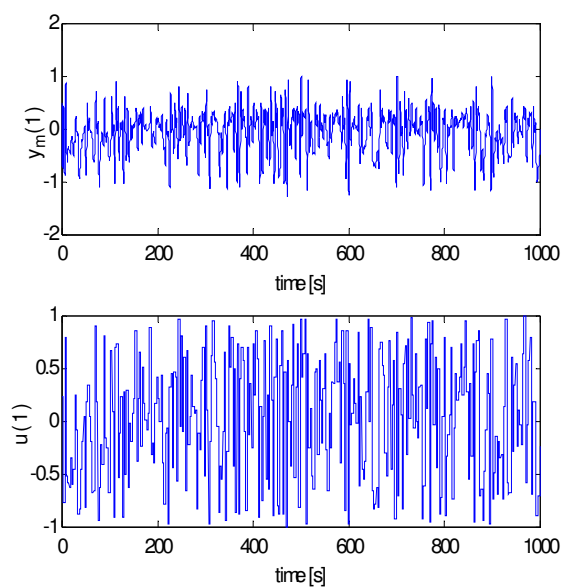


Figure 7: The result of a PlotModelOutput command.

### 7.7 Simulated model output together with data

In order to plot the simulated model output signal over time, together with the corresponding measured data, the user is required to

1. Execute the script **PlotSystemAndModelOutput.m**. The output signals of the model and the system are then plotted as a function of time, in the same plots. Note that the time scale is assumed to be seconds. In case another time scale is required, the figure needs to be edited after plotting, or the script needs modification.

*Example 13:* The MATLAB command window command is

```
» PlotSystemAndModelOutput
```

```
»
```

The following plot is generated

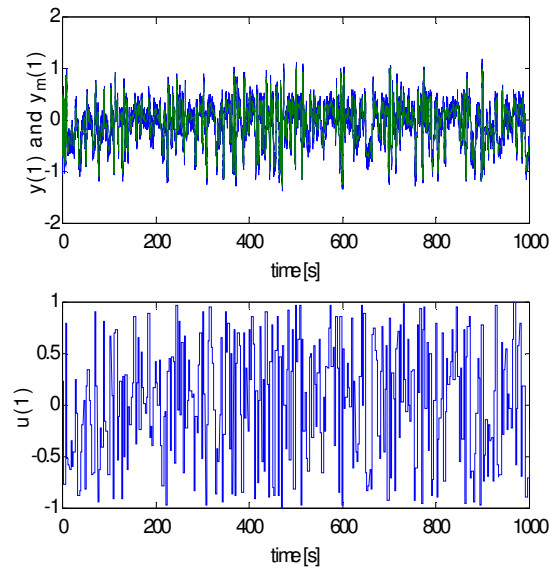


Figure 8: The result of a `PlotSystemAndModelOutput` command.

### 7.8 Residual errors

In order to plot the prediction errors, obtained from the simulated model output signal using parameters at the end of the identification run, the user is required to

1. Execute the script **PlotResidualErrors.m**. The errors are then plotted as a function of time. Note that the time scale is assumed to be seconds. In case another time scale is required, the figure needs to be edited after plotting, or the script needs modification.

*Example 14:* The MATLAB command window command is

```
» PlotResidualErrors
```

```
»
```

The following plot is generated

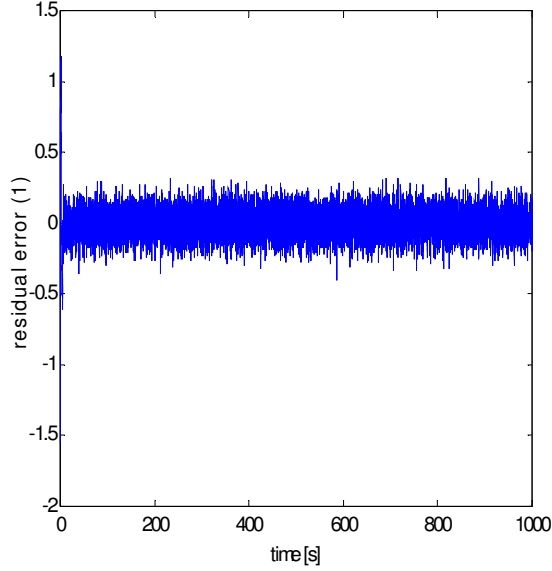


Figure 9: The result of a PlotResidualErrors command.

