A new code for orbit analysis and Schwarzschild modelling of triaxial stellar systems

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ABSTRACT

We review the methods used to study the orbital structure and chaotic properties of various galactic models and to construct self-consistent equilibrium solutions by Schwarzschild’s orbit superposition technique. These methods are implemented in a new publicly available software tool, SMILE, which is intended to be a convenient and interactive instrument for studying a variety of 2D and 3D models, including arbitrary potentials represented by a basis-set expansion, a spherical-harmonic expansion with coefficients being smooth functions of radius (splines) or a set of fixed point masses. We also propose two new variants of Schwarzschild modelling, in which the density of each orbit is represented by the coefficients of the basis-set or spline spherical-harmonic expansion, and the orbit weights are assigned in such a way as to reproduce the coefficients of the underlying density model. We explore the accuracy of these general-purpose potential expansions and show that they may be efficiently used to approximate a wide range of analytic density models and serve as smooth representations of discrete particle sets (e.g. snapshots from an $N$-body simulation), for instance, for the purpose of orbit analysis of the snapshot. For the variants of Schwarzschild modelling, we use two test cases – a triaxial Dehnen model containing a central black hole and a model re-created from an $N$-body snapshot obtained by a cold collapse. These tests demonstrate that all modelling approaches are capable of creating equilibrium models.

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

1 INTRODUCTION

The study of galactic structure relies heavily on construction and analysis of self-consistent stationary models, in which stars and other mass components (i.e. dark matter) move in the gravitational potential $\Phi$, related to their density distribution $\rho$ by the Poisson equation

$$\nabla^2 \Phi(r) = 4\pi \rho(r),$$

in such a way that the density profile remains unchanged. The evolution of distribution function $f(r, v)$ of stars moving in the smooth potential is described by the collisionless Boltzmann equation (CBE):

$$\frac{\partial f}{\partial t} + v \frac{\partial f}{\partial r} - \frac{\partial \Phi}{\partial r} \frac{\partial f}{\partial v} = 0.$$  (2)

To construct a stationary self-consistent model corresponding to a given density profile $\rho(r)$, one needs to find a function $f$ which is

the solution of (2) with a potential $\Phi$ related to $\rho$ via (1), such that $\rho = \int f(r, v) \, dv$. Various approaches to this problem may be broadly classified into methods based on the distribution function, Jeans equations, orbit superposition and iterative $N$-body schemes (see introduction sections in Dehnen (2009); Morganti & Gerhard (2012) for a nice summary). Of these, the first two are dealing with a smooth description of the system, but are usually restricted to sufficiently symmetric potentials (or special cases like a triaxial fully integrable model of van de Ven et al. 2003), while the latter two techniques represent the stellar system in terms of Monte Carlo sampling of the distribution function by orbits or $N$-body particles, thus achieving more flexibility at the expense of lack of smoothness.

In this paper, we discuss the Schwarzschild’s orbit superposition method, first introduced by Martin Schwarzschild (1979). It consists of two steps: first a large number of trajectories are computed numerically in the given potential $\Phi$ and their properties (most importantly, spatial shape) are stored in some way, then these orbits are assigned non-negative weights so that the density $\rho$, which corresponds to this potential via the Poisson equation (1), is given by a weighted sum of densities of individual orbits. In the classical

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Schwarzschild method, the density is represented in a discrete way by dividing the configuration space into a 3D grid and computing the mass in each grid cell, both for the underlying density profile to be modelled and for the fraction of time that each orbit spends in each cell. Then the contribution of each orbit to the model is obtained by solving a linear system of equations, subject to the condition that all orbit weights are non-negative. We propose two additional formulations of the Schwarzschild method, in which the density is represented not on a grid, but as coefficients of expansion over some basis.

This method was first applied to demonstrate that a triaxial self-consistent model of a galaxy can be constructed, and has been used to study various triaxial systems with central density cores (Statler 1987) and cusps (Merritt & Fridman 1996; Merritt 1997; Stiopis 1999; Terzić 2002; Capuzzo-Dolcetta et al. 2007; Thakur et al. 2007). The essential questions considered in these and other papers are whether a particular potential–density pair can be constructed by orbit superposition, what are the restrictions on the possible shapes of these models, how important are different types of orbits, including the role of chaotic orbits.

The Schwarzschild method is also very instrumental in constructing mass models of individual galaxies. In this application, the density model is obtained from observational data of surface brightness profile, which, however, does not have a unique deprojection in a non-spherical case. Kinematic constraints also come from observations. Typically one constructs a series of models with varying shape, mass of the central black hole (BH), etc. and evaluates the goodness of fit to the set of observables; the χ² statistics is then used to find the range of possible values for the model parameters allowed by the observations. Due to complicated geometry of the triaxial case, involving two viewing angles, most of the studies concentrated on the axisymmetric models. There exist several independent implementations of the axisymmetric modelling technique (e.g. Cretton et al. 1999; Gebhardt et al. 2000; Valluri, Merritt & Emsellem 2004) and at least one for triaxial models (van den Bosch et al. 2008). Observation-driven Schwarzschild method was used for constructing models of the Galactic bulge (Zhao 1996b; Häfner et al. 2000; Wang et al. 2012), constraining mass-to-light ratio and shape of galaxies (Thomas et al. 2004; van den Bosch et al. 2008), and for estimating the masses of central BHs (Gebhardt et al. 2003; Valluri et al. 2005; van den Bosch & de Zeeuw 2010). Related techniques are the made-to-measure method (Syer & Tremaine 1996b; de Lorenzi et al. 2007; Dehnen 2009; Long & Mao 2010) or the iterative method (Rodionov, Athanassoula & Sotnikova 2009), in which an N-body representation of a system is evolved in such a way as to drive its properties towards the required values, by adjusting masses or velocities of particles ‘on-the-fly’ in the course of simulation.

Of these two flavours of the Schwarzschild method, this paper deals with the first. We continue and extend previous theoretical studies of triaxial galactic models with arbitrary density profiles and, possibly, a central massive BH. For these potentials which support only one classical integral of motion – the energy¹ (in the time-independent case) – orbital structure is often quite complicated and has various classes of regular and many chaotic orbits, therefore it is necessary to have efficient methods of orbit classification and quantifying the chaotic properties of individual orbits and the entire potential model. These orbit analysis methods are often useful by themselves, besides construction of self-consistent models, for instance, for the purpose of analysing the mechanisms driving the evolution of shape in N-body simulations (Valluri et al. 2010) or the structure of merger remnants (Hoffman et al. 2010).

We have developed a new publicly available software tool, SMILE,² for orbit analysis and Schwarzschild modelling of triaxial stellar systems, intended to address a wide variety of ‘generic’ questions from a theorist’s perspective. This paper presents an overview of various methods and algorithms (including some newly developed) used in the representation of potential, orbit classification, detection of chaos and Schwarzschild modelling, that are implemented in SMILE.

In the Section 2, we introduce the basic definitions for the systems being modelled, and describe the main constituents and applications of the software. Section 3 presents the potentials that can be used in the modelling, including several flexible representations of an arbitrary density profile, Section 4 is devoted to methods for analysis of orbit class and its chaotic properties, and Section 5 describes the Schwarzschild orbit superposition method itself. In the remainder of the paper, we explore the accuracy of our general-purpose potential expansions (Section 6) and the efficiency of constructing a triaxial model using all variants of Schwarzschild method considered in the paper (Section 7).

2 THE SCOPE OF THE SOFTWARE

SMILE is designed in a modular way, allowing for different parts of the code to be used independently in other software (for instance, the potential represented as a basis-set expansion (BSE) could be used as an additional smooth component in an external N-body simulation program (e.g. Lowing et al. 2011), or the orbit analysis could be applied to a trajectory extracted from an N-body simulation). There are several principal constituents, which will be described in more detail in the following sections.

The first is the potential-density pair, which is used to numerically integrate the orbits and represent the mass distribution in Schwarzschild model. There are several standard analytical mass models and three more general representations of an arbitrary density profile, described in Section 3. The second fundamental object is an orbit with given initial conditions, evolved for a given time in this potential. The orbit properties are determined by several orbit analysis and chaos detection methods, presented in Section 4. The third constituent is the orbit library, which is a collection of orbits with the same energy, or covering all possible energies in the model. In the first case, these orbits may be plotted on the Poincaré surface of section (for 2D potentials), or on the frequency map (for the 3D case, discussed in Section 4.4), to get a visual insight into the orbital structure of the potential. In the second case, this collection of orbits is the source component of the Schwarzschild model. The target of the model may consist of several components: the density model corresponding to the potential in which the orbits were computed (there exist several possible representations of the density model, introduced in Section 5.2), kinematic information (e.g. the velocity anisotropy as a function of radius) or the observational constraints

¹ Axisymmetric Schwarzschild models are often called three-integral models, to distinguish them from simpler approaches involving only two classical integrals; however, this is not quite correct since not all orbits respect three integrals of motion even in the axisymmetric case.

² The acronym stands for ‘Schwarzschild Modelling Interactive exploratory Environment’. The software can be downloaded at http://hd.ipl.ru/~eugvas/smile/
in the form of surface density and velocity measurements with associated uncertainties. The latter case is not implemented in the present version of the software, but may be described within the same framework. The module for Schwarzschild modelling takes together the source and target components and finds the solution for orbit weights by solving an optimization problem (Section 5.3).

The tasks that can be performed using SMILE include:

(i) visualization and analysis of the properties of individual orbits;
(ii) exploration of the orbital structure of a given potential (one of well-studied analytical models, or an N-body snapshot with no a priori known structure);
(iii) construction of self-consistent Schwarzschild models for the given density profile, with adjustable properties (e.g. velocity anisotropy or the fraction of chaotic orbits) and their conversion to an N-body representation.

Many of these tasks can be performed interactively (hence the title of the program) in the graphical interface; there is also a scriptable console version more suitable for large-scale computations on multicore processors.

3 POTENTIAL MODELS

A number of standard non-rotating triaxial mass models are implemented.

(i) Logarithmic: \( \Phi(\tilde{r}) = \ln(\tilde{r}^2 + \tilde{p}^2) \), which was studied in Miralda-Escudé & Schwarzschild (1989); Schwarzschild (1993); Papaphillipou & Laskar (1998).

(ii) Triaxial generalization of Dehnen (1993) double power-law model:

\[
\rho(\tilde{r}) = \frac{3 - \gamma}{4\pi pq} \tilde{r}^{-\gamma}(1 + \tilde{r})^{-(4 - \gamma)},
\]

studied extensively by a number of authors (Merritt & Fridman 1996; Valluri & Merritt 1998; Wachlin & Ferraz-Mello 1998; Siopis 1999).

(iii) Scale-free (single power-law): \( \rho(\tilde{r}) = \tilde{r}^{-\gamma} \), studied in Terzić (2002).

(iv) Anisotropic harmonic oscillator: \( \Phi(\tilde{r}) = \tilde{r}^2 \), used in Kandrup & Sideris (2002).

Here, \( \tilde{r} = (x^2 + y^2 + z^2 + p^2)^{1/2} \) is the elliptical radius, and axis ratios \( q = y/x, p = z/x (p \leq q \leq 1) \) are defined for the potential (in the case of logarithmic and harmonic potentials) or for the density (in other cases). \( x, y \) and \( z \) are the longest, intermediate and short axes, correspondingly. A central supermassive BH may be added to any potential (actually it is modelled as a Newtonian potential, since general relativistic effects are presumably not important for the global dynamics in the galaxy).

In addition to these models, there are several more flexible options. One is a representation of a potential–density pair in terms of a finite number of basis functions with certain coefficients, and evaluation of forces and their derivatives as a sum over these functions, for which analytical expressions exist. For not very flattened systems, an efficient choice is to write each member of the basis set as a product of a function depending on radius and a spherical harmonic:

\[
\Phi(r, \theta, \phi) = \sum_{n=0}^{n_{\text{max}}} \sum_{l=0}^{l_{\text{max}}} \sum_{m=-l}^{l} A_{nlm} \Phi_{nlm} Y_{l}^{m}(\theta, \phi). \tag{4}
\]

The idea to approximate a potential–density pair by a finite number of basis functions with given coefficients goes back to Clutton-Brock (1973), who used a set of functions based on the Plummer model, suitable to deal with cored density profiles. Later, Hernquist & Ostriker (1992) introduced another class of basis functions, adapted for the cuspy Hernquist (1990) profile, to use in their self-consistent field (SCF) N-body method. Zhao (1996a) introduced a generalized \( \alpha \)-model including two previous cases; there exist also other bases sets for near-spherical (Allen, Palmer & Papaloizou 1990; Rahmati & Jalali 2009) or flattened near-axisymmetric (e.g. Brown & Papaloizou 1998) systems. We use the basis set of Zhao (1996a), which is general enough to accommodate both cuspy and cored density profiles, with a suitably chosen parameter \( \alpha \). (For a discussion on the effect of different basis sets see Carpineto & Wachlin 2006; Kalapotharakos, Efthymiopoulos & Voglis 2008). This formalism and associated formulae are presented in the Appendix A2, and tests for accuracy and recipes for choosing \( \alpha \) are discussed in Section 6. A variation of this approach is to numerically construct a basis set whose lowest order function is specifically tuned for a particular mass distribution, and higher order terms are derived according to a certain procedure involving orthogonalization of the whole set (Saha 1993; Brown & Papaloizou 1998; Weinberg 1999). We propose another, conceptually simple approach described below.

Instead of requiring radial functions to form an orthogonal basis set, we may represent them as arbitrary smooth functions, namely, splines with some finite number of nodes:

\[
\Phi(r, \theta, \phi) = \sum_{l=0}^{l_{\text{max}}} \sum_{m=-l}^{l} \Phi_{lm}(r) Y_{l}^{m}(\theta, \phi). \tag{5}
\]

The advantage over the basis set approach is its flexibility – one may easily adapt it to any underlying potential model; the number and radii of nodal points may be chosen arbitrary (however, it is easier to have them equal for all spherical harmonics); the evaluation of potential at a given point depends only on the coefficients at a few nearby nodes rather than on the whole basis set. Derivatives of the potential up to the second order are continuous and easily evaluated. A similar approach was used for Schwarzschild modelling in Valluri et al. (2004) and Siopis et al. (2009) for an axially symmetric potential. In the context of N-body simulations, a related ‘spherical-harmonic expansion’ method, pioneered by Aarseth (1967); van Albada & van Gorkom (1977); White (1983), computes coefficients at each particle’s location, optionally introducing softening to cope with force divergence as two particles approach each other in radius. In the variant proposed by McGlynn (1984); Sellwood (2003), coefficients of angular expansion are evaluated at a small set of radial grid points; radial dependence of forces is then linearly interpolated between grid nodes while the angular dependence is given by truncated spherical harmonic expansion. This potential solver is also used in the made-to-measure method of de Lorenzi et al. (2007).

Both basis-set (BSE) and spline expansions may be constructed either for a given functional form of density profile (not requiring the corresponding potential to be known in a closed form), or from a finite number of points representing Monte Carlo sampling of a density model. In the former case, one may use an analytic density model from a pre-defined set (e.g. Dehnen, Plummer, isochrone, etc.), or a flexible smooth parametrization of an arbitrary density profile by a Multi-Gaussian expansion (Emsellem, Monnet & Bacon 1994; Cappellari 2002). In the case of initialization from an N-body snapshot, the spline coefficients are calculated by a method.
suggested in Merritt (1996), similar to non-parametric density estimators of Merritt & Tremblay (1994); first the angular expansion coefficients are evaluated at each particle’s radius, then a smoothing spline is constructed which approximates the gross radial dependence of these coefficients while smoothing local fluctuations. More details on the spline expansion method are given in the Appendix A3, and the tests are presented in Section 6.

Yet another option is to use directly the potential of N-body system of ‘frozen’ (fixed in place) bodies. We use the potential solver based on the Barnes & Hut (1986) tree-code algorithm. It can represent almost any possible shape of the potential, but is much slower and noisier in approximating a given smooth density profile. It may use a variable softening length, the optimal choice for which is discussed in Section 6.3.

4 ORBIT ANALYSIS

4.1 Orbit integration

Orbit integration is performed by the dop853 algorithm (Hairer, Nørsