Chaos detection tools: application to a self-consistent triaxial model

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ABSTRACT
Together with the variational indicators of chaos, the spectral analysis methods have also achieved great popularity in the field of chaos detection. The former are based on the concept of local exponential divergence. The latter are based on the numerical analysis of some particular quantities of a single orbit, e.g. its frequency. In spite of having totally different conceptual bases, they are used for the very same goals such as, for instance, separating the chaotic and the regular components. In fact, we show herein that the variational indicators serve to distinguish both components of a Hamiltonian system in a more reliable fashion than a spectral analysis method does.

We study two start spaces for different energy levels of a self-consistent triaxial stellar dynamical model by means of some selected variational indicators and a spectral analysis method. In order to select the appropriate tools for this paper, we extend previous studies where we make a comparison of several variational indicators on different scenarios. Herein, we compare the average power-law exponent (APLE) and an alternative quantity given by the mean exponential growth factor of nearby orbits (MEGNO): the MEGNO’s slope estimation of the largest Lyapunov characteristic exponent (SE/LCE). The spectral analysis method selected for the investigation is the frequency modified Fourier transform (FMFT).

Besides a comparative study of the APLE, the fast Lyapunov indicator (FLI), the orthogonal fast Lyapunov indicator (OFLI) and the MEGNO/SE/LCE, we show that the SE/LCE could be an appropriate alternative to the MEGNO when studying large samples of initial conditions. The SE/LCE separates the chaotic and the regular components reliably and identifies the different levels of chaoticity. We show that the FMFT is not as reliable as the SE/LCE to describe clearly the chaotic domains in the experiments. We use the latter indicator as the main variational indicator to analyse the phase space portraits of the model under study.

Key words: methods: numerical – galaxies: kinematics and dynamics.

1 INTRODUCTION
The understanding of a realistic dynamical model is strongly related to the capability of identifying the chaotic and the regular nature of its orbits.

One one hand, one of the main features of chaotic orbits is their high sensitivity to the initial conditions (hereinafter i.c.), and the concept of local exponential divergence applies perfectly to identify such correspondence. On the other hand, the regular orbits do not show this behaviour. A seminal contribution to the field of chaos detection has been made by Lyapunov when he introduced the Lyapunov characteristic exponents, hereinafter LCEs (Oseledc 1968; Lyapunov 1992). The LCEs are theoretical quantities that measure the local rate of exponential divergence. Therefore, the LCEs are appropriate to distinguish between chaotic and regular motion.

The LCEs constitute the backbone of variational indicators of chaos (VICs). Moreover, their numerical implementation (Benettin et al. 1980; Froeschlé 1984; Tancredi, Sánchez & Roig 2001; Skokos 2010), i.e. the Lyapunov indicators (LIs), was a major contribution to the development of both fast and easy-to-compute new VICs. Nowadays, there is a plethora of VICs in the literature, such as the smaller alignment index, SALI (Skokos 2001; Skokos et al. 2004; Szell et al. 2004; Bountis & Skokos 2006; Carpintero 2008; Antonopoulos, Vasileios & Bountis 2010), and its generalized version, the generalized alignment index, GALI (Skokos, Bountis & Antonopoulos 2007, 2008; Manos & Athanassoula 2011); the mean exponential growth factor of nearby orbits, MEGNO (Cincotta & Simó 2000; Cincotta, Giordano & Simó 2003;
Giordano & Cincotta 2004; Goździewski, Konacki & Wolszczan 2005; Gayon & Bois 2008; Lemaître, Delisle & Valk 2009; Hinc et al. 2010; Compère, Lemaître & Delisle 2011; Maffione, Giordano & Cincotta 2011a); the Fast Lyapunov Indicator, FLI (Froeschlé, Gončzi & Lega 1997a; Froeschlé, Lega & Gončzi 1997b; Froeschlé & Lega 1998, 2000, 2006; Lega & Froeschlé 2001; Guzzo, Lega & Froeschlé 2002; Paleari, Froeschlé & Lega 2008; Todorović, Lega & Froeschlé 2008; Lega, Guzzo & Froeschlé 2010); its variant, the orthogonal fast Lyapunov indicator, OFLI (Fouchard et al. 2002); and a second-order variant of the OFLI, the OFLIT (Barrio 2005; Barrio, Blesa & Serrano 2009a; Barrio, Borczyk & Breiter 2009b; Barrio, Blesa & Serrano 2010). Furthermore, we can include the VICs based on the properties of dynamical spectra like the invariant spectra of stretching numbers or local Lyapunov characteristic numbers, LLCNs (Contopoulos et al. 1997; Lega & Froeschlé 1998), the invariant spectra of helicity or twist angles (Voglis & Contopoulos 1994; Contopoulos & Voglis 1996, 1997; Contopoulos et al. 1997; Froeschlé & Lega 1998; Lega & Froeschlé 1998; Voglis, Contopoulos & Efthymiopoulos 1998) and the spectral distance (Voglis, Contopoulos & Efthymiopoulos 1999). Finally, we include the relative Lyapunov indicator, RLI (Sándor, Erdi & Efthymiopoulos 2000; Sándor et al. 2004, 2007; Szell et al. 2004), which is not based on the evolution of the solution of the first variational equations as the others, but on the evolution of two different but very close orbits.

Notwithstanding the large number of VICs cited, we could also include alternative methods, for instance, the average power-law exponent, APLE (Lukes-Gerakopoulos, Voglis & Efthymiopoulos 2008), which is based on the concept of Tsallis entropy; evolutionary algorithms (Petalas et al. 2009); and the record could indeed continue.

Together with the VICs, the spectral analysis methods (SAMs), based on the numerical analysis of some particular quantities of a single orbit, e.g. its frequency, have also achieved great popularity in the field of chaos detection. Among the SAMs we find the method outlined by Laskar (1990): the frequency map analysis, FMA (Laskar, Froeschlé & Celletti 1992; Laskar 1993; Papaplioppou & Laskar 1996, 1998) and a modification of the latter in order to improve the precision in the computed frequencies and amplitudes, the frequency modified Fourier transform, FMFT (Sidlóchovský & Nesvorný 1997) or other alternatives using fast Fourier transform (FFT) techniques (Michtchenko & Ferraz-Mello 1995; Ferraz-Mello et al. 2005; Michtchenko et al. 2010), among others. As an easy identification of the important resonances (regions of phase space occupied by orbits with commensurable orbital frequencies) is possible with the SAMs, they are able to identify the major resonant orbit families. The relative importance of each family regarding the phase space can be assessed from the number of orbits associated with the particular resonance the family belongs to. That is, the SAMs are not only suitable to determine the resonance web like the VICs (Kaneko & Konishi 1994; Cincotta et al. 2003; Froeschlé, Lega & Guzzo 2006; Lukes-Gerakopoulos et al. 2008) but also to identify the individual resonances by their frequency vectors.

Though the conceptual bases of the VICs and the SAMs are totally different, many researchers use both types of techniques for similar goals. Nevertheless, we show herein that this seems to be a no reliable approach to study some important aspects of the dynamics of a Hamiltonian system. Moreover, in the literature there are plenty of evidence that the complementary use of both types of techniques results in an efficient and solid way to gather dynamical information from a Hamiltonian system (Froeschlé et al. 1997b; Froeschlé & Lega 1998; Guzzo et al. 2002; Kalapotharakos & Voglis 2005; Barrio et al. 2009b). For instance, in Froeschlé et al. (1997b) the authors compare the sensitivity of the FLI and the FMA on the standard map and on the Hénon & Heiles (1964) potential. In Guzzo et al. (2002) the authors use the FLI and the analytically filtered Fourier analysis, AFFA (Guzzo & Benettin 2001), another example of SAM, to determine the value of the critical parameter at which the transition from the unfortunately called Nekhoroshev to the Chirikov regime occurs in a quasi-integrable Hamiltonian model and a standard four-dimensional map. In Kalapotharakos & Voglis (2005), the authors use a measure related with the LCEs, the alignment indices (Voglis et al. 1998, 1999; Skokos 2001) and an index computed with the FMFT to study two different N-body models simulating elliptical galaxies.

One of the aspects that make a research accurate is the reliability of the tools available for the analysis. This reliability is a measure of, for instance, the confidence and efficiency of such tools. Regarding chaos detection, we believe that it is the diversity of the employed techniques that the fruitfulness of the research relies on. Therefore, we started a series composed of four papers in order to achieve the best combination of tools to study a given dynamical problem. In the first paper, Maffione et al. (2011a), we test the MEGNO against the most widely used chaos detection technique in the literature, which is the LI, on a somewhat realistic problem: a self-consistent triaxial stellar (ScTS) dynamical model (Muzzio, Carpintero & Wachlin 2005; Cincotta, Giordano & Muzzio 2008). In the second and third papers, Maffione et al. (2011b, hereinafter M11) and Darriba et al. (2012, hereinafter D12), we make a comparison of most of the VICs mentioned earlier on rather simple problems, i.e. on mappings and on the model of Hénon & Heiles (1964), respectively. Our purpose here is to extend the previous comparisons of VICs to alternative techniques, including a SAM. Further, this last paper of the series is regarded as a closing report on the subject and the conclusions comprise the whole investigation. We continue with the ScTS model and study two start spaces for four different energy levels by means of several chaos detection tools. In order to select appropriate VICs for the study, we consider the results of M11 and D12 and add a rather new technique, the APLE, and an alternative quantity given by the MEGNO: the MEGNO’s slope estimation of the largest LCE (SE/LCE) to their comparison. The MEGNO is part of the suggested VICs to compose the ClsF<sup>1</sup> in D12. In fact, we are interested in studying if its variant, i.e. the SE/LCE, is more reliable than the MEGNO on its own.

The study is organized as follows: in Section 2 we describe the ScTS model and introduce the <span>\( p_{x_0} - p_{z_0} \) and the <span>\( x_0 - z_0 \) start spaces. In Section 3, we briefly introduce the chaos detection techniques to be used within the reported research. We start with the VICs: the LI, the MEGNO and the SE/LCE (Section 3.1); the FLI and the OFLI (Section 3.2) and the APLE (Section 3.3). We finish with a short FMFT introduction (Section 3.4). In Section 4, we deal with the capability of the FMFT to resolve the different structures in a divided phase space (Section 4.1) and compare its performance with the SE/LCE in both start spaces of the ScTS model (Section 4.2). In Section 5, we study the efficiency of several VICs on the <span>\( p_{x_0} - p_{z_0} \) start space to decide which ones are appropriate to investigate the dynamics of the ScTS model (Sections 5.1 and 5.2). In Section 6 we analyse for four different energy surfaces, the <span>\( p_{x_0} - p_{z_0} \) (Section 6.1) and the <span>\( x_0 - z_0 \) (Section 6.2) start spaces by means of a complementary use of the VICs selected in the previous section.