## 8. Model validation – ODE case

Two scripts are provided for model validation purposes (in addition to the commands of section 7). The first method studies the value of the loss function that is minimized by the RPEM-algorithm. Since the measurement covariance matrix is estimated, the loss function contains an additional term on top of the sample average of the squared prediction errors.

### 8.1 RPEM loss function

The RPEM loss function that is computed is given by

$$V(\hat{\theta}(t_0 + NT_s)) = \frac{1}{2} \frac{1}{N} \sum_{i=1}^N \varepsilon(t_0 + iT_s, \hat{\theta}(t_0 + NT_s))^T \varepsilon(t_0 + iT_s, \hat{\theta}(t_0 + NT_s)) + \frac{1}{2} \log \det(\hat{\Lambda}(t_0 + NT_s)) \quad (29)$$

The user is referred to [ 1 ], [ 6 ] and the references therein for further details. In order to compute the loss function, using parameters at the end of the identification run, the user is required to

1. Execute the script **ComputeRPEMLossFunction.m**. The loss function is then computed.

*Example 15:* The MATLAB command window command is

```
» ComputeRPEMLossFunction
```

```
...
```

```
percentReady =
```

```
100
```

```
V =
```



-1.7094

»

Note: Another relevant measure to use is the sum of the squared prediction errors.

## 8.2 Mean residual analysis

Mean residual analysis is a method that evaluates the obtained static characteristics of an identified model of any kind. It operates by sorting residual errors into bins, the bin being decided by the value of the measured output signal with the same time index as the residual error. The mean of the residuals are then computed, in each bin, and plotted against the range of the output signal. The number of samples of each bin is also plotted. The user is referred to [ 7 ] for further details. In order to perform mean residual analysis, the user is required to

1. Execute the script **MeanResidualAnalysis.m**.
2. Provide the intervals used by the method when prompted for *intervals*.

*Example 16:* This example performs mean residual analysis using about 40 intervals, each with an output amplitude width of 0.1. The MATLAB command window command is

```
» meanResidualAnalysis
```

```
ans =
```

```
Input intervals for division into bins
```

```
K» intervals=(-2:0.1:2);
```

```
K» return
```

```
»
```

The following plot results

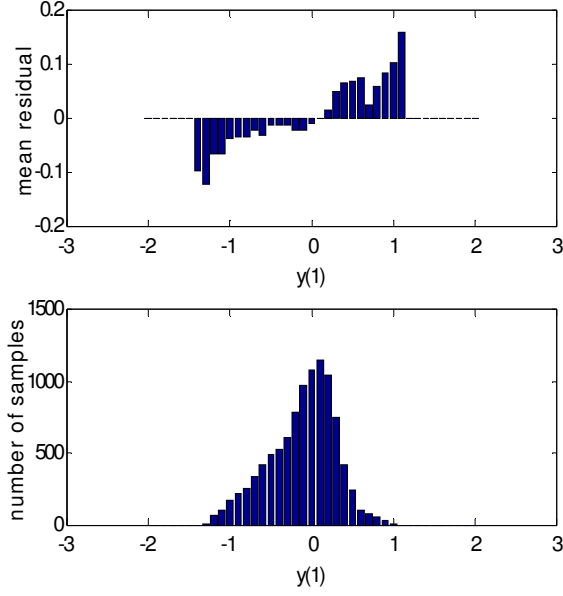


Figure 10: The result of a MeanResidualAnalysis command.

## 9. The RPEM for identification of static nonlinear MISO systems

The RPEM for identification of MISO static nonlinear systems is based on the model

$$\begin{aligned}
 & f(u_1, \dots, u_1^{(n_1)}, \dots, u_k, \dots, u_k^{(n_k)}, \theta) \\
 &= \sum_{i_{u_1}=0}^{I_{u_1}} \dots \sum_{i_{u_1}^{(n_1)}=0}^{I_{u_1}^{(n_1)}} \dots \sum_{i_{u_k}=0}^{I_{u_k}} \dots \sum_{i_{u_k}^{(n_k)}=0}^{I_{u_k}^{(n_k)}} \theta_{i_{u_1}^{(n_1)} \dots i_{u_k}^{(n_k)}} (u_1)^{i_{u_1}} \dots \\
 & \dots (u_1^{(n_1)})^{i_{u_1}^{(n_1)}} \dots (u_k)^{i_{u_k}} \dots (u_k^{(n_k)})^{i_{u_k}^{(n_k)}} = \varphi^T(\mathbf{u}) \theta.
 \end{aligned} \tag{30}$$

