Matlab code for Lyapunov exponents of fractional order systems

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In this paper the Benettin-Wolf algorithm to determine all Lyapunov exponents for a class of fractional-order systems modeled by Caputo's derivative and the corresponding Matlab code are presented. First it is proved that the considered class of fractional-order systems admits the necessary variational system necessary to find the Lyapunov exponents. The underlying numerical method to solve the extended system of fractional order, composed of the initial value problem and the variational system, is the predictor-corrector Adams-Bashforth-Moulton for fractional differential equations. The Matlab program prints and plots the Lyapunov exponents as function of time. Also, the programs to obtain Lyapunov exponents as function of the bifurcation parameter and as function of the fractional order are described. The Matlab program for Lyapunov exponents is developed from an existing Matlab program for Lyapunov exponents of integer order. To decrease the computing time, a fast Matlab program which implements the Adams-Bashforth-Moulton method, is utilized. Four representative examples are considered.

Keywords: Lyapunov exponents, Benettin-Wolf algorithm, Fractional-order dynamical system

1. Introduction

Despite a long history, the doubts that fractionalorder (FO) derivatives have no clear geometrical interpretations (see e.g. [Podlubny, 2002]), was one of the several reasons that fractional calculus was not used in physics or engineering. However, during the last more than 10 years, fractional calculus starts to attract increasing attention. There are nowadays more and more works on FO systems and their related applications in physics, engineering, mathematics, finance, chemistry, and so on. For the theory on the existence, uniqueness, continuous dependence on parameters and asymptotic stability of solutions of FDEs with general nonlinearities see, for example, [Oldham & Spanier, 1974; Caputo, 1967], and [Diethelm & Ford, 2002; Kilbas & Trujillo, 2001; Podlubny, 1999].

The Lyapunov exponents (LEs) measure the average rate of divergence or convergence of orbits starting from nearby initial points. Therefore, they can be used to analyze the stability of limits sets and to check sensitive dependence on initial conditions, that is, the presence of potential chaotic attractors. On the other side, in [Cvitanović *et al.*, 2016] Cvitanović et al. do not recommend the evaluation

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of the LEs and recommend to: "Compute stability exponents and the associated covariant vectors instead. Cost less and gets you more insight. [...] we are doubtful of their utility as means of predicting any observables of physical significance". Moreover, we additionally note here, Perron's counterexample [Leonov & Kuznetsov, 2007] which shows actually that the use of LEs, obtained via the linearization procedure, for the study of the behaviour of nonlinear system requires a rigorous justification. However, determining LEs remains the subject of many works and grown into a real software industry for modern nonlinear physics (see, e.g. [Hegger et al., 1999; Barreira & Pesin, 2001; Skokos, 2010; Czornik et al., 2013; Pikovsky & Politi, 2016; Vallejo & San-[uan, 2017] and others).

Nowadays there are two widely used definitions¹, of the LEs: via the exponential growth rates of norms of the fundamental matrix columns [Lyapunov, 1892] and via the exponential growth rates of the sigular values of fundamental matrix [Oseledets, 1968]. Corresponding approaches for the LEs computation and their difference are discussed, e.g., in [Kuznetsov *et al.*, 2018a, 2016].

Remark that in numerical experiments we can consider only finite time, and, thus, the numerically computed values of LEs can differ significantly from the limit values (e.g. if the considered trajectory belongs to a *transient chaotic set*), and are often referred as *finite-time LEs*.

Applying the statistical physics approach and assuming the ergodicity (see, e.g. [Oseledets, 1968]), the LEs for a given dynamical system are often estimated by local LEs along a "typical" trajectory. However, in numerical experiments, the rigorous use of the ergodic theory is a challenging task (see, e.g. [Cvitanović *et al.*, 2016, p.118]). If the LEs are the same for any trajectory, then Frederickson *et al.* [Frederickson *et al.*, 1983, p.190] suggested to call them as *absolute* ones and wrote that such absolute values rarely exist. For example, substantially different values of the local LEs can be obtained along trajectories on coexisting nonsymmetric attractors in the case of multistability² (see, e.g. such corresponding examples for the classical Lorenz system and Henon map [Leonov *et al.*, 2016; Kuznetsov *et al.*, 2018b]).

In order to study the chaoticity of an attractor in numerical experiments, one has to consider a grid of point covering the attractor and compute corresponding finite-time local LEs for a certain time. Remark that while the time series obtained from a *physical experiment* are assumed to be reliable on the whole considered time interval, the time series produced by the integration of *mathematical dynamical model* can be reliable on a limited time interval only due to computational errors.

Taking into account the above discussion further we consider finite-time local LEs and their computation in Matlab by the analog of the Bennetin-Wolf algorithm.

2. Benettin-Wolf Algorithm for LEs of FO

The determination of LEs of a system of integer or fractional order with the Benettin-Wolf algorithm, requires the numerical integration of differential equations of integer or fractional order. Because the purpose of this paper is to present a Matlab code for LEs of systems of FO, in the next subsections, only the most important steps (such as the existence of variational equations of FO) used to implement the algorithm in Matlab language are presented (theoretical details can be found in the related references).

2.1. Numerical integration of FDEs

The autonomous FO systems considered in this paper are modeled by the following Initial Value Problem (IVP) with Caputo's derivative

$$D_*^q x = f(x), x(0) = x_0,$$
(1)

for $t \in [0,T]$, $q \in (0,1)$, $f : \mathbb{R}^n \to \mathbb{R}^n$ and D^q_* , Caputo's differential operator of order q with starting

¹Relying on the Oseledec ergodic theorem [Oseledets, 1968] the above definitions often do not differ (see, e.g. Eckmann & Ruelle [Eckmann & Ruelle, 1985, p.620, p.650], Wolf et al. [Wolf *et al.*, 1985a, p.286, p.290-291], and Abarbanel et al. [Abarbanel *et al.*, 1993, p.1363, p.1364]), however in general case, they may lead to different values [Kuznetsov *et al.*, 2018a], [Bylov *et al.*, 1966, p.289], [Leonov & Kuznetsov, 2007, p.1083].

 $^{^{2}}$ While trivial attractors (stable equilibrium points) can be easily found analytically or numerically, the search of all periodic and chaotic attractors for a given system is a challenging problem. See, e.g. famous 16th Hilbert problem [Hilbert, 1901-1902] on the number of coexisting periodic attractors in two dimensional polynomial systems, which was formulated in 1900 and is still unsolved, and its generalization for multidimensional systems with chaotic attractors [Leonov & Kuznetsov, 2015].

 $^{^{3}}$ Based on philosophical arguments rather than a mathematical point of view, some researchers questioned the appropriateness

point 0^3

$$D^{q}_{*}x(t) = \frac{1}{\Gamma(1-q)} \int_{0}^{t} (t-\tau)^{-q} x'(\tau) d\tau,$$

with Γ the known Euler function.

Properties of the Caputo's differential operator, D^q_* , are discussed in [Podlubny, 1999; Gorenflo & Mainardi, 1997].

Under Lipschitz continuity of the function f, the IVP (1) admits a unique solution [Diethelm *et al.*, 2002].

Remark 2.1.

i) In the case of an integer-order dynamical system, denoting the solution of the underlying IVP as $x(t, x_0)$, one has $x_s \circ x_t = x_{t+s}$ (see e.g. [Zhou, 2016]). Due to the memory dependence of the derivatives, this does not hold in the case of systems modeled by FDEs. However, motivated by the numerical character of this paper, the definition of integer order dynamical systems which states that if the underlying IVP admits unique solutions existing on infinite time interval, the problem defines a dynamical system (see [Stuart & Humphries, 1998, Definition 2.1.2]) is adopted.

ii) Even fractional-order dynamics describe a real object more accurately than classical integer-order dynamics, systems modeled by the IVP (1) cannot have any non-constant periodic solution (see e.g. [Tavazoei & Haeri, 2009]). However, a solution may be asymptotically periodic [Danca *et al.*, 2018a]. These trajectories are called *numerically periodic*, in the sense that the trajectory, from numerical point of view, can be an extremely-near periodic with respect to, e.g., Euclidean norm [Danca *et al.*, 2018a]. A numerically periodic trajectory refers to as a closed trajectory in the phase space in the sense that the closing error is within a given bound of 1E - n, with n being a sufficiently large positive integer.

The numerical integrations required by the LEs algorithm for FO systems are performed in this paper with the predictor-corrector Adams-Bashforth-Moulton (ABM) method for FDEs, proposed Diethelm et al. [Diethelm *et al.*, 2002] which is constructed for the fully general set of equations without any special assumptions, being easy to implement in any language. Let us next assume that we are working on a uniform grid $\{t_j = jh : n = 0, 1, ..., N + 1\}$ with some integer N with the step size h and the case of $q \in (0, 1)$. Then, the predictor form, x^P , at the point t_{j+1} , is the fractional variant of the Adams-Bashforth method

$$x^{P}(t_{n+1}) = x_{0} + \frac{1}{\Gamma(q)} \sum_{j=0}^{n} b_{j,n+1} f(x(t_{j})),$$

while the corrector formula (the fractional variant of the one-step implicit Adams Moulton method) reads

$$x(t_{n+1}) = x_0 + \frac{h^q}{\Gamma(q+2)} f\left(x^P(t_{n+1})\right) + \frac{h^q}{\Gamma(q+2)} \sum_{j=0}^n a_{j,n+1} f(x(t_j)),$$

where a and b are the corrector and predictor weights respectively given by the following formula

$$a_{j,n+1} = \begin{cases} n^{q+1} - (n-q) (n+1)^q & \text{if } j = 0, \\ (n-j+2)^{q+1} + (n-j)^{q+1} & \\ -2 (n-j+1)^{q+1} & \text{if } 1 \le j \le n, \\ 1 & \text{if } j = n+1, \end{cases}$$

and

$$b_{j,n+1} = \frac{h^q}{q} \left((n+1-j)^q - (n-j)^q \right)$$

The predictor-corrector ABM method has an error which is roughly proportional to h^2 . Thus, to obtain an error of, e.g., 1.0E - 6, a step size close to h = 1.0E - 3 should be considered.

Because due to the long memory processes, the utilized ABM method as described in [Diethelm *et al.*, 2002], is time consuming. Therefore, a fast optimized ABM method, FDE12.m [Garrappa, 2012] is used.

FDE12.m is called by the following command line:

[t,x] = FDE12(q,fcn,t0,tf,x0,h);

where q represents the commensurate fractionalorder, fcn.m is the file with the function to be integrated, t0 and tf define the time span, x0 are the initial conditions, and h represents the integration step-size.

of using initial conditions of the classical form in the Caputo derivative [Hartley *et al.*, 2013]. However, it should be emphasized that, in practical (physical) problems, physically interpretable initial conditions are necessary and Caputo's derivative is a fully justified tool [Diethelm, 2014].

For example, consider the integration of the FO Rabinovich-Fabrikant (RF) system [Danca, 2016], which has the following Matlab function, RF.m

```
function dx = RF(t,x)
dx=[x(2)*(x(3)-1+x(1)*x(1))+0.1*x(1);
x(1)*(3*x(3)+1-x(1)*x(1))+0.1*x(2);
-2*x(3)*(p+x(1)*x(2))];
```

with the bifurcation parameter p = 0.98. With the following parameters, one obtains the chaotic attractor in Fig. 1 (a):

[t,x]=FDE12(0.999,@RF,0,1500,[0.1;0.1;0.1],... 0.01);

To note that FDE12.m requires the entry, x0, as a column vector [x10,x20,x30]' or, similarly, [x10;x20;x30]. Also, the returned exit, x, is a column vector.

2.2. Algorithm for LEs of the FO system (1)

Notation 2.1. Hereafter the finite-time local LEs of FO are called LEs.

The existence of the variational equations necessary to determine LEs is ensured by the following Theorem [Li *et al.*, 2010]

Theorem 1. System (1) has the following variational equations which define the LEs

$$D_*^q \Phi(t) = D_x f(x) \Phi(t),$$

$$\Phi(0) = I,$$
(2)

where Φ is the matrix solution of the system (1), D_x is the Jacobian of f and I is the identity matrix.

Further we assume that for the matrix $\Phi(t)$ the cocycle property takes place.

Therefore, the algorithm to determine the LEs of the system (1) becomes similar to the case of integer order.

Lyapunov exponents measure the exponential growth, or decay, of infinitesimal phase-space perturbations of a chaotic dynamical system.

The algorithm for numerical evaluation of LEs utilized in this paper, has been proposed in the seminal works of Benettin et al. [Benettin *et al.*, 1980] (see also [Shimada & Nagashima, 1979]), one of the the first work to propose a Gram-Schmidt orthogonalization procedure to compute LEs for continuous systems of integer order, as described in [Eckmann & Ruelle, 1985]), and by Wolf et al. [Wolf *et al.*, 1985b] (see also [Eckmann *et al.*, 1986]).

The algorithm to find all LEs, described as a Fortran code by Wolf et al. [Wolf *et al.*, 1985b], and also as a Basic code in [Baker & Gollub, 1990], solves the equations of motion under perturbations and periodic orthonormalization.

To note that the accuracy and reliability of numerically determined LEs depend on initial conditions, on the selection of the perturbations, performances of the utilized integration numerical method and also on orthonormalization step size. A longtime numerical calculation of the leading Lyapunov exponent requires rescaling the distance between nearby trajectories, in order to keep the separation within the linearized flow range. To avoid overflow, one calculates the divergence of nearby trajectories for finite timesteps and renormalizes to unity after a finite number of steps (Gram-Schmidt procedure [Eckmann & Ruelle, 1985; Christiansen & Rugh, 1997]).

Therefore, the main steps to determine numerically the LEs are: numerical integration of the FO system (1) together with the variational system (2) (i.e. the extended system), Gram-Schmidt procedure and picking up the exponents during the renormalization procedure, the LEs being determined as the average of the logarithm of the stretching factor of each perturbation, steps presented in Algorithm 1.

3. The Matlab code for LEs

Consider the following general assumptions:

- The considered systems, modeled by the IVP (1), are autonomous;
- The system (1) is of commensurate order: $q_1 = q_2 = \dots = q_{ne} = q;^4$
- In the case of chaotic behavior, the fractionalorder has been chosen close to 1, such that chaos is significant.
- The right hand side of system (1), f, is smooth enough.
- Because of the space restrictions, only the main programs code are presented, and indications on how to write the other ones.

⁴The incommensurate case can be treated similarly, the only difference referring to the utilized numerical method for FDEs.

Algorithm 1: Algorithm for LEs of FO system (1)

Input:							
-ne	\triangleright number of equations						
-x_start	$\triangleright n_e$ initial conditions of (1)						
$-t_start, t_end$	\triangleright time span						
-h_norm	\triangleright Normalization step-size						
$n_it \leftarrow (t_end - t_start)/h_norm$	\triangleright iterations number						
for $i \leftarrow ne+1$ to $ne(ne+1)$ do							
x(i) = 1.0	\triangleright initial conditions of (2)						
end							
$t \leftarrow t_start$							
for $i \leftarrow 1$ to $n_i t$ do							
$x \leftarrow \text{integration of FO systems (1)-(2)}$							
$t \leftarrow t + h_norm$							
$zn(1),, zn(ne) \leftarrow Gram-Schmidt pro$	cedure						
$s(1) \leftarrow 0$							
for $k \leftarrow 1$ to ne do	for $k \leftarrow 1$ to ne do						
$ s(k) \leftarrow s(k) + \log(zn(k))$	\triangleright vector magnitudes						
$LE(k) \leftarrow s(k)/(t - t_start)$	ightarrow LEs						
end							
end							
Output:LE							

A simple way to build in Matlab language the algorithm for FO systems, the program FO_Lyapunov.m (Appendix A), was to modify either some existing program, e.g., the program lyapunov.m [Govorukhin, 2004], which is a Matlab variant of the original LEs algorithm proposed in [Benettin *et al.*, 1980] or [Wolf *et al.*, 1985b] or, similarly, to translate in Matlab the BASIC program [Baker & Gollub, 1990], a close variant of the original algorithm) or, also, the Fortran code proposed by Wolf et all in [Wolf *et al.*, 1985b], and modify it for FDEs.

The program, called FO_Lyapunov.m, is launched with the following command line:

```
[t,LE]=F0_Lyapunov(ne,@ext_fcn,t_start,h_norm,...
t_end,x_start,h,q,out);
```

where **ne** represents the equations (and state variables) number, **ext_fcn.m** the function containing the extended system (1)-(2), **t_start** and **t_end** the time span, **h_norm** the normalization step, **x_start** the initial condition (as column vector), **h** the step size of FDE12.m, and **out** indicates the steps number when intermediate values of time and LEs are printed (for **out=0**, no intermediate results will be printed out).

As shown in the algorithm for LEs (Algorithm 1),

it is necessary to solve the extended system (1)-(2) of FO (yellow line) which is given in the function ext_fcn.m. In this file, beside the right hand side function f of the system (1), the Jacobi matrix J should be included (see Appendix B where the function for the RF system is presented).

The program plots the time evolutions of the LEs.

4. Numerical tests

Beyond numerical artifacts that might occur when numerically integrating a system of ODEs of integer order, notions such as "shadowing time" and "maximally effective computational time" reveal that it is possible to have reliable numerical simulations only on a relative finite-time interval (see, e.g., [Sarra & Meador, 2011; Wang *et al.*, 2012]). The case of FO systems is even more delicate. In this paper we have considered generally $t \in [0, 300]$ and, for the Lorenz system, $t \in [0, 500]$.

Let consider the function for the RF system, LE_RF.m, which include the extended system (1)-(2) (Appendix B). Because ne=3, beside the 3 variables x(1),x(2),x(3) required by the numerical solution of the original system (1), the matrix solution of the system (2) requires other more ne×ne=9 variables from

the total of ne(ne+1)=12 variables: x(1:12), f=zeros(size(x))=zeros(12), where f loads the first ne=3 righthand side expressions of system (1), and the $ne \times ne=9$ righthand side expressions of the variational system (2).

For example, for the RF system, with the following command line:

```
[t,LE]=F0_Lyapunov(3,@LE_RF,0,0.02,300,...
[0.1;0.1;0.1],0.005,0.999,1000);
```

one obtains the intermediary results printed every out=1000 h_norm steps, presented in Table 1 (see also Fig. 1 (b) where the dynamics of the LEs are drawn).