<sup>1</sup>ClsF: ‘chaos indicators’ function’. It is a minimal and efficient package of variational indicators to study a general Hamiltonian.
Finally, Section 7 summarizes the main conclusions of the whole investigation.

As the extensive use of acronyms in the text might be overwhelming for the reader, in Table 1 we list in alphabetical order the most important acronyms used in this paper.

### Table 1. List of the most important acronyms in alphabetical order.

| Acronym | Definition |
|---------|------------|
| APLE | Average power-law exponent |
| CIoF | Chaos indicators function |
| FLI | Fast Lyapunov indicator |
| FMA | Frequency map analysis |
| FMFT | Frequency modified Fourier transform |
| LCEs | Lyapunov characteristic exponents |
| LIs | Lyapunov indicators |
| MEGNO | Mean exponential growth factor of nearby orbits |
| OFLI | Orthogonal fast Lyapunov indicator |
| SE/LCE | Slope estimation of the largest LCE |
| ScTS model | Self-consistent triaxial stellar model |
| VICs | Variational indicators of chaos |

2 THE MODEL AND THE START SPACES

In order to obtain a fairly realistic dynamical model of an elliptical galaxy, in Muzzio et al. (2005) the authors follow the cold dissipationless collapse of 100 000 particles randomly distributed following a 1/r law within a sphere. The velocities were randomly chosen from a spherical Gaussian distribution but only their tangential component was retained. After the system had relaxed, there remained 86 818 particles resembling an elliptical galaxy: the ScTS model. The system observes the de Vaucouleurs law shown by Muzzio et al. (2005) in their fig. 2. The model reproduces many dynamical characteristics of real elliptical galaxies, such as mass distribution, flattening, triaxiality and rotation (Muzzio 2006). The ScTS model has a strong triaxiality and a flattening that increases from the border of the system to its centre (see table 1 in the same paper). The resulting triaxial potential has semi-axes $X$, $Y$, $Z$ satisfying the condition $X > Y > Z$, and its minimum, which is close to $-7$, matches the origin. The potential is less flattened than the mass distribution, as expected. See table 1 in Muzzio et al. (2005).

The ScTS model is an $N$-body potential which is neither smooth nor stationary. As the authors needed to compute the orbits and their corresponding LIs, they froze the potential and represented it with a quadrupolar approximation. The equation that reproduces the potential is

$$V(x) = - f_n(x) - f_i(x) \cdot (x^2 - y^2) - f_z(x) \cdot (z^2 - y^2)$$

with $x = (x, y, z)$ and where

$$f_n(x) = \frac{\alpha_n}{[p_n^{0.085} + \delta_n^{0.04}]^{\alpha_n}},$$

(1)

with $p_n^2$ the square of the softened radius given by $p_n^2 = x^2 + y^2 + z^2 + \varepsilon$ when $n = 0$, or $p_n^2 = x^2 + y^2 + z^2 + 2 \cdot \varepsilon^2$ for $n = x, z$, and $\alpha_n, \delta_n, \alpha_i, \alpha_z$ are constants.

The selected value for the softening parameter is $\varepsilon \simeq 0.01$ for any $n$. The adopted values for the constants $\alpha_n, \delta_n, \alpha_i, \alpha_z$ and $ac_n$ are given in Table 2.

The stationary character of the parameters given in Table 2 was tested by performing several fits at different times after virialization, resulting in a precision of 0.1 per cent. For further details on the ScTS model, refer to Muzzio et al. (2005) and Cinccotta et al. (2008).

We have already used the same model to test the MEGNO in a previous study (Maffione et al. 2011a).

The spaces of i.c. used in this investigation are the $p_{10} - p_{20}$ and the $x_0 - z_0$ start spaces for four energy surfaces: $-0.1, -0.3, -0.5$ and $-0.7$. The $p_{20} - p_{30}$ start space is a set of i.c. of the form $(p_{20}, p_{30})$ with $x_0 = y_0 = z_0 = 0$ while $p_{20}$ is restrained by the energy condition (Papaphilippou & Laskar 1998). The $x_0 - z_0$ start space is a set of i.c. of the form $(x_0, z_0)$ with $y_0 = 0$, $p_{20} = p_{30} = 0$ and $p_{20}$ is restrained by the energy condition (Schwarzschild 1993). On one hand, the combination of both start spaces is an adequate choice to sample many of the different orbital families within the triaxial model (Schwarzschild 1993; Papaphilippou & Laskar 1998). On the other hand, the stable boxlets (i.e. resonant box orbits) which are centrophobic are not well represented in our $p_{20} - p_{30}$ start space.

Therefore, the $p_{10} - p_{20}$ and the $x_0 - z_0$ start spaces of the ScTS model seem to provide fairly realistic scenarios for testing many characteristics of both the VICs of the package and the FMFT.

### Table 2. Adopted values for the coefficients of the functions $f_n$ given by equation (1).

| $n$ | $\alpha$ | $\alpha_i$ | $\alpha_z$ | $ac_n$ |
|-----|----------|------------|------------|--------|
| 0   | 0.920 126 57 | 1.15 | 0.1340 | 1.037 665 79 |
| x   | 0.085 265 04 | 0.97 | 0.1283 | 4.615 715 81 |
| z   | $-0.058 710 11$ | 1.05 | 0.1239 | 4.420 309 43 |

3 THE TECHNIQUES

The aim of this section is to briefly introduce the techniques we will use in the forthcoming study.

3.1 The LI, the MEGNO and the SE/LCE

Considering a continuous dynamical system defined on a differentiable manifold $S$, where $\Phi(x) = x(t)$ characterizes the state of the system at time $t$, being the state of the system at time $t = 0$, $x(0) = x_0$. The state of the system after two consecutive time steps $t$ and $t'$ will be given by the composition law $\Phi^{t+t'} = \Phi^t \circ \Phi^{t'}$.

Moreover, the tangent space of $x$ maps on to the tangent space of $\Phi(x)$ according to the operator $d_x \Phi$ and following the rule $u(t) = d_x \Phi (u(0))$, where $u(0)$ is an initial deviation vector (hereinafter i.d.v.). The action of such operator at consecutive time intervals satisfies the equation

$$d_x \Phi^{t+t'} = d_x \Phi^{t'} \circ d_x \Phi^t.$$

If we suppose that our manifold $S$ has some norm denoted by $\| \cdot \|$, we can define the useful quantity

$$\lambda_i(x) = \frac{\| d_x \Phi^t (u) \|}{\| u \|}$$

called ‘growth factor’ in the direction of $u$.

Let $H(p, q)$ with $p, q \in \mathbb{R}^N$ be an $N$-dimensional Hamiltonian, that we suppose autonomous just for the sake of simplicity. Introducing the following notation:

$$x = (p, q) \in \mathbb{R}^{2N}, \quad f(x) = (-\partial H/\partial q, \partial H/\partial p) \in \mathbb{R}^{2N},$$

the equations of motion can be written in a simple way as

$$\dot{x} = f(x).$$

(2)
Let γ(x0; t) be an arc of an orbit of the flow (2) over a compact energy surface: M0 ⊂ ℝ2N, M0 = {x : H(p, q) = h}, then
\[ γ(x0; t) = \{x(x0; t') : x_0 \in M_0, 0 \leq t' < t}. \]

We can gain fundamental information about the Hamiltonian flow in the neighbourhood of any orbit γ through the largest LCE: ILCE, defined as
\[ χ_γ = χ[γ(x0; t)] = \lim_{t \to \infty} \frac{1}{t} \ln λ_1[γ(x0; t)] \]
with
\[ λ_1[γ(x0; t)] = \frac{∥d_x \Phi^t(\mathbf{w})∥}{∥\mathbf{w}∥}, \]
where \( ∥d_x \Phi^t(\mathbf{w})∥ \) is an ‘infinitesimal displacement’ from γ at time t. The fact that the ILCE (and its truncated value, the so-called LI = \( \lim_{t \to T} \frac{1}{t} \ln λ_1[γ(x0; t)] \) for T finite) measures the local mean exponential rate of divergence of nearby orbits is clearly understood when equation (3) is written in an integral fashion:
\[ χ_γ = \lim_{t \to \infty} \frac{1}{t} \int_0^t \frac{∥d_x \Phi'(\mathbf{w})∥}{∥d_x \Phi'(\mathbf{w})∥} \, dt' = \frac{∥d_x \Phi'(\mathbf{w})∥}{∥d_x \Phi'(\mathbf{w})∥}, \]
where the bar denotes time average.

Finally, the orbit is classified as regular or chaotic if the LI tends to zero (as \( \ln (t)/t \)) or to a positive value, respectively [a detailed discussion on the theory and computation of the LCEs can be found in the review of Skokos (2010)].

Now, we can introduce the quantity \( Y[γ(x_0; t)] \) through the expression
\[ Y_γ = Y[γ(x_0; t)] = \frac{2}{t} \int_0^t \frac{∥d_x \Phi'(\mathbf{w})∥}{∥d_x \Phi'(\mathbf{w})∥} \, dt', \]
which is related to the integral in equation (4), i.e. in the case of an exponential increase of \( ∥d_x \Phi'(\mathbf{w})∥, \frac{∥d_x \Phi'(\mathbf{w})∥}{∥d_x \Phi'(\mathbf{w})∥} = ∥\mathbf{w}∥ \cdot \exp(\chi_γ/t) \), the quantity \( Y_γ \) can be considered as a weighted variant of the integral in equation (4). Note that the quantity \( \hat{χ}_γ = Y_γ/t \) verifies \( \hat{χ}_γ \sim 2/t \) for regular motion and \( \hat{χ}_γ \sim χ_γ \) for chaotic motion, with \( t \to \infty \), which show that, in the case of regular motion \( \hat{χ}_γ \) converges to 0 faster than \( χ_γ \) does (which goes to zero as \( \ln t/t \)), while for chaotic motion both magnitudes approach the positive ILCE at a rather similar rate.

Instead of using the instantaneous rate of increase, \( χ_γ \), we average the logarithm of the growth factor, \( \ln(λ_1) = χ_γt \). Introducing the time average (MEGNO)
\[ \overline{Y}_γ = \overline{Y}[γ(x_0; t)] = \frac{1}{t} \int_0^t Y_γ \, dt', \]
we notice that the time evolution of the MEGNO can be briefly described in a suitable and unique expression for all kinds of motion. Indeed, its asymptotic behaviour can be summarized as follows:
\[ \overline{Y}_γ \approx a_γt + b_γ, \]
where \( a_γ = 0 \) and \( b_γ \approx 2 \) for quasi-periodic motion, while \( a_γ = χ_γ/2 \) and \( b_γ \approx 0 \) for irregular, stochastic motion. Deviations from the value \( b_γ \approx 2 \) (and \( a_γ = 0 \)) indicate that γ is close to some particular objects in phase space, being \( b_γ \leq 2 \) or \( b_γ > 2 \) for stable periodic orbits (resonant elliptic tori) or unstable periodic orbits (hyperbolic tori), respectively.

Refer to figs 1(c) and (d) from Cincotta et al. (2003) to see the general behaviour of the MEGNO for regular and chaotic orbits.

It is possible to estimate the LI of the orbit from equation (5) if we apply a simple linear least-squares fitting on the MEGNO (Cincotta & Simó 2000; Cincotta et al. 2003). This estimation is the earlier mentioned SE/LCE indicator. The least-squares fitting at the end of the integration process uses the last 80 per cent of the points in order to avoid the initial transient. Thus, the SE/LCE almost considers the full history of the orbit which makes it a very sensitive VIC.

The performances of the MEGNO’s time evolution curves to characterize the different levels of stability and chaoticity of the orbits shown in M11 and D12 were excellent. On the other hand, the description of the regular components in a divided phase space by means of the final values of the MEGNO (i.e. the values of the indicator at the end of the total integration time) was not satisfactory. The SE/LCE seems to be an appropriate alternative to solve the problem.

Fig. 1 shows the performances of the LI, the MEGNO and the SE/LCE for two sets of i.c., one of them located in the \( p_{x_0} - p_{z_0} \) start space (top panel) and the other set located in the \( x_0 - z_0 \) start space (bottom panel). In order to ensure that the VICs are well computed for a given orbit, the total integration time should verify \( T > T_0(E) \), where the function \( T_0(E) \) is some characteristic time-scale which depends on the energy surface \( E \). The function \( T_0(E) \) is taken as the period of the stable X-axis periodic orbit and we consider the energy surface \( E = -0.7 \) for both sets of i.c. \( (T_0(-0.7) \approx 7 \) units of time — hereinafter u.t. — see Maffione et al. 2011a). Therefore, on fixing the condition \( T \geq 10^3 T_0(E) \) to obtain reliable values for the VICs, we conclude that a total integration time of \( 7 \times 10^3 \) u.t. would be appropriate for the experiment (Maffione et al. 2011a). In the \( p_{x_0} - p_{z_0} \) start space, the set is composed of 790 i.c. of the

![Figure 1. Final values of the LI, the MEGNO and the SE/LCE. The i.c. were taken in the \( p_{x_0} - p_{z_0} \) start space (top panel) and in the \( x_0 - z_0 \) start space (bottom panel). Both sets of i.c. were taken on the energy surface \( -0.7 \) and the VICs were integrated for \( 7 \times 10^3 \) u.t. The values of the indicators are in logarithmic scale.](http://mnras.oxfordjournals.org/)

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form \( p_{x_0} \in [0 : 2.5] \) and \( p_{y_0} = 0 \). In the \( x_0 - z_0 \) start space, the set is composed of 646 i.c. of the form \( z_0 \in [0 : 0.7] \) and \( x_0 = 0 \).

In Fig. 1, the SE/LCE and the LI show very similar final values for chaotic orbits. The MEGNO grows exponentially fast and reaches higher final values (\( \gtrsim 1 \)) than the SE/LCE or the LI. The MEGNO clearly distinguishes between different levels of chaoticity. Consequently, the SE/LCE seems to be an unnecessary alternative to the MEGNO for describing the chaotic component. In the case of a divided phase space, where regular and chaotic components are mixed, the asymptotic behaviour of MEGNO’s final values for regular orbits may hide their different stability levels (see M11 and D12). The SE/LCE tends to zero for regular orbits. Moreover, the SE/LCE tends to decrease faster than the LI for regular motion (Cincotta & Simó 2000; Cincotta et al. 2003). So, no matter if the phase space is strongly divided, this behaviour allows the SE/LCE’s final values to show certainly (and faster than the LI) different levels of stability for the regular orbits. Thus, it seems to be appropriate to use the time evolution curves of the MEGNO for the individual analysis of the orbits (M11 and D12) and the final values of the SE/LCE as a global VIC to obtain the portraits of divided phase spaces (see Sections 4.2, 6.1 and 6.2).

The MEGNO has a low computational cost and the linear least-squares fitting in the computation of the SE/LCE does not require a significant amount of extra time. Hence, the SE/LCE is an alternative quantity that also has a low computational cost.

### 3.2 The FLI and the OFLI

The FLI is a quantity intimately related to the ILCE (Froeschlé et al. 1997a,b; Froeschlé & Lega 2000, 2006; Lega & Froeschlé 2001; Guzzo et al. 2002) and the MEGNO (Mestre, Cincotta & Giordano 2011). The FLI is able to distinguish between regular and chaotic (weakly chaotic) motion (Froeschlé et al. 1997a,b) and also between resonant and non-resonant motion (Froeschlé & Lega 2000; Lega & Froeschlé 2001; Guzzo et al. 2002) using only the first part of the computation of the ILCE.

Here, we use the initial definition of the FLI given in Froeschlé et al. (1997a,b) like in D12, i.e. we use a basis of i.d.v. for its computation. The definition given in Froeschlé & Lega (2000) uses only one i.d.v., the results do not change. Nevertheless, the computing time is obviously reduced (Froeschlé & Lega 2000).

On an \( N \)-dimensional Hamiltonian \( H \), we follow the time evolution of \( 2N \) unit i.d.v. (e.g. the ScTS model is three dimensional and the basis consists of six i.d.v.). The FLI at time \( t \) is defined by the highest norm among the unit i.d.v. of the basis, as follows:

\[
FLI(t) = \sup_i \| u(t)_i \|, \| u(t)_2 \|, \ldots, \| u(t)_{2N} \|.
\]

For further details, refer to Froeschlé et al. (1997a,b).

For both regular and chaotic motion, the FLI tends to infinity as time increases, but with completely different rates. The FLI grows linearly with time for regular motion and it grows exponentially fast for chaotic motion. Furthermore, the FLI grows linearly with time, but with different rates in the case of resonant regular orbits and non-resonant ones. Although the curves corresponding to resonant and non-resonant motions are separated, the oscillations due to the distortion of the orbits themselves may hide the distinction between the libration islands and the tori. Therefore, in Froeschlé & Lega (2000) the authors not only introduce the FLI using only one i.d.v. but also replace the definition of the FLI by its running average. We also applied an average to smooth the curves of the FLI.

In the case of the OFLI\(^2\) we take the orthogonal component for the flow for each unit i.d.v. \( u(t)_i \), \( i = 1, \ldots, 2N \) of the basis at every time step and retain the largest component among them:

\[
OFLI(t) = \sup_{i} \left[ u(t)_1^2, u(t)_2^2, \ldots, u(t)_{2N}^2 \right].
\]

This modification makes the OFLI a VIC which can easily distinguish periodicity among the orbits of the regular component. The OFLI for periodic orbits oscillates around a constant value, while for quasi-periodic motion and chaotic motion it has the same behaviour as the FLI.

Refer to figs 5 and 7 from Fouchard et al. (2002) to see the general behaviour of the FLI and the OFLI for regular and chaotic orbits.

### 3.3 The APLE

In the early study of Tsallis, Plastino & Zheng (1997) the authors show that when a non-linear dynamical system is in the regime of the so-called edge of chaos, the \( q \)-entropy rate of increase remains constant for a long time (metastable states). Moreover, they discuss that this behaviour may be associated with a power law rather than with exponential sensitivity of the orbits to the i.c. The interested reader may refer to Tsallis et al. (2002) for a ‘pedagogical guided tour’ on the subject.

In Lukes-Gerakopoulos et al. (2008) the authors find that the time evolution of the i.d.v. for a weakly chaotic orbit \( y(x_0; t) \) (i.e. a combination of a linear and an exponential law: \( y(t) = c_y t + e^{p_y t} \), with \( c_y \gg 1 \) and \( p_y \ll 1 \)) justifies theoretically why metastable states with a constant rate of increase of the \( q \)-entropy appear in the non-linear dynamical system. They argue that this law is reflected upon a transient behaviour in which the growth of the i.d.v. is almost a power law: \( y(t) \propto t^{p_y} \), with \( p_y > 1 \) (and the exponent \( p_y \)) can be associated with a \( q \)-exponent via the relationship \( p_y = 1/(1-q) \).

In fact, they introduce a method to compute the \( q \)-exponent and use it as a VIC (the APLE) in order to distinguish regular orbits from nearby weakly chaotic orbits.

For an \( N \)-dimensional Hamiltonian \( H \), consider a partitioning of the \( 2N \)-dimensional phase space \( \mathcal{S} \) into a large number of volume elements of size \( \delta^{2N} \) for some small \( \delta \) and let \( x(0) \) be the i.c. of an orbit located in a particular volume element. The authors introduce the APLE, \( p_y \), as follows:

\[
p_y = \frac{\ln \left( \frac{\| u(t) \|}{\| u(t)|^2} \right)}{2 \ln (\frac{1}{\delta})},
\]

where \( |u(t)|^2 = \sum_{k=1}^{m} |w_k(t)|^2 \) and \( w_k(t) \) is one of the \( m \) deviation vectors of an orthogonal basis \( \{w_k(t)\} \) of the tangent space to \( \mathcal{S} \) at the initial point \( x(0) \). Every \( w_k(t) \) has length greater than or equal to \( \delta \) and \( t_i \) is a transient initial time of orbit evolution.