Here

$$\begin{aligned}
 \theta &= \left( \theta_{0 \dots 0} \quad \dots \quad \theta_{0 \dots I_{u_k}^{(n_k)}} \quad \theta_{0 \dots 010} \quad \dots \quad \theta_{0 \dots 01 I_{u_k}^{(n_k)}} \quad \dots \right. \\
 & \quad \left. \theta_{0 \dots 0 I_{u_k}^{(n_k-1)} 0} \quad \dots \quad \theta_{0 \dots 0 I_{u_k}^{(n_k-1)} I_{u_k}^{(n_k)}} \quad \dots \quad \theta_{I_{u_1}^{(n_1)} \dots I_{u_k}^{(n_k)}} \right)^T \\
 \varphi &= \left( 1 \quad \dots \quad \left( (u_k^{(n_k)})^{I_{u_k}^{(n_k)}} \right) \quad u_k^{(n_k-1)} \quad \dots \quad \left( u_k^{(n_k-1)} (u_k^{(n_k)})^{I_{u_k}^{(n_k)}} \right) \quad \dots \quad (u_k^{(n_k-1)})^{I_{u_k}^{(n_k-1)}} \quad \dots \right)
 \end{aligned}$$

$$\left( \left( u_k^{(n_k-1)} \right)^{I_{u_k^{(n_k-1)}}} \left( u_k^{(n_k)} \right)^{I_{u_k^{(n_k)}}} \right) \dots \left( \left( u_1 \right)^{I_{u_1}} \dots \left( u_k^{(n_k)} \right)^{I_{u_k^{(n_k)}}} \right) \right)^T \quad (31)$$

The recursive algorithm is then developed exactly as in [1] and [2]. The main difference is that the state equation iteration and the corresponding state gradient iteration disappears. The end result is the following algorithm where all variables are explained in detail in [1] and [2]

$$\begin{aligned} \boldsymbol{\varepsilon}(t) &= \mathbf{y}_m(t) - \mathbf{y}(t) \\ \mathbf{A}(t) &= \mathbf{A}(t - T_s) + \frac{\mu(t)}{t} \left( \boldsymbol{\varepsilon}(t) \boldsymbol{\varepsilon}^T(t) - \mathbf{A}(t - T_s) \right) \\ \mathbf{R}(t) &= \mathbf{R}(t - T_s) + \frac{\mu(t)}{t} \left( \boldsymbol{\psi}(t) \mathbf{A}^{-1}(t) \boldsymbol{\psi}^T(t) - \mathbf{R}(t - T_s) \right) \\ \hat{\boldsymbol{\theta}}(t) &= \left[ \hat{\boldsymbol{\theta}}(t - T_s) + \frac{\mu(t)}{t} \mathbf{R}^{-1}(t) \boldsymbol{\psi}(t) \mathbf{A}^{-1}(t) \boldsymbol{\varepsilon}(t) \right] \\ \boldsymbol{\varphi}(t) &= \begin{pmatrix} 1 & \dots & \left( \left( u_k^{(n_k-1)}(t) \right)^{I_{u_k^{(n_k-1)}}} \right) & u_k^{(n_k-1)}(t) & \dots & \left( \left( u_k^{(n_k-1)}(t) \right) \left( u_k^{(n_k)}(t) \right)^{I_{u_k^{(n_k)}}} \right) & \dots \\ \dots & \left( u_k^{(n_k-1)}(t) \right)^{I_{u_k^{(n_k-1)}}} & \dots & \left( \left( u_k^{(n_k-1)}(t) \right)^{I_{u_k^{(n_k-1)}}} \left( u_k^{(n_k)}(t) \right)^{I_{u_k^{(n_k)}}} \right) & \dots & \left( \left( u_1(t) \right)^{I_{u_1}} \dots \left( u_k^{(n_k)}(t) \right)^{I_{u_k^{(n_k)}}} \right) \end{pmatrix}^T \\ y(t + T_s) &= \boldsymbol{\varphi}^T(t) \boldsymbol{\theta} \\ \boldsymbol{\psi}(t + T_s) &= \boldsymbol{\varphi}(t + T_s). \end{aligned} \quad (32)$$

It should be noted that this algorithm collapses to a least-squares case. Hence it will always converge.

## 10. Software – Static case

All software for identification of nonlinear static systems fits into the framework of Figure 1. As much as possible of the software has been kept usable for both the nonlinear dynamic case and the nonlinear static case. However, static versions of the following 7 scripts replace or add to the dynamic counterparts, in case a static model is identified: **dhdthetaStatic.m** (new m-file), **GenerateDataStatic.m**, **hStatic.m**, **RPEMStatic.m**, **SetupDataStatic.m**, **SetupRPEMStatic.m** and **SimulateModelOutputstatic.m**. These files are now briefly described.

**SetupDataStatic.m**: This m-file sets up the parameters needed for static data generation. The file has been modified by selection of parameters needed for dynamic data generation consistent with the static case. Most variables are however left as dummy variables, so that **SetupDataStatic.m** does not prevent the successful execution of existing scripts later in the data processing chain.

**GenerateDataStatic.m:** This file has been created from **GenerateData.m** by deletion of the state iteration. Note that this script makes use of **h.m** that must be based only on input signals in the static case. The state argument in the call is an empty matrix.

**SetupRPEMStatic.m:** This file has been created from **SetupRPEM.m** by setting the state dimension equal to zero, by setting state initial values equal to a 0x1 matrix, and by setting the sampling period scale factor scalingTs equal to 1. The latter is important since no scaling shall be applied in the static case.

**RPEMStatic.m:** This file has been created from **RPEM.m** by deletion of the parts of the algorithm that are not relevant in the static case. This is true for the state variable iteration and the corresponding state gradient iteration. The exact details follow by a comparison of the RPEM of [2] with (32) above. A further modification is the use of calls to the functions **hStatic\_m.m** and **dhdthetaStatic\_m.m**, in order to compute output predictions and gradients.

**hStatic\_m.m:** This function computes (30) above. It was created by small modifications of **f\_m.m**. The order of the output prediction (1) is e.g. included in the function call.

**DhdthetaStatic\_m.m:** This function generates  $\phi$  of (31). It was created by small modifications of **dfdtheta\_m.m**. The order of the output prediction (1) is e.g. included in the function call.

**SimulateModelOutputStatic.m.** This file has been created from **SimulateModelOutput.m** by deletion of the state variable iteration and by a call to the function **hStatic\_m.m** for computation of the output prediction.

## 11. Kalman filter based initialization algorithm

The Kalman filter based algorithm for initialization of the RPEM (dynamic case) is based on (3) just like the RPEM. The recursive algorithm is then developed exactly as in [ 5 ] and [ 6 ]. The end result is the following scheme where all variables are explained in detail in [ 5 ] and [ 6 ]

Initiate

$$\bar{x}(0| -T_s), P(0| -T_s)$$

Iterate

$$K(t) = P(t|t - T_s)H^T (HP(t|t - T_s)H^T + R_2)^{-1}$$

$$\bar{x}(t|t) = \bar{x}(t|t - T_s) + K(t)(y_m(t) - H\bar{x}(t|t - T_s))$$

$$P(t|t) = P(t|t - T_s) - K(t)HP(t|t - T_s)$$

$$\bar{x}(t + T_s | t) = F(t, \xi(t), u(t))\bar{x}(t) \quad (33)$$

$$P(t + T_s | t) = F(t, \xi(t), u(t))P(t | t)F^T(t, \xi(t), u(t)) + R_1$$

It should be noted that this algorithm is similar to a least squares problem, solved recursively. Hence (33) will always converge. The key idea to obtain the algorithm is to replace the estimated states of the RPEM by an approximation  $\xi(t)$ , obtained by repeated differentiation of the output signal. Note that the scaled sampling period needs to be applied in the preceding differentiation step, cf. [ 5 ] and [ 6 ].

## 12. Software – Kalman filter based initialization algorithm

To use the Kalman filter based initialization algorithm the following steps are required. Please note, that since this algorithm is based on the same model structure as the dynamic RPEM in section 6 some of the information is identical.

### 12.1 Initialization setup

The preparation for the identification run requires that the user

1. Modifies the output equation and the corresponding derivative of the underlying ODE model, as given by **h\_m.m** and **dhd\_x\_m.m**. The function **h\_m.m** implements the output equation of the model. Note that this function is allowed to be a nonlinear function of the state and input. The function is not allowed to be dependent on the estimated parameters, it must be known a priori. Note also that the derivative of the function, with respect to the estimated state, needs to be supplied in the function **dhd\_x\_m.m**.
2. Provides further input data in the script **SetupInit.m**. The parameters that define the data generation are directly written into this script. These parameters define the dimension of the system, the initial value used in the ODE model, the size of the initial value of the  $P$ -recursion of (33), the magnitude of the covariance matrices for the state and parameter vector system disturbances, ( $R_{1,x}$  and  $R_{1,\theta}$  respectively), the magnitude of the measurement noise covariance,  $R_2$ , the scale factor, as well as the down-sampling period used to avoid too large logs during long runs with high degree models. The reader is referred to [ 1 ] - [ 6 ] for details on these parameters, as well as on their use.
3. Executes **SetupInit.m**. This loads the necessary parameters into the MATLAB workspace.

*Example 17:* The system used above is to be identified with a second order model, which implies that the true parameter and corresponding regressor vectors are

$$\theta = (0 \quad 1 \quad -1 \quad 0 \quad -2 \quad -1 \quad 0 \quad 0)^T \quad (34)$$

$$\varphi(x, u) = (1 \quad u \quad x_2 \quad x_2 u \quad x_1 \quad x_1 u \quad x_1 x_2 \quad x_1 x_2 u)^T. \quad (35)$$

The scale factor is selected equal to 2. The magnitude of the state and parameter disturbance covariance are selected to 0.01 and 0.001 respectively, while the measurement covariance matrix is selected equal to 1. The initial value of the  $P$  - recursion is selected equal to 0.1. The functions **h\_m.m** and **dhd\_x\_m.m** become

**h\_m.m**

```
function [h_m]=h_m(x_m,u);
    h_m=x_m(1,1);
end
```

**dhd\_x\_m.m**

```
function [dhd_x_m]=dhd_x_m(x,u);
    dhd_x_m=[1 0];
end
```

The setup script **SetupInit.m** becomes

```
nx=2;
x_m_0=[1 0]';

%
% Remaining initial values
%

y_m_0=[x_m_0(1)];
scaleFactorP=1e-1;

R1_x=1e-1;           % Scaling factor for state disturbance
R1_theta=1e-3;       % Scaling factor for parameter vector disturbance
R2=1e-2;             % Scaling factor for measurement noise

%
% Parameters
%

downSampling=10; % The downsampling factor used when data from the
run is saved
scalingTs=2; % The scaling factor with which the sampling period is
multiplied during identification
```

Finally the user executes **SetupInit.m** in the MATLAB command window

```
>> setupInit  
>>
```

## 12.2 Initialization command window control and estimated parameters

In order to perform an identification run the user is required to execute and provide input to the script **Init.m**. The execution of this script makes use of four additional functions, implementing the polynomial model applied for modeling of the RHS of the ODE. These functions are **F\_m.m**, **dfdx\_m.m**, **differentiateY.m** and **GenerateIndices.m**. The latter function generates the exponents of all factors of all terms of the polynomial expansion. The generation of these indices involves nested loops. They are therefore calculated in advance and used in repetitive calls in the form of a table.

To identify the system, the user is required to

1. Execute the script **Init.m**
2. Provide the degrees of the polynomial model (*polynomialOrders*) when prompted. The *polynomialOrders* variable is a column vector with the first element corresponding to the maximal degree of  $x_1$ , the second element corresponding to the maximal degree of  $x_2$  and so on. The last element corresponds to the maximum degree of the derivative of highest degree of the last input signal component. In the present example, *polynomialOrders* = [1 2 3]' would mean that the highest degree term of the polynomial expansion is  $\theta_{123}x_1x_2^2u^3$ .
3. Provide a list of indices that are not to be used (*notUsedIndices*) by the algorithm. The indices exclude terms in the polynomial expansions. Providing an empty matrix ([]) indicates that no terms shall be excluded. The list of not used indices are to be provided as rows in a matrix, where the number of rows equals the number of terms that are to be excluded from the model. In the present example *notUsedIndices* = [0 0 0; 1 1 1] would mean that the terms  $\theta_{000}$  and  $\theta_{111}x_1x_2u$  are to be excluded from the model.
4. Provide the initial parameter vector (*theta\_0*). Note that unlike in the dynamic RPEM case, this parameter vector does *not* necessarily need to correspond to a linearized system with all poles within the unit circle. It is for example perfectly all right to choose *theta\_0*=0. However, when using the initialization algorithm in combination with the dynamic RPEM it is important to check that the *final* parameter vector obtained with **Init.m** corresponds to a linearized model with poles within the stability region as described in section 6. If this is not the case it will not be possible to use the obtained parameter vector for initializing the RPEM. Observe that the scale factor of the sampling period needs to be accounted for - it is a part of the linearized model, cf. [1].

*Example 18:* In this example the same system as in Example 17 is studied. The algorithm in this example is initialized with

$$\hat{\theta}(0) = (0 \ 0 \ 0 \ 0 \ 0 \ 0 \ 0 \ 0)^T. \quad (36)$$

This corresponds to the model

$$\varphi(\mathbf{x}, u) = (1 \ u \ x_2 \ x_2 u \ x_1 \ x_1 u \ x_1 x_2 \ x_1 x_2 u)^T. \quad (37)$$

In this example comments and explanations have been added. To distinguish these from the actual commands the comments are in italics. The command sequence applied in the MATLAB command window is

```
>> Init
ans =
Input polynomialOrders and notUsedIndices - The script asks for the max degrees of states and inputs
K>> polynomialOrders=[1 1 1]'
polynomialOrders =
    1
    1
    1
K>> notUsedIndices=[]
notUsedIndices =
    []
K>> return
allIndices = - The script returns the degrees of all included terms, input degrees to the right
    0    0    0
    0    0    1
    0    1    0
    0    1    1
    1    0    0
    1    0    1
    1    1    0
    1    1    1
ans =
Input theta_0 - The script asks for an initial parameter vector
K>> theta_0=zeros(8,1)
theta_0 =
    0
    0
```



```

0
0
0
0
0
0
K>> return
...
percentReady = - The script displays the fraction of the processing that is completed.
100
LinearizedPoleRadii=
0.9956
0.9956
ans = - The script displays the identified parameters – scaled parameters to the left
0.0000 0.0000
0.2450 0.9798
-0.4900 -0.9800
-0.0004 -0.0007
-0.4900 -1.9600
-0.2449 -0.9797
-0.0002 -0.0003
-0.0006 -0.0012
»

```

The estimated parameters of the left column correspond to the ones obtained directly from the RPEM, i.e. these are scaled parameters. The right column contain parameters that are recomputed to correspond to the original sampling period. Note that the exact result depends on the generated input signal. This may differ between systems and execution occasions since the seed for the random number generator may differ. Hence, slight variations of the estimated parameters are normal.

### 12.3 Re-initiation, multiple runs and iterative refinement

The script **Init.m** produces initial parameters for the RPEM. To run the RPEM from the end results of the initialization algorithm, the function **reInitiate.m** is required. In order to perform an RPEM run, the user is required to

1. Run the script **reInitiate.m**. That script prompts the user for *polynomialOrders* and *notUsedIndices*. The parameter vector at the end of the run, together with the previous and new degrees, are then used to re-initiate all relevant quantities of the RPEM.
2. Run the **SetupRPEM.m** file, as described in section 6.
3. Rerun **RPEM.m**. Note that the RPEM does not need to prompt the user for any further information this time.

Here, **reinitiate.m** is used in exactly the same way as described in section 6.3. Note, however, that the model structure is normally not changed when the initialization algorithm is used in combination with the RPEM, as the initial parameters are given for a specific model structure. By changing the model structure between the initialization and RPEM algorithms, the benefits of an initialization algorithm may be lost or significantly reduced.

### 13. Summary

This report describes a software package for identification of nonlinear systems. Future work in this field, that result in useful MATLAB routines, will be integrated with the presently available functionality. Updated versions of this report will then be made available.

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