10.00	0.1611	0.0660	-2.1614	
20.00	0.1923	0.0069	-2.0503	
30.00	0.0984	-0.7397	-1.1817	
40.00	0.0248	0.0542	-1.9761	
50.00	0.0440	-0.0168	-1.9867	
60.00	0.0697	-0.0006	-2.0701	
70.00	0.0354	-0.0116	-2.0167	
80.00	0.0331	-0.0618	-1.9360	
90.00	0.0201	0.0125	-1.9754	
100.00	0.0429	0.0112	-1.9794	
110.00	0.0337	0.0252	-1.9698	
120.00	0.0262	-0.0213	-1.9038	
130.00	0.0624	-0.0043	-1.9537	
140.00	0.0660	-0.0029	-1.9811	
150.00	0.0645	-0.0010	-2.0008	
160.00	0.0743	0.0018	-2.0304	
170.00	0.0710	-0.0009	-2.0394	
180.00	0.0583	0.0139	-2.0548	
190.00	0.0678	0.0042	-2.0665	
200.00	0.0662	-0.0217	-2.0498	
210.00	0.0628	-0.0155	-2.0622	
220.00	0.0602	-0.0125	-2.0715	
230.00	0.0570	-0.0222	-2.0666	
240.00	0.0643	-0.0222	-2.0813	
250.00	0.0639	-0.0079	-2.1020	
260.00	0.0623	0.0045	-2.1121	
270.00	0.0630	0.0006	-2.0987	
280.00	0.0551	-0.0036	-2.0771	
290.00	0.0761	0.0009	-2.0937	
300.00	0.0749	0.0018	-2.0850	

Table 1. For $t \in [0, 300]$, the RF system has LE = (0.0749, 0.0018, -2.0850) (last line, blue).

2. If one considers the Lorenz system

$$D_{*}^{q}x_{1} = \sigma(x(2) - x(1)),$$

$$D_{*}^{q}x_{2} = -x(1)x(3) + px(1) - x(2) \qquad (3)$$

$$D_{*}^{q}x_{3} = x(1)x(2) - \beta x(3);$$

with q = 0.985 and the standard parameters $\sigma = 10, \beta = 8/3$ and the bifurcation parame-

ter p = 200, after some neglected transients, one obtains an apparently stable cycle (see Fig. 2 (a) and Remark 2.1 (*ii*)).

To obtain the LEs one write the following command line:

[t,LE]=F0_Lyapunov(3,@LE_Lorenz,0,5,500,... [0.1;0.1;0.1],0.001,0.985,10);

which gives LE=(-0.0026, -0.0870, -1.6225). The function LE_Lorenz.m can be obtained similarly with LE_RF.m. The time evolution of the LEs is presented in Table 2 (see also Fig. 2 (b)).

50.00	0.1759	-0.1591	-1.5683	
100.00	0.0611	-0.1108	-1.6050	
150.00	0.0346	-0.0927	-1.6300	
200.00	0.0215	-0.0877	-1.6288	
250.00	0.0135	-0.0866	-1.6269	
300.00	0.0082	-0.0865	-1.6255	
350.00	0.0043	-0.0866	-1.6244	
400.00	0.0014	-0.0867	-1.6236	
450.00	-0.0008	-0.0869	-1.6230	
500.00	-0.0026	-0.0870	-1.6225	

Table 2. Time evolution of the LEs for the Lorenz system. LE=(-0.0026, -0.0870, -1.6225).

3. Consider next the non-smooth 4-dimensional system [Danca *et al.*, 2018a]

$$D_{*}^{q}x_{1} = -x_{1} + x_{2}, D_{*}^{q}x_{2} = -x_{3}\operatorname{sgn}(x_{1}) + x_{4}, D_{*}^{q}x_{3} = |x_{1}| - a, D_{*}^{q}x_{4} = -bx_{2}.$$
(4)

with a = 1, b = 0.5 and q = 0.98. With the command line:

[t,LE]=F0_Lyapunov(4,@LE_4d,0,0.02,300,... [0.1;0.1;0.1],0.005,0.98,1000);

one obtains LE=(0.1262, 0.0846, 0.0778, -1.5244) (Fig. 3).

For this example, ne=4, the number of the variables is 20 and, therefore, compared with systems with ne=3, the function file, LE_4d.m, must be modified accordingly (compare the red line in LE_RF.m). Also, in order to obtain the printed intermediated values of LEs, in the FO_Lyapunov.m, the fprintf command, must be modified by adding one supplementary specifier %10.3f.

Because from the four LEs, the system admits

three positive LEs, it can be considered as hyperchaotic (see [Danca *et al.*, 2018b] for a discussion about the number of positive LEs of hyperchaotic systems).

Remark 4.1. Note that because the system is not smooth and also discontinuous, the correct numerical integration required for this system cannot be done without a previous smooth approximation [Danca *et al.*, 2018a]. Thus, like all numerical methods for FDEs which are designed for continuous dynamical systems, the integrator FDE12 cannot be utilized in this case without the mentioned approximation. Also, without a smooth approximation, the Jacobian matrix, utilized in the Benettin-Wolf algorithm, cannot be determined.

4. LEs can be also plotted as function of the bifurcation parameter p for $p \in [p_{min}, p_{max}]$ (program run_Lyapunov_p.m, Appendix C). The program uses a slightly modified variant of FO_Lyapunov.m, which is called FO_Lyapunov_p.m. To obtain FO_Lyapunov_p.m from FO_Lyapunov.m, the following modifications have to be done:

(a) The header line of the code FO_Lyapunov.m is replaced with the following line (to note that the output t and the input out are no more necessary)

function LE=F0_Lyapunov_p(ne,ext_fcn_p,...
t_start,h_norm,t_end,x_start,h,q,p);

(b) the printing and plotting lines (*) (red color) are deleted;

(c) the rest of the lines remain the same;

The m-file containing the extended system, ext_fcn_p.m, for the RF system, is presented in Appendix D. Passing the parameter p between these codes, can be easily realized due of the facilities of the program FDE12 (see FDE12).

For example, for the RF system, for $p \in [1.1, 1.3]$, for 1000 values of p (n = 1000), with the following command

run_F0_Lyapunov_p(3,@LE_RF_p,0,0.02,200,... [0.1;0.1;0.1],0.002,0.998,1.1,1.3,800)

one obtains the evolution of the LEs drawn in Fig. 4 (a).

5. LEs can be plotted also as function of the frac-

tional order q (program run_Lyapunov_q.m in Appendix E). The used program FO_Lyapunov_q, is obtained from FO_Lyapunov with the following modifications:

(a) The header line of the code is replaced with the following line (to note that the input **out** is no more necessary);

function [t,LE]=F0_Lyapunov_q(ne,ext_fcn,...
t_start,h_norm,t_end,x_start,h,q);

(b) the printing and plotting lines (*) (red color) are deleted;

(c) the rest of the lines remain the same;

The function containing the extended system, ext_fcn.m, does not require any modification. For example, for the RF system, with the command

run_F0_Lypaunov_q(3,@LE_RF,0,0.05,150,... [0.1;0.1;0.1],0.002,0.9,1,800)

one obtains the LEs plotted in Fig. 4 (b).

Remark 4.2. The presented programs can be optimized especially in the cases of FO_Lyapunov_p.m and FO_Lyapunov_q.m. A simple improvement was to use while loop (instead for) which for these two programs reduce substantially the computational time. However, every parameter p (or order q) step, several constant parameters are shared between programs, fact which slows significantly the programs. Therefore, supplementary optimization should be done.

6. Another potential application of the proposed LEs algorithm, is to represent the LEs as function of two variables: the order q and the bifurcation parameter p.

Let the Chen system

$$D_*^q x_1 = a(x_2 - x_1), D_*^q x_2 = (p - a)x_1 - x_1 x_3 + c x_2, D_*^q x_3 = x_1 x_2 - b x_3,$$

with parameters a = 35, b = 3 and q and pvariables. By considering $S_i := LE(q, c)$, for i = 1, 2, 3, for $q \in [0.9, 1]$ and $p \in [20, 30]$, the obtained surfaces are plotted in Fig. 5 (see [Danca *et al.*, 2018b] where the algorithm to obtain LEs as surfaces is described). One can see that there exists a unique positive LE (red surface, S_1) for

all values of the considered parameter c but only for some fractional order values q, for which S_1 is situated over the horizontal plane LE=0, where LEs are zero. Also, one can see that S_2 , which is almost identic to S_1 when the underlying LE1,2, are negative, becomes zero for a relative large values of q and p, when S_2 separates from S_1 and identify (with the underlying numerical error) with the plane LE=0.

5. Conclusion and discussion

Starting from an existing variant for integer-order system, in this paper, based on the Benettin-Wolf algorithm, we proposed a Matlab program to determine numerically finite-time local LEs for FO systems.

Due to inherent numerical errors, the algorithm should be utilized with precaution. As known, among the numerical errors, the results depends strongly on the initial conditions, time integration interval and, especially, on the renormalization step size h_norm .

The relative large numerical errors of the Benettin-Wolf algorithm, which for the considered examples was of order of 1.E - 2, can be observed in the cases when one knows that the system presents a numerically periodic cycle (see Remark 2.1 (*ii*)), when the maximum LE should be zero. For example, the Lorenz system, which for p = 200 presents such apparently stable cycle (see Fig. 2 (a)), has the maximum LE zero only with two precise decimals. Actually, in general, three precise decimals for zero LEs are extremely rare. Note that this zero value, appears only for h_norm=5 and a smaller integration step size, h = 0.001 and only for a larger time interval [0,500]. Therefore, if no special improvements of numerical integration are implemented, with Benettin-Wolf algorithm, for integer but also for fractional order systems, two or three decimals are the maximal expected number of decimals.

One of the most important algorithm variable, is h_norm, which determines the normalization moments influences the results. This dependence can be also deduced from the example of the RF system. With h=0.005 and h_norm=0.005 one obtains LEs=(0.0631, 0.0031, -2.0774), while with the same stapsize, h=0.005, but h_norm=0.5, LEs=(0.0643, 0.0026, -1.8254).

Since on our best knowledge, there is not criterion to choose precisely h_norm, the recommended way would be to realize several tests to choose the value for which slightly modifications does not change significantly the results.

In the case of fixed step-size integration numerical methods, like the considered FDE12, h_norm can be chosen as depending on the step size h. Therefore, as Fig. 6 shows, h_norm could be multiple of integration step size h (ε represents the perturbation between nearly two trajectories x and \bar{x}).

Regarding the Gram-Schmidt procedure, in order to speed up the code, one can use the QR decomposition based on the Householder transformation: [Q, R] = qr(A). To have matrix Rwith positive diagonal elements, one can additionally use Q = Q*diag(sign(diag(R))); R =R*diag(sign(diag(R))). See also [Ramasubramanian & Sriram, 2000].

The integration step-size of FDE12, h, plays an important role. As the documentation of the program specifies, there exists the possibility to increase the performances of the numerical integration, but in time integration detriment.

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Fig. 1. (a) A chaotic attractor of the RF system of FO, for q = 0.999. (b) Dynamics of the LEs.



Fig. 2. (a) An apparently stable cycle of the generalized Lorenz system of FO, for q = 0.985. (b) Dynamics of the LEs.

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Fig. 3. Dynamics of the LEs of the 4-dimensional system of FO (4).



Fig. 4. Perturbation and rescaling of a nearby trajectory, after every h_{norm} steps, considered as multiple of h (here $h_{norm} = 2h$, sketch).



Fig. 5. LEs of RF system. (a) LEs as function of q for $q \in [0.9, 1]$. (b) LEs as function of p for $p \in [1.1, 1.3]$.



Fig. 6. LEs of Chen's system of FO represented by function of two variables: q and parameter p. Surfaces S_i , for i = 1, 2, 3, represents LE(q, p).

Appendices

A. Program for LEs of FO

```
function [t,LE]=F0_Lyapunov(ne,ext_fcn,t_start,
    h_norm,t_end,x_start,h,q,out);
%
%
    Program to compute LEs of systems of FO.
%
%
    The program uses a fast variant of the
%
    predictor-corrector Adams-Bashforth-Moulton,
%
    "FDE12.m" for FDEs, by Roberto Garrappa:
%
%
    https://goo.gl/XScYmi
%
%
    m-file required: FDE12 and
%
    the function containing the extended system
%
    (see e.g. LE_RF.m).
%
%
    FO_Lyapunov.m was developed, by
%
    modifying the program Lyapunov.m,
%
    by V.N. Govorukhin:
%
%
    https://goo.gl/wZVCtg
%
%
    FO_Lyapunov.m, FDE12.m and LE_RF.m
%
    must be in the same folder.
%
%
    How to use it:
%
      [t,LE]=FO_Lyapunov(ne,ext_fcn,t_start,...
%
      h_norm,t_end,x_start,h,q,out);
%
%
    Input:
%
      ne - system dimension;
%
      ext_fcn - function containing the extended
%
      system;
%
      t_start, t_end - time span;
      h_norm - step for Gram-Schmidt
%
%
      renormalization:
%
      x_start - initial condition;
%
      outp - priniting step of LEs;
%
      ioutp==0 - no print.
%
%
    Output:
%
      t - time values;
%
      LE Lyapunov exponents to each time value.
%
%
    Example of use for the RF system:
%
     [t,LE]=F0_Lyapunov(3,@LE_RF,0,0.02,300,...
%
     [0.1;0;1;0.1], 0.005, 0.999, 1000);
%
%
    The program is presented in:
%
%
    Marius-F. Danca and N. Kuznetsov,
%
    Matlab code for Lyapunov exponents of
%
    fractional order systems
%
%
hold on;
```

```
% Memory allocation
x=zeros(ne*(ne+1),1);
x0=x;
c=zeros(ne,1);
gsc=c; zn=c;
n_it = round((t_end-t_start)/h_norm);
% Initial values
   x(1:ne)=x_start;
    i=1;
    while i<=ne
       x((ne+1)*i)=1.0;
        i=i+1;
    end
    t=t_start;
% Main loop
it=1;
while it<=n_it</pre>
%
       Solution of extended ODE system
        [T,Y] = FDE12(q,ext_fcn,t,t+h_norm,x,h);
       t=t+h_norm;
       Y=transpose(Y);
       x=Y(size(Y,1),:); %solution at t+h_norm
       i=1;
       while i<=ne</pre>
            j=1;
            while j<=ne;</pre>
               x0(ne*i+j)=x(ne*j+i);
                j=j+1;
            end;
           i=i+1;
       end:
%
       orthonormal Gram-Schmidt basis
       zn(1)=0.0;
       j=1;
       while j<=ne</pre>
           zn(1)=zn(1)+x0(ne*j+1)*x0(ne*j+1);
           j=j+1;
       end;
       zn(1)=sqrt(zn(1));
       j=1;
       while j<=ne</pre>
           x0(ne*j+1)=x0(ne*j+1)/zn(1);
            j=j+1;
        end
        j=2;
       while j<=ne</pre>
           k=1;
           while k<=j-1
               gsc(k)=0.0;
               1=1;
               while l<=ne;</pre>
                    gsc(k)=gsc(k)+x0(ne*l+j)*x0(ne*l+
                        k):
                   1=1+1;
                end
               k=k+1;
            end
```

B. Function LE RF.m

```
while k<=ne
               1=1:
               while l<=j-1
                   x0(ne*k+j)=x0(ne*k+j)-gsc(1)*x0(
                       ne*k+l);
                   1=1+1:
               end
               k=k+1;
           end:
           zn(j)=0.0;
           k=1;
           while k<=ne
               zn(j)=zn(j)+x0(ne*k+j)*x0(ne*k+j);
               k=k+1;
           end
           zn(j)=sqrt(zn(j));
           k=1;
           while k<=ne
               x0(ne*k+j)=x0(ne*k+j)/zn(j);
               k=k+1:
           end
           j=j+1;
       end
       update running vector magnitudes
       k=1;
       while k<=ne;</pre>
           c(k)=c(k)+log(zn(k));
           k=k+1;
       end;
       normalize exponent
       k=1:
       while k<=ne</pre>
           LE(k)=c(k)/(t-t_start);
           k=k+1;
       end
       i=1;
       while i<=ne</pre>
           j=1;
           while j<=ne;</pre>
               x(ne*j+i)=x0(ne*i+j);
               j=j+1;
           end
           i=i+1;
       end;
       x=transpose(x);
       it=it+1;
       print and plot the results
       if (mod(it,out)==0) %
                                  (*)
           fprintf('%10.2f %10.4f %10.4f...
            %10.4f\n',[t,LE]); % (*)
                                   (*)
       end; %
       plot(t,LE) %
                                   (*)
end
% displays the box outline around axes
xlabel('t','fontsize',16) %
                                   (*)
ylabel('LEs','fontsize',14) %
                                   (*)
set(gca,'fontsize',14)%
                                   (*)
box on;%
                                   (*)
line([0,t],[0,0],'color','k')%
                                  (*)
```

k=1.

%

%

%

```
function f=LE_RF(t,x)
```

```
%Output data must be a column vector
f=zeros(size(x));
```

%variables allocated to the variational equations X = [x(4), x(7), x(10);x(5), x(8), x(11); x(6), x(9), x(12)];

```
%RF equations
f(1)=x(2)*(x(3)-1+x(1)*x(1))+0.1*x(1);
f(2)=x(1)*(3*x(3)+1-x(1)*x(1))+0.1*x(2);
f(3) = -2 \times (3) \times (0.98 + \times (1) \times (2));
```

```
%Jacobian matrix
```

```
J=[2*x(1)*x(2)+0.1, x(1)*x(1)+x(3)-1, x(2);
    -3*x(1)*x(1)+3*x(3)+1,0.1,3*x(1);
    -2*x(2)*x(3),-2*x(1)*x(3),-2*(x(1)*x(2)+0.98)];
```

```
%Righthand side of variational equations
f(4:12)=J*X; % To be modified if ne>3
```

C. Program for LEs as function on p

```
function run_Lyapunov_p(ne,ext_fcn,t_start,h_norm,
    t_end,x_start,h,q,p_min,p_max,n);
hold on;
p_step=(p_max-p_min)/n
p=p_min;
while p<=p_max</pre>
   LE=F0_Lyapunov_p(ne,ext_fcn,t_start,h_norm,t_end
        ,x_start,h,q,p);
   p=p+p_step
   plot(p,LE);
end
```

D. Function LE_RF_p.m

```
function f=LE_RF_p(t,x,p)
%p is the parameter
f=zeros(size(x));
X = [x(4), x(7), x(10);
x(5), x(8), x(11);
x(6), x(9), x(12)];
%RF equations
f(1)=x(2)*(x(3)-1+x(1)*x(1))+0.1*x(1);
f(2)=x(1)*(3*x(3)+1-x(1)*x(1))+0.1*x(2);
f(3) = -2 * x(3) * (p + x(1) * x(2));
%Jacobian matrix
J=[2*x(1)*x(2)+0.1, x(1)*x(1)+x(3)-1, x(2);
-3*x(1)*x(1)+3*x(3)+1,0.1,3*x(1);
-2*x(2)*x(3), -2*x(1)*x(3), -2*(x(1)*x(2)+p)];
f(4:12)=J*X; % To be modified if ne>3
```

E. Program for LEs as function of q

while q<q_max
[t,LE]=F0_Lyapunov_q(ne,ext_fcn,t_start,h_norm,
 t_end,x_start,h,q);
q=q+q_step;
fprintf('q=%10.4f\n %10.4f', q);
plot(q,LE);
end</pre>