Therefore, in the case of regular motion, \( p_y \) tends to 1, while for chaotic motion it grows exponentially fast. Moreover, the APLE shows oscillations around a constant value for a transient time interval before the exponential growth for weakly chaotic orbits. The length of the time interval is directly connected with the hyperbolicity of the local phase space.

Refer to fig. 2 from Lukes-Gerakopoulos et al. (2008) to see the general behaviour of the APLE for regular and chaotic orbits.

\(^2\) Let us remark that the following definition of the OFLI is slightly different from the definition of the OFLI given in Fouchard et al. (2002) where the authors use only one i.d.v.
3.4 The FMFT

The detailed description of the FMFT\(^3\) has been published in \(\text{Sidlichovský \& Nesvorný} (1997).\) However, we should mention the conceptual aspects of this technique. Since the main advantage of the FMFT is that it improves the computation of the amplitudes and the frequencies given by the FMA (Laskar 1990, 1993, 1999, 2003; Laskar et al. 1992; Papaphilippou \& Laskar 1996, 1998), we decided to introduce only a short description of the latter.

Let \(\mathcal{H}\) be a non-integrable Hamiltonian which is not non-degenerate:

\[
\det \left( \frac{\partial v(I)}{\partial I} \right) = \det \left( \frac{\partial^2 \mathcal{H}(I)}{\partial I^2} \right) \neq 0,
\]

with \(I \in B \subset \mathbb{R}^N\) the action-like variables and \(v\) the frequency vector. The Kolmogorov-Arnold-Moser theory states that for sufficient small values of the perturbation parameter \(\epsilon\), there exists a Cantor set \(\Omega\) of \(v\) that satisfies the Diophantine condition:

\[
\|k \cdot v\| > \frac{\kappa_v}{|k|^{m}},
\]

where \(\kappa_v\) and \(m\) are constants and \(|k|\) is the norm of \(k\). Furthermore, the system still possesses invariant tori with linear flow (KAM tori). The solutions lie on these invariant tori and are given in complex variables through their Fourier series:

\[
z_j(t) = z_0e^{i\nu_j\cdot xt} + \sum_m a_m(v)e^{i(m\cdot v)t},
\]

where the coefficients \(a_m(v)\) depend smoothly on the frequencies \(\nu_i, i = 1, \ldots, N\). If we keep all the angle-like variables fixed (\(\theta \in \Pi^N, \Pi^N\) is the \(N\)-dimensional torus, fixed to \(\theta = \theta_0\)), we obtain a frequency map \(\mathcal{F}_{\theta_0} : B \rightarrow \Omega_{\nu}; \ I \rightarrow p_2(\Psi^{-1}(\theta_0, I))\),

\[
\mathcal{F}_{\theta_0} : B \rightarrow \Omega_{\nu}; \ I \rightarrow p_2(\Psi^{-1}(\theta_0, I)),
\]

where \(p_2\) is the projection on \(\Omega_{\nu}\). \(\Psi(\phi, \nu) = v\). The FMA numerically gives through an iterative process the frequency map \(\mathcal{F}\) (i.e., amplitudes and frequencies) defined over the whole domain \(B\), which coincides, within the numerical precision adopted, with the \(\mathcal{F}_{\theta_0}\) of equation (6) in the set of KAM tori. The frequency map \(\mathcal{F}\) is obtained seeking for the quasi-periodic approximations of the solutions over a finite time-span through a finite series of terms:

\[
z_j(t) = z_0e^{i\nu_j\cdot xt} + \sum_{k=1}^s a_m(v)e^{i(m\cdot v)t},
\]

Once we obtain the quasi-periodic approximation from equation (7), we are able to establish a correspondence between the action-like variables \((I \in B)\) and the rotation numbers (or the frequency vector \(v \in \Omega_{\nu}\)). This correspondence is the so-called FMA (Laskar 1990). Regular regions of the phase space allow a very accurate estimation of real rotation numbers in a given time interval (for instance, \(\alpha 1/T^3\) using the MFT with the Hanning window, \(\alpha 1/T^2\) using the MFT without the Hanning window, \(\alpha 1/T\) using the usual FFT techniques, where \(\{-T; T\}\) is set as the whole time-span), and should not differ from the rotation numbers obtained in another time interval, considering a certain accuracy. Strictly speaking, the frequencies are only defined on KAM tori, the FMA numerically estimates over a finite time-span a frequency vector for any i.c., though. If the estimates of the rotation numbers given by the FMA vary (above the required accuracy) between different time periods, it is implied that the corresponding KAM tori are destroyed. Thus, the study of this frequency map \(\mathcal{F}\) (i.e. the FMA) regarding its constancy in time provides important clues about the dynamics of the system.

For a comprehensive discussion of the FMA, refer to Laskar (2003).

In order to calculate the frequencies with the FMFT, we need to compute the equations of motion in the ScTS model. We use the Taylor method (Jorba \& Zou 2005) that proved to be very convenient for the task in D12. The precision in the computation of the coordinates is \(10^{-15}\). The numerical computation of the VICs was done using the DOPRI8 routine (Prince \& Dormand 1981). This routine achieved better results than Taylor when we required the simultaneous integration of the coupled system of equations of motion and variational equations in the ScTS model (see also D12). The conservation of the energy with the DOPRI8 routine was \(\sim 10^{-13}, 10^{-14}\).

All the computations in the forthcoming investigation were done using the following configuration. Hardware: CPU, 2×Dual XEON 5450, Dual Core 3.00 GHz; M.B., Intel S5000VSA; RAM, 4GB(4×1GB), Kingston DDR-2, 667MHz, Dual Channel. Software: GFORTRAN 4.2.3.

4 THE FMFT AS A GLOBAL INDICATOR OF CHAOS

In this section, we compare the performances of the FMFT and the SE/LCE as global indicators of chaos.

4.1 Accuracy in the estimation of the frequencies

In order to analyse the nature of each orbit, we apply the FMFT to the following function in each degree of freedom (hereinafter d.o.f.):

\[
\psi_j(t_i) = \psi_j^x(t_i) + i\psi_j^y(t_i)
\]

with \(\psi_j\) \((j = x, y, z)\) a complex function, where \(\psi_j^x\) (the real part of \(\psi_j\)) corresponds to the Cartesian coordinates \((x, y, z)\) and \(\psi_j^c\) (the conjugate part of \(\psi_j\)) corresponds to the Cartesian velocities \((v_x, v_y, v_z)\). The \(t_i\) with \(k = 1, \ldots, n\) are the trajectory samplings. Every set of points \(\psi_j\) generates a quasi-periodic approximation and we keep the frequency \(\nu_j(\neq 0)\) with the maximum amplitude. We call this frequency the fundamental frequency of the system in the j/d.o.f.

Theoretically speaking, if the fundamental frequencies correspond to a regular orbit, they should remain constant in time. Numerically speaking, the variation exists and it is due to the precision of the computation. Thus, in order to determine such a precision for our particular experiment, we take samples of regular orbits, according to both convergent values of the LI and the MEGNO (< ln(t)/T or < 2.01, respectively, see Maffione et al. 2011a).

We are aware that samples obtained from the use of VICs can only be statistically rich in regular orbits. That is, the actual values of the VICs are reached only at infinite time and as we have just truncated approximations of those values, some of the orbits might be misclassified. In order to be as certain as possible about the dominant type of orbits from the samples, we follow their behaviour for \(10^4 T(E)\) (see Section 3.1). In a galaxy like the one represented by the ScTS model, the crossing time is 0.5 u.t. If we suppose that the Hubble time is of the order of \(10^3\) crossing times (a high value

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\(^3\) There is a downloadable version of the FMFT in the personal web page of David Nesvorný: http://www.boulder.swri.edu/~davidn/
There are 100 i.c. in each sample and we integrate the energy surfaces to Section 3.4 and Muzzio (2006). The concomitant results for the times for the energy surfaces \( -0.1 \) and \( -0.5 \) on two 50 per cent overlapping time intervals of 3 \( \times \) \( 10^2 \) u.t.\(^4\) results in the determination of the fundamental frequencies with the FMFT.

In Table 3 we show the corresponding arithmetic means and standard deviations of the variations of the fundamental frequencies for the energy surfaces: \(-0.1\), \(-0.3\), \(-0.5\) and \(-0.7\) on the \( p_{x_0} - p_{z_0} \) start space. Thus, there exist a range of values where the regular orbits could not be reliably distinguished from the chaotic orbits by means of the computation of the fundamental frequencies with the FMFT.

\(^4\)The time intervals are \([0: 3 \times 10^2 T_c(E)]\) and \([1.5 \times 10^2 T_c(E): 4.5 \times 10^2 T_c(E)]\).

Figure 2. Variations of the fundamental frequencies (in logarithmic scale) computed with the FMFT for four samples of 100 regular orbits on each d.o.f.: on the \( p_{x_0} - p_{z_0} \) start space and on the energy surfaces \(-0.1\) (top-left panel) and \(-0.7\) (top-right panel), and on the \( x_0 - z_0 \) start space and on the energy surfaces \(-0.1\) (bottom-left panel) and \(-0.7\) (bottom-right panel). The missing points are due to the computational precision.

In Fig. 2 we show that the differences in the computed values of the fundamental frequencies between both overlapping time intervals for regular orbits (actually, some of them might be chaotic) are not generally below the theoretical value, particularly in the \( p_{x_0} - p_{z_0} \) start space. Therefore, we apply the FMFT on two 50 per cent overlapping time intervals of 3 \( \times \) \( 10^2 \) \( T_c(E) \) for every i.c. Although \( 10^3 T_c(E) \) is the time-span we use to compute the VICs reliably, the results in the determination of the fundamental frequencies with the FMFT are not very different using only 3 \( \times \) \( 10^2 \) \( T_c(E) \). Besides, the CPU time is certainly reduced. Therefore, we apply the FMFT on two 50 per cent overlapping time intervals of 3 \( \times \) \( 10^2 \) \( T_c(E) \) (Wachlin & Ferraz-Mello 1998).\(^4\) The number of points used in the estimation of the frequencies is 8192 which makes \( \sim 27 \) points for each \( T_c(E) \). Finally, we estimate the fundamental frequencies \( \nu_{j}^{(i)} \) for each time interval and calculate the differences [the accuracy in the determination of the frequencies should be \( \sim 10^{-9} \) according to Section 3.4 and Muzzio (2006)]. The concomitant results for the energy surfaces \(-0.1\) and \(-0.7\) for both the \( p_{x_0} - p_{z_0} \) and the \( x_0 - z_0 \) start spaces are shown in Fig. 2.

### Table 3. Arithmetic means and standard deviations of the differences of the fundamental frequencies between both overlapping time intervals for the samples of regular orbits used in Fig. 2. The values are ordered by energy surface and d.o.f.

| Energy   | \( x \)     | \( y \)     | \( z \)     |
|----------|-------------|-------------|-------------|
| \(-0.1\) | \(~1.2^{-9}\) | \(~2.3^{-7}\) | \(~3.4^{-7}\) |
| Standard deviations | \(~1.2^{-7}\) | \(~2.5^{-5}\) | \(~1.7^{-5}\) |
| \(-0.3\) | \(~1.8^{-8}\) | \(~1.2^{-6}\) | \(~5.9^{-6}\) |
| Standard deviations | \(~4.6^{-6}\) | \(~4.3^{-4}\) | \(~5.9^{-4}\) |
| \(-0.5\) | \(~5.3^{-9}\) | \(~1.9^{-8}\) | \(~1.1^{-7}\) |
| Standard deviations | \(~5.3^{-7}\) | \(~2.7^{-6}\) | \(~1.3^{-5}\) |
| \(-0.7\) | \(~3.5^{-7}\) | \(~6.9^{-8}\) | \(~1.9^{-5}\) |
| Standard deviations | \(~1.6^{-5}\) | \(~1.3^{-5}\) | \(~1.2^{-3}\) |
start space. The arithmetic means show an average conservation between the seventh and the eighth decimal in the fundamental frequencies. Nevertheless, the standard deviations oscillate around an average of the fifth decimal. Therefore, there is an interval where the FMFT cannot distinguish clearly between regular and chaotic orbits in the experiment. However, we can infer certain limits which can help us to identify if the orbit is regular or chaotic using the FMFT. For instance, the variations in the computed fundamental frequencies for the regular orbits only affect decimals higher than the fourth (see Fig. 2). Thus, if there is a difference in the computation of the fundamental frequencies of the orbit in decimals lower than the fourth, it is highly probable that the orbit is chaotic. This result is in complete agreement with the one shown in Muzzio (2006) for the same ScTS model and with the practical limit used also in Aquilano et al. (2007) and Muzzio, Navone & Zorzi (2009).

The results for the \( x_0 - \Delta_1 \) start space are similar to those for the \( p_{\Delta_0} - p_{\Delta_0} \) start space. Nevertheless, for lower energy values (Fig. 2, bottom-right panel), the averaged precision in the computation of the frequencies improved substantially.

We also make some experiments with the FMFT, using synthetic signals for verifying the precision in the computation of the fundamental frequencies. The results were remarkably better and are closer to the theoretical estimates (Muzzio 2006). Nevertheless, this is not the case for the relatively complex ScTS model and the precision in the estimations is clearly diminished. Thus, we use the same model and the same spaces we study to calibrate the FMFT.

4.2 A comparative evaluation between the FMFT and the SE/LCE

We are interested in testing the performance of the FMFT as a global indicator of chaos. Thus, we apply it together with a variational indicator in order to characterize the phase space portraits of two different regions. One of the regions is located in the \( p_{\Delta_0} - p_{\Delta_0} \) start space (the region is identified in the top panel of Fig. 3), whereas the other is located in the \( x_0 - \Delta_1 \) start space (which is identified in the bottom panel of the same figure). Both regions belong to the energy surface \( -0.7 \). The VIC selected for this study is the SE/LCE (see Section 3.1).

In order to apply the FMFT as a global indicator of chaos, we follow the same procedure as described in Section 4.1. Thus, we consider two overlapping time intervals of \( 10^3 \) periods. The first time interval is \([0: 7 \times 10^3]\) u.t. and the second time interval is \([3.5 \times 10^3: 10.5 \times 10^3]\) u.t. According to Section 3.4, the accuracy in the determination of the frequencies should be \( \sim 10^{-11} \). The computation is over 624 100 i.c., within the \( p_{\Delta_0} - p_{\Delta_0} \) start space, covering the region depicted in the top panel of Fig. 3 and over 596 258 i.c., within the \( x_0 - \Delta_1 \) start space, covering the region depicted in the bottom panel of the same figure.

To use the FMFT as a global indicator of chaos and compute the corresponding phase space portraits, we need to define a new quantity (Wachlin & Ferraz-Mello 1998). Such a quantity is \( \log(\Delta F) \), \( \Delta F \) being \( |v^{(i)}(t) - v^{(i)}(t)| + |v^{(i)}(t) - v^{(j)}(t)| + |v^{(i)}(t) - v^{(j)}(t)| \), where \( v^{(i)}(t) \) is the frequency (computed with the FMFT) associated with the d.o.f \( j \) for the time interval \( t(i) \). All the fundamental frequencies have to be computed in both intervals. Finally, the phase space portraits shown by the quantity \( \log(\Delta F) \) consist of 622 521 i.c. for the \( p_{\Delta_0} - p_{\Delta_0} \) start space and 594 690 i.c. for the \( x_0 - \Delta_1 \) start space.

In order to make appropriate comparisons on each start space, we need to evaluate the \( \log(\Delta F) \) and the SE/LCE on equal standings. Thus, it is necessary to decide a cut-off level for the distributions of their values to use the same scales for both indicators.

In Fig. 4 we show the histograms according to the final values of the techniques. The histograms in the top panel correspond to the \( p_{\Delta_0} - p_{\Delta_0} \) start space and the histograms in the bottom panel correspond to the \( x_0 - \Delta_1 \) start space. A clear bimodal distribution (one peak corresponding to regular orbits, and the other to chaotic orbits) is shown for both indicators. In the case of the \( p_{\Delta_0} - p_{\Delta_0} \) start space, the indicators have less than 1 per cent values for the orbits below \( 10^{-8} \) (pointed out in the top panel of the same figure). Thus, the scales on the \( p_{\Delta_0} - p_{\Delta_0} \) start space go from \( 10^{-8} \) to 0. The similarities between both histograms on the \( p_{\Delta_0} - p_{\Delta_0} \) start space make easier the selection of a common cut-off level. However, there are important differences between the histograms on the \( x_0 - \Delta_1 \) start space. The SE/LCE shows a bimodal distribution within the interval \([\sim 10^{-10}: 0]\) with its peaks well separated. The peaks in the distribution on the \( x_0 - \Delta_1 \) start space match the ones found for the same indicator, but on the \( p_{\Delta_0} - p_{\Delta_0} \) start space (see both the panels in Fig. 4). This is not the case for the \( \log(\Delta F) \). The
peak that corresponds to the regular component is beyond the value $10^{-10}$ (see Fig. 4, bottom panel and Fig. 2, bottom-right panel).

Nevertheless, we find that the number of orbits with values of the SE/LCE below the value $10^{-10}$ are negligible and the log ($\Delta F$) does not show further structure below that value. Thus, we decided to use that value as the cut-off level in the distributions on the start space to build the scales.

Note also that according to log ($\Delta F$) on the $p_{x_0} - p_{z_0}$ start space, the peak corresponding to regular orbits is $\sim 3.1 \times 10^{-6}$. This value confirms the conclusions drawn in Section 4.1 with a sample composed only of 100 orbits for the same start space. That is, if we use the arithmetic means for the energy surface $-0.7$ (Table 3), we arrive at a similar value of $\sim 1.9 \times 10^{-7}$. Furthermore, the minimum between both peaks (which gives a clear distinction between the regular and the chaotic components) in the $p_{x_0} - p_{z_0}$ start space is taken around the value $10^{-4}$. This value also matches the one considered to separate regular from chaotic orbits for the same start space in Section 4.1.

In Fig. 5 we present the phase space portraits given by the SE/LCE (left-hand panels) and the log ($\Delta F$) (right-hand panels) for both selected regions in the $p_{x_0} - p_{z_0}$ (top panels) and the $x_0 - z_0$ (bottom panels) start spaces.

The phase space portraits of the region inside the $p_{x_0} - p_{z_0}$ start space given by the SE/LCE and the log ($\Delta F$) present the same structures. In spite of this high level of coincidence in the description of the $p_{x_0} - p_{z_0}$ start space, there are differences in the $x_0 - z_0$ start space. In the bottom-right panel of Fig. 5, the quantity log ($\Delta F$) shows an amount of spurious structures\(^5\) not shown by the variational indicator (bottom-left panel of the same figure). In fact, the spurious structures make difficult to select a threshold to distinguish between regular and chaotic orbits. Still, it is possible to define a cut-off value for the log ($\Delta F$), but the procedure is more hand-made. For example, if we take as regular orbits those preserving $\geq 4$ decimals in their fundamental frequencies (see Section 4.1 and the previous discussion), we recover the portrait given by the SE/LCE for the $x_0 - z_0$ start space. However, the SE/LCE (as well as most of the VICs studied in previous papers, for instance M11 and D12) shows a large separation of the different kinds of motion in both portraits (see also Fig. 4), and the choice of a threshold (if it is not already defined) is easier.

These results, which are not as clear as those given by the variational indicators in terms of motion separation, make the FMFT a second choice to study the global dynamics in a divided phase space.

The procedure used to determine the chaoticity or regularity of the orbits with the FMFT is standard. The deficiency in the descriptions is basically due to the method’s sensitivity to its parameters. Therefore, the reliability of the FMFT as a global indicator of chaos is, comparatively speaking, limited.

### 4.2.1 The computing times

The VICs need to compute the equations of motion together with the variational equations to determine the nature of the orbits. This process may take a large amount of time. On the other hand, the FMFT works with the computation of the frequencies, which is a fast process, but the determination of the log ($\Delta F$) to characterize the nature of the orbits involves further calculations. For example, the computation of the SE/LCE (a fast VIC) to obtain the top and bottom-left panels of Fig. 5 took $\sim 670$ and $\sim 330$ h, respectively, by using the DOPRI8 routine. Instead, the computation of the fundamental frequencies (nothing more than the line of biggest amplitude in each d.o.f. was computed) with the FMFT for 624 100 and 594 690 i.c. took $\sim 30$ h for both experiments, after the integration of the equations of motion. The time spent in this integration for $7 \times 10^3$ u.t. was of $\sim 570$ and $\sim 280$ h for the regions in the $p_{x_0} - p_{z_0}$ and the $x_0 - z_0$ start spaces, respectively. In both cases, we used the Taylor method (see Section 3.4). Furthermore, if we want to use the FMFT as a global indicator of chaos, we need to compute the log ($\Delta F$), i.e. we need to compute the frequency vector for two time intervals. If we overlap the time intervals in 50 per cent, we economize on computing time. The total time interval must be $1.05 \times 10^4$ u.t. under these circumstances. That is, 50 per cent larger than the one used by the variational indicators. Finally, the time required by the FMFT extends from $\sim 600$ ($\sim 570 + \sim 30$) to $\sim 885$ h and from $\sim 310$ ($\sim 280 + \sim 30$) to $\sim 450$ h to obtain the portraits in the top and bottom-right panels of Fig. 5, respectively.

The advantage of the fast computation of the frequencies is certainly lost in the time-consuming process involved in the computation of the log ($\Delta F$).

In conclusion, the FMFT as a global indicator of chaos seems to be rather inconvenient when there are other alternatives such as fast VICs like the SE/LCE.

\(^5\) Some of them are due to the phenomenon of Moiré, which is inherent in discrete Fourier transform techniques (Barrio et al. 2009b).
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Figure 5. Phase space portraits corresponding to both regions within the $p_{x_0} - p_{z_0}$ (top panels) and the $x_0 - z_0$ start (bottom panels) spaces on the energy surface $-0.7$. The samples used to compute the SE/LCE consist of 624 100 and 596 258 (top and bottom-left panels, respectively) i.c. and $7 \times 10^3$ u.t. The samples used to compute the log ($\Delta F$) consist of 622 521 and 594 690 (top and bottom-right panels, respectively) i.c. and two overlapping time intervals of $7 \times 10^3$ u.t. each. Chaotic regions are identified by the values of the indicators that are higher than $\sim -4, \sim -3$ (note that the values of the indicators are in logarithmic scale). See the text for further details.

5 THE SELECTION OF PROPER VICs

The aim of this section is to select the proper VICs to be used in the forthcoming experiments on the $p_{x_0} - p_{z_0}$ and the $x_0 - z_0$ start spaces of the ScTS model (Section 6).

We first study a small region in the $p_{x_0} - p_{z_0}$ start space (within the intervals $p_{x_0} \in [0.87 : 1.02]$ and $p_{z_0} \in [1.39 : 1.54]$) on the energy surface $-0.7$, shown in the red square in Fig. 6). We use the MEGNO, the SE/LCE, the FLI, the OFLI and the APLE. This first experiment provides us with a comparison between quite new techniques (the SE/LCE and the APLE) and VICs with excellent performances shown in previous studies (the MEGNO and the FLI/OFLI, e.g. in M11 and D12).

In addition, we use the same $10^3$ characteristic times throughout the experiments.

We are interested in the resolving power and the speed of convergence of the VICs. Therefore, we compute 10 201 i.c. within the region marked in Fig. 6 for $7 \times 10^3$ u.t.

5.1 The resolving power of the VICs

First, we briefly discuss the resolving power of the VICs under the circumstances of the experiment.

As we are looking for VICs to study big samples of i.c., we consider the final values of the indicators rather than their time evolution curves. Furthermore, when the indicator grows or decreases exponentially, it is convenient to stop the integration process at a particular saturation value. Thus, instead of simply registering the final values of the indicator, we can record the time needed to reach such a saturation value, i.e. the saturation times (Skokos et al. 2007).

The final values together with the saturation times of the VIC should be considered for the description of the phase space portrait (M11 and D12).
Figure 7. We show the phase space portraits of the different VICs using a total integration time of $7 \times 10^3$ u.t. for the region displayed on Fig. 6. We present the final values of the MEGNO (top-left panel); the final values of the SE/LCE (top-right panel); the final values (middle-left panel, in logarithmic scale) and the saturation times (middle-right panel) of the OFLI; and the final values (bottom-left panel) and the saturation times (bottom-right panel) of the APLE. The cold colours correspond to regular regions while the warm colours correspond to chaotic domains for the final values. The colour scale is inverted for the saturation times.

In Fig. 7 we present the phase space portraits given by the final values of the MEGNO and the SE/LCE (top-left and top-right panels, respectively); the OFLI final values (middle-left panel) and the saturation times (middle-right panel); and the APLE final values (bottom-left panel) and the saturation times (bottom-right panel). The phase space portraits given by the FLI are very similar to those shown with the OFLI and they are not included for the sake of simplicity.

The region under analysis has a regular component (the main central structure) and a chaotic component (surrounding the main central structure). Moreover, there are two different regions of chaos. The region that surrounds the regular component is composed of very strong chaotic orbits while the band on the right-hand side of every panel in Fig. 7 is filled with moderate-to-strong chaotic orbits.

The values of the MEGNO and the SE/LCE do not grow exponentially for chaotic orbits as it happens, for instance, with the OFLI (and the FLI) or the APLE. Then, a saturation value is not defined and a saturation time is not recorded in the computing process. However, the values of the MEGNO and the SE/LCE can be very large for strong chaotic orbits and very different from their values for regular and mild chaotic orbits. In a divided phase space with regions of very strong chaos, the large values of these indicators for strong chaotic orbits may hide the small differences between regular and mild chaotic orbits. Indeed, the differences among strong chaotic orbits are generally not as important as the differences between strong and mild chaotic orbits or between mild chaotic orbits and regular orbits. Thus, a saturation value for the MEGNO or the SE/LCE is also very useful in this kind of scenario. As shown in
In the case of the OFLI (and the FLI) or the APLE, the chaotic orbits grow exponentially fast and a saturation value is needed to avoid the propagation of errors in their computation. The saturation value in Fig. 7 for the OFLI (middle panels) is \( -\ln(10^2) \) (see Section 3.2). The saturation value in Fig. 7 for the APLE (bottom panels) is \( -\ln(10^2)/\ln(t) \) (see Section 3.3).

In the top panels of Fig. 7, we can see the regular and the chaotic components well separated by the MEGNO (left-hand panel) and the SE/LCE (right-hand panel). Furthermore, the two different chaotic regions are clearly shown: the region of strong chaos (surrounding the central structure with regular orbits) and the band on the right-hand side of the panels filled with moderate-to-strong chaotic orbits. There is no structure at all in the main central structure composed of regular orbits (top-left panel) because of the MEGNO’s asymptotic threshold (Section 3.1). Also, there is no further structure in the strong chaotic region due to the selected saturation value. The SE/LCE shows a similar portrait (top-right panel).

In the middle panels of Fig. 7, we use the OFLI to describe the region under analysis. On one hand, we recover the phase space portraits (middle-left panel) given by the MEGNO and the SE/LCE in the top panels of the same figure. On the other hand, the band on the right-hand side of both panels shows a more detailed structure with the OFLI. This is due to the fact that the saturation value taken for the OFLI covers a wider range of chaotic orbits than the values chosen for the MEGNO or the SE/LCE. The saturation times (middle-right panel) enhance the information given with the final values of the OFLI, specially within the strong chaotic region.

In the bottom panels of Fig. 7, we present the phase space portraits given by the APLE final values (left-hand panel) and saturation times (right-hand panel). The description is very similar to those given by the other VICs if we consider both the final values and the saturation times of this indicator. For example, the identification of the band on the right-hand side of the panels needs the information of the saturation times. The saturation times show a clear distinction between the region of strong chaos surrounding the main central structure and the region with mild and strong chaotic orbits in the band on the right-hand side of the panels. This is not the case for the other VICs (the MEGNO, the SE/LCE, the OFLI and the FLI) for which the saturation times play a complementary role and the old values are enough for such identification. Nevertheless, the degree of coincidence in the descriptions of the four VICs (plus the FLI) seems to be high. Thus, there is no decisive advantage in favour of one of them.

### 5.2 The speed of convergence of the VICs: thresholds

Now we focus on another important characteristic of the VICs which is fundamental for an efficient study of large samples of i.e.: the speed of convergence.

We use the same total integration time (7 × 10^3 u.t.) and the same i.e. (10 201) used in Section 5.1. The initial 10^2 u.t. of the time interval does not give reliable information for our purposes and it is not considered in this experiment. The remaining 6.9 × 10^3 u.t. are divided into 70 sub-intervals of ~98.57 u.t. each. We compute the number of chaotic orbits in each sub-interval. Before that, we had to identify the chaotic orbits. So, we needed accurate thresholds for each VIC. Many of them have theoretical or empirical estimations of their thresholds. Yet, they are only approximations and should be adjusted to each particular situation. Hence, we generalize the experiments shown in M11 and D12 and consider a wide range of thresholds for the different VICs. Finally, for each sub-interval, we compute the number of chaotic orbits for every VIC according to the different thresholds.

Performing several tests (i.e. considering different thresholds and VICs), we find that a percentage of ~83 per cent for the chaotic component is stable and prevails in the experiment. We take it as the ‘true’ percentage for the chaotic component. Having such a ‘true’ percentage, we can adjust the thresholds of the VICs under consideration. The fastest convergence of the VIC to a stable percentage of chaotic orbits close to the ‘true’ percentage gives the most appropriate threshold for that particular indicator.

In order to identify such appropriate thresholds for every VIC under consideration, we define the ‘selected thresholds’. These thresholds allow the associated indicator to reach the ‘true’ percentage of chaotic orbits within the total integration time.\(^6\)

In Fig. 8 we present the variation of the selected thresholds with the integration time for the APLE (top-left panel), the MEGNO (top-right panel), the FLI (bottom-left panel) and the OFLI (bottom-right panel). The SE/LCE is not included in the experiment because it does not have a defined mechanism to compute its threshold. The filled circles in Fig. 8 indicate the selected thresholds that allow the convergence of the corresponding VIC to a stable, close to the ‘true’ percentage of chaotic orbits. The filled squares in the same figure indicate the selected thresholds that offer the fastest convergence. In other words, the filled squares indicate the most appropriate thresholds for a given indicator.

The APLE has the theoretical time-independent threshold \( \sim 1 \) [see Section 3.3 and Lukes-Gerakopoulos et al. (2008) for further details]. However, we consider threshold values between \(~1.2\) and \(~1.5\). The threshold values below \(~1.2\) characterize 100 per cent of the orbits as chaotic orbits for the whole time interval, and thus, they are discarded. The threshold values above \(~1.5\) do not arrive at the ‘true’ percentage of the chaotic component within the time interval, and thus, they are also discarded. If we use the threshold values in the interval \( \sim[1.4: 1.5]\), the APLE converges to stable percentages of the chaotic component close to the ‘true’ percentage (see the top-left panel of Fig. 8). If we use the threshold values around 1.4, the APLE’s convergence is the fastest (approximately by \(5.5 \times 10^3\) u.t.).

The MEGNO, like the APLE, has also a time-independent threshold: \( \sim 2 \) [see Section 3.1 and Cincotta et al. (2003) for further details]. For the same reasons given for the APLE, we consider threshold values above the theoretical approximation, i.e. between \( \sim 2.8\) and \( \sim 4\). The threshold values below \( \sim 2.8\) characterize 100 per cent of the orbits as chaotic orbits for the whole time interval, and thus, they are discarded. The threshold values above \( \sim 4\) are much larger than the theoretical asymptotic estimation of \( \sim 2\), and thus, they are also discarded. If we use the threshold values close to \( \sim 4\), the MEGNO reaches percentages of the chaotic component close to the ‘true’ percentage more rapidly (see the top-right panel of Fig. 8). Nevertheless, using this range of selected thresholds the percentages are not convergent. Moreover, the percentages oscillate within the interval \( \sim [84\text{ per cent} : 87\text{ per cent}]\) depending on the threshold.

\(^6\) We can estimate the saturation value \( \langle \tilde{t} \rangle \) for the SE/LCE from the expression \( 20 \sim \tilde{t}/2 \times t \), with \( t = 7 \times 10^3 \) u.t. (see Section 3.1).

\(^7\) Actually, the selected thresholds allow the associated VIC to reach a percentage of chaotic orbits between 82 and 84 per cent within the total integration time.
Figure 8. We present the selected thresholds for the APLE (top-left panel), the MEGNO (top-right panel), the FLI (bottom-left panel) and the OFLI (bottom-right panel). The sample consists of 10 201 i.c. for $7 \times 10^3$ u.t. in the region shown in Fig. 6. See the text for further details.

selected. In other words, the MEGNO gives a higher percentage of chaotic orbits than the prevailing percentage shown by the other VICs. This experiment confirms the results yielded in M11 and D12.

The FLI and the OFLI have both similar behaviour and time-dependent thresholds to distinguish chaotic motion from regular motion [see Section 3.2 and Fouchard et al. (2002) for further details]. After computing the logarithm in their definitions, a tentative threshold for them is $\ln (t)$ with $t$ the integration time. We consider the general expression $a \times \ln (t)$ in order to study a variation of the thresholds changing the values of the parameter $a$.

The values of the parameter $a$ for the FLI belong to the interval $\sim [1.2: 1.6]$. The theoretical threshold with $a \sim 1$ is not efficient because it does not arrive at stable percentages of the chaotic component close to the ‘true’ percentage for the whole time interval. The same applies to the thresholds with $a \lesssim 1.2$ so they are discarded. Furthermore, the threshold values with $a \gtrsim 1.6$ do not reach at the ‘true’ percentage of the chaotic component within the time interval, and thus, they are also discarded. If we use the threshold values in the interval $\sim [1.2: 1.6]$ in the time interval $\sim [5.7 \times 10^3: 7 \times 10^3]$ u.t., the FLI converges to stable percentages of the chaotic component close to the ‘true’ percentage (see the bottom-left panel of Fig. 8). If we use the thresholds with $a$ in the interval $\sim [1: 1.4]$ and within the time interval $\sim [5.7 \times 10^3: 7 \times 10^3]$ u.t., the OFLI converges to stable percentages of the chaotic component close to the ‘true’ percentage (see the bottom-right panel of Fig. 8). If we use the threshold values with $a \sim 1.07$, the OFLI’s convergence is the fastest [approximately by $5.7 \times 10^3$ u.t., i.e. $1.07 \times \ln (5.7 \times 10^3) \sim 9.2$ in the figure]. These results are very similar to those obtained with the FLI as mentioned before. Yet, once again there is an important difference: the threshold selected for the OFLI ($a \sim 1.07$) is closer to the theoretical approximation ($a \sim 1$) than the threshold selected for the FLI ($a \sim 1.3$). Therefore, this indicates that the OFLI may have a better theoretical approximation of its threshold than the APLE.

Finally, we also observe similar results for the OFLI. However, the parameter $a$ varies from $\sim 0.5$ to $\sim 1.5$, i.e. for the first time the interval of selected thresholds includes the theoretical estimation ($a \sim 1$). That is, if we use the theoretical approximation, the OFLI reaches a stable percentage of chaotic orbits close to the ‘true’ percentage of 83 per cent. Nevertheless, the convergence arrives at the end of the time interval, i.e. $\sim 6.9 \times 10^3$ u.t. The threshold values below $\sim 0.5$ characterize 100 per cent of the orbits as chaotic orbits for the whole time interval, and thus, they are discarded. The threshold values above $\sim 1.5$ do not arrive at the ‘true’ percentage of the chaotic component within the time interval, and thus, they are also discarded. If we use the thresholds with $a$ in the interval $\sim [1: 1.4]$ and within the time interval $\sim [5.7 \times 10^3: 7 \times 10^3]$ u.t., the OFLI converges to stable percentages of the chaotic component close to the ‘true’ percentage (see the bottom-right panel of Fig. 8). If we use the threshold values with $a \sim 1.07$, the OFLI’s convergence is the fastest [approximately by $5.7 \times 10^3$ u.t., i.e. $1.07 \times \ln (5.7 \times 10^3) \sim 9.2$ in the figure]. These results are very similar to those obtained with the FLI as mentioned before. Yet, once again there is an important difference: the threshold selected for the OFLI ($a \sim 1.07$) is closer to the theoretical approximation ($a \sim 1$) than the threshold selected for the FLI ($a \sim 1.3$). Therefore, this indicates that the theoretical approximation of the thresholds of the OFLI may work better.

6 GLOBAL ANALYSIS OF THE START SPACES

In this section we proceed to study some dynamical aspects of the ScTS model by means of the complementary use of the VICs selected in Section 5.
We consider samples of around one million i.c. on the $p_{z_0} - p_{z_0}$ and the $x_0 - z_0$ start spaces for four different energy surfaces. Even though we do not study the remaining five start spaces indicated in Schwarzschild (1993) and Papaphilippou & Laskar (1998), we gain sufficient information in order to briefly discuss some main features of the dynamics of the system.

The total integration times extend to $1.17 \times 10^5$ u.t. for the energy surface $-0.1$ (Section 4.1). The CPU times become critical and will be considered a priority in the selection of the VICs. The FLI family of indicators presents the most versatile indicators, showing good performances in all the experiments (see also M11 and D12 in which they are included in the corresponding CIvF). The MEGNO is also another candidate because of its good performance (also included in the CIvF of M11 and D12). The SE shows a lower computational cost in the experiment than that of the FLI family of indicators. Therefore, we select the LI (the least time-consuming indicator given a fixed total integration time; see e.g. D12) and the pair MEGNO/SE/LCE as the VICs to study the ScTS model.

### 6.1 The $p_{z_0} - p_{z_0}$ start space

We have already mentioned that the SE/LCE has certain advantages when we study big samples of orbits (see Section 3.1). Nevertheless, it does not have any defined procedure to determine a threshold in order to distinguish between regular and chaotic motion. Indeed, this fact is a serious problem if we want to separate regular motion from weak chaotic motion. We need a threshold for the SE/LCE to correctly describe the regular component (red, blue and black colours) increases for lower energy values. However, this increment is not considerable yet.

In Fig. 9 we observe that independently of the energy surface, the chaotic component is the dominant one on the phase space portraits (yellow and orange colours; see also columns 3 and 5 in Table 4). The regular component (red, blue and black colours) increases for values of $p_{z_0}$ from $-0.3$ to $-0.7$.

In the top-left panel of Fig. 9, we observe a clear separation of both components. We show a fully connected chaotic component for values of $p_{z_0} \lesssim 1.7$ and the regular component for values of $p_{z_0} \gtrsim 1.7$, besides some hyperbolic structures. These structures grow as we move to lower energy surfaces. They are resonances that start to overlap with each other and occupy the regular component. We can also distinguish a separation inside the chaotic component. The fully connected chaotic domain moves back to lower values of $p_{z_0}$. The region is occupied by another chaotic domain characterized by a resonance overlapping regime and a lower Lyapunov number (top-right panel of Fig. 9). On the energy surface $-0.5$ (bottom-left panel of Fig. 9), the fully connected chaotic domain has noticeably moved backwards and the resonances

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8 In this test on computing times, we consider the definition of the FLI given in Froeschl´e & Lega (2000), where the authors compute the evolution of the FLI using one deviation vector.

9 The theoretical estimations: $9.9743 \times 10^{-5}$ (u.t.)$^{-1}$ for the energy surface $-0.1$; $4.2024 \times 10^{-4}$ (u.t.)$^{-1}$ for $-0.3$; $7.82722 \times 10^{-4}$ (u.t.)$^{-1}$ for $-0.5$ and $1.264809 \times 10^{-3}$ (u.t.)$^{-1}$ for $-0.7$, which are considerably smaller than the thresholds of the LI, because the estimation of the ILCE by the slope of the MEGNO converges more rapidly to the ILCE than the standard procedure (see Section 3.1; Cincotta et al. 2003).

10 We use a cut-off level around $10^{-9}$ to have the same scales in Fig. 9 for every energy surface. Then, the percentages of orbits that we do not consider are negligible. Those percentages are $\sim 0.77$ per cent in the energy surface $-0.1$, $\sim 0.06$ per cent in $-0.3$, $\sim 0.03$ per cent in $-0.5$ and $\sim 0.02$ per cent in $-0.7$.

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| Energy surface | Threshold (LI) | Chaotic orbits (LI) | Threshold (SE/LCE) | Chaotic orbits (SE/LCE) |
|---------------|----------------|---------------------|--------------------|------------------------|
| $-0.1$        | $1.2 \times 10^{-4}$ | $\sim 71.6$ per cent | $1.4 \times 10^{-5}$ | $\sim 66.8$ per cent |
| $-0.3$        | $5.8 \times 10^{-4}$ | $\sim 65.8$ per cent | $6.4 \times 10^{-5}$ | $\sim 65.9$ per cent |
| $-0.5$        | $1.25 \times 10^{-3}$ | $\sim 61.7$ per cent | $1.3 \times 10^{-4}$ | $\sim 63.0$ per cent |
| $-0.7$        | $1.7 \times 10^{-3}$ | $\sim 65.0$ per cent | $2.7 \times 10^{-4}$ | $\sim 62.9$ per cent |
occupy extended regions of the $p_{x_0} - p_{z_0}$ start space. Furthermore, a major resonance is seen around the value $p_{x_0} \sim 0.5$ and minor resonances are seen around the values $p_{x_0} \sim 1$ and $p_{x_0} \sim 1.3$. In the bottom-right panel of Fig. 9, for the energy surface $-0.7$, the portrait shows how the resonances occupy most of the regular component. Moreover, major resonances grow within the chaotic domains. The most evident resonance is that located at the value $p_{x_0} \sim 0.5$, which has already been identified in outer energy surfaces.

In spite of the short discussion of the $p_{x_0} - p_{z_0}$ start space given above, the efficiency of variational indicators to describe the global characteristics of the phase space is evident. The VICs easily identify the regular and the chaotic domains and their mutual interplay, along with phase space mixing processes like resonance overlapping. However, the identification of the underlying resonant structure by the frequency vectors is not readily understood. We should study the dimensionality of the tori (e.g. with the GALI method; Skokos et al. 2007; D12) to identify resonant orbits, locate the periodic orbits (e.g. with the OFLI; Fouchard et al. 2002; D12) and analyse their stability (e.g. with the MEGNO; Cincotta et al. 2008) to look for resonant orbit families. The process could be time consuming and rather inefficient since the VICs are not the most suitable tools for the task. In fact, the SAMs are far better options (see Section 1 and references therein).

The dynamics of the $p_{x_0} - p_{z_0}$ start space can be briefly described as a dominant chaotic component based mostly on a resonance overlapping regime. This phase mixing process produces a more homogeneous chaotic component for the energy surface $-0.1$.

For further studies concerning the $p_{x_0} - p_{z_0}$ start space of the ScTS model, refer to Muzzio et al. (2005) and Cincotta et al. (2008).

Now, we apply the same package of techniques to the $x_0 - z_0$ start space, where the dynamics is totally different since it is dominated by the regular orbits.

6.2 The $x_0 - z_0$ start space

Here, we implement the same procedure applied in Section 6.1 to study the $x_0 - z_0$ start space. We obtain the phase space portraits with the SE/LCE for four energy surfaces, calibrating the thresholds with the assistance of the LI, and we further discuss the global dynamics of the $x_0 - z_0$ start space.

In order to determine the percentages of the chaotic component, we proceed like we did in Section 6.1. We present the results ordered by energy surface on Table 5.

In Fig. 10, we present the phase space portraits of the $x_0 - z_0$ start space$^{11}$ for the same energy surfaces considered in Section 6.1. Also the total integration times remain the same and the VICs used for the study guarantee reliable and stable pictures. The numbers of orbits in the samples are different and depend on the energy surface: 998 938 i.c. for the energy surface $-0.1$, 998 948 for $-0.3$, 999 400 for $-0.5$ and 998 626 for $-0.7$.

There is a significant change in the dynamics of the $x_0 - z_0$ start space with respect to the results for the $p_{x_0} - p_{z_0}$ start space. The

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$^{11}$ We use a cut-off level of around $10^{-10}$ to have the same scales in Fig. 10 for every energy surface. Then, the percentages of orbits that we do not consider are negligible. Those percentages are $\sim 0.22$ per cent in the energy surface $-0.1$, $\sim 0.05$ per cent in $-0.3$, $\sim 0.02$ per cent in $-0.5$ and $\sim 0.02$ per cent in $-0.7$. 

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Figure 9. Phase space portraits of the SE/LCE for a sample of 1000444 i.c. on the $p_{x_0} - p_{z_0}$ start space and for four energy surfaces. Top-left panel: phase space portrait corresponding to the energy surface $-0.1$ and using a total integration time of $1.17 \times 10^4$ u.t. Top-right panel: $-0.3$ and $2.4 \times 10^4$ u.t. Bottom-left panel: $-0.5$ and $1.2 \times 10^4$ u.t. Bottom-right panel: $-0.7$ and $7 \times 10^3$ u.t. Chaotic regions are identified with values of the indicator higher than $\sim 5$ or $\sim 4$ for the energy surfaces $-0.1$ and $-0.3$ or $-0.5$ and $-0.7$, respectively. See Table 4 for further details. Note that the values of the indicator are in logarithmic scale.
Table 5. For every energy surface in the \( x_0 - z_0 \) start space, we present the threshold associated with the LI and the estimated percentage of chaotic orbits; the threshold associated with the SE/LCE and the estimated percentage of chaotic orbits.

| Energy surface | Threshold (LI) | Chaotic orbits (LI) | Threshold (SE/LCE) | Chaotic orbits (SE/LCE) |
|----------------|---------------|---------------------|--------------------|------------------------|
| \(-0.1\)       | \(1.25 \times 10^{-4}\) | \(\sim 3.76\) per cent | \(9.4 \times 10^{-6}\) | \(\sim 4.65\) per cent |
| \(-0.3\)       | \(5.3 \times 10^{-4}\)  | \(\sim 8.15\) per cent | \(3.2 \times 10^{-5}\) | \(\sim 9.2\) per cent |
| \(-0.5\)       | \(9.4 \times 10^{-4}\)  | \(\sim 15.06\) per cent | \(6.5 \times 10^{-5}\) | \(\sim 12.74\) per cent |
| \(-0.7\)       | \(1.7 \times 10^{-3}\)  | \(\sim 16.37\) per cent | \(9.5 \times 10^{-5}\) | \(\sim 16.75\) per cent |

Figure 10. Phase space portraits of the SE/LCE for the \( x_0 - z_0 \) start space and for four energy surfaces. Top-left panel: phase space portrait corresponding to the energy surface \(-0.1\) for 998,938 i.c. and using a total integration time of \(1.17 \times 10^5\) u.t. Top-right panel: \(-0.3\), with 998,948 i.c. and \(2.4 \times 10^4\) u.t. Bottom-left panel: \(-0.5\), with 999,400 i.c. and \(1.2 \times 10^4\) u.t. Bottom-right panel: \(-0.7\), with 998,626 i.c. and \(7 \times 10^3\) u.t. Chaotic regions are identified with indicator values higher than \(\sim -5\) or \(\sim -4\) for the energy surfaces \(-0.1\), \(-0.3\) and \(-0.5\) or \(-0.7\), respectively. See Table 5 for further details. Note that the values of the indicator are in logarithmic scale.

corresponding phase space portraits presented in Fig. 10 show a dominant regular component (see Table 5, columns 3 and 5) instead of the rather large chaotic component seen for the \( p_{x_0} - p_{z_0} \) start space. The percentages of the smaller component have a substantial increment towards the innermost energy surface. Indeed, in this case, the expansion of the chaotic component is due to the growth of the different orbital families’ separatrices. This fact is clearly seen in Fig. 10. In the top-left panel of Fig. 10, we can see a very weak separatrix that separates the short-axis tubes from the outer long-axis tubes. In the top-right panel of the same figure, we can see how the former separatrix is connected to the others which separate the inner long-axis tubes and the box orbits [see Papaphilippou & Laskar (1998) for further details on the different orbital families and their locations in phase space]. This becomes evident in the bottom panels of Fig. 10. Furthermore, a strong chaotic domain appears in the crossing of the main separatrices. Therefore, as the dominant component comprises regular orbits, the main contribution to the chaotic component emerges from the separatrices of resonances.

Finally, the dynamics of the \( x_0 - z_0 \) start space can be briefly described as a dominant regular component. The major resonances that separate the main orbital families have separatrices which grow when the negative energies increase. Furthermore, the growth of the separatrices increases the percentage of chaotic orbits very fast.

7 CONCLUSIONS

With this report we conclude a series of investigations (reported in Maffione et al. 2011a, M11 and D12) towards an efficient choice of a minimal package of techniques to study a general Hamiltonian.

Here, we apply both VICs and an SAM (the FMFT; Sidlichovský & Nesvorný 1997) to study a fairly realistic dynamical model of an elliptical galaxy (the ScTS model; Muzzio et al. 2005).

In order to select the appropriate VICs for the experiments, we not only consider previous comparisons, but also extend them with new indicators: the SE/LCE and the APLE (Lukes-Gerakopoulos et al. 2002).
et al. 2008). The SE/LCE is an efficient estimation of the ILCE by means of the MEGNO (Cincotta & Simó 2000) and works as its reliable alternative in the case of studying big samples of orbits. The APLE has not shown advantages over the FLI/OFLI for the purposes of the experiment, but it behaves similarly.

The SE/LCE seems to improve the performance of the MEGNO, in particular situations. Therefore, it should be considered in the CI sF presented in D12 as a suitable alternative.

On the other hand, we consider the performance of the FMFT as the representative SAM to compare with the VICs. The FMFT is an improvement of the FMA outlined by Laskar (1990), which is widely used by the scientific community.

The main advantage of the SAMs is a fast computation of the frequencies. However, the performance of the FMFT as a global indicator of chaos is not as efficient as the performances shown by the VICs in the experiments. The SAMs are designed to compute the frequencies, which are quantities strictly related to regular motion. Using such techniques as global indicators of chaos involves forcing the methods to do something for which they are not designed. The VICs are based on the concept of local exponential divergence. Hence, the detection of chaos is their main purpose. A natural consequence of this is that an efficient application of those techniques is based on their complementary implementation (see Section 1 for references). The SAMs are of use to describe the resonance web while the VICs to study the interplay of regular and chaotic domains (see Section 6).

Finally, the recommended CI sF for the analysis of a general Hamiltonian is composed of the MEGNO/SE/LCE, the FLI/OFLI and the GALI 2LCE: the MEGNO/SE/LCE and the FLI/OFLI as global indicators of chaos to obtain the phase space portraits and display the interaction between regular and chaotic orbits; the GALI 2LCE to analyse small samples within regions of complex dynamics and regions where the chaotic component is dominant (see Skokos et al. 2007 and D12). Furthermore, the CI sF can be used with a SAM (e.g. the FMFT) in order to characterize the resonance web with the frequency vectors.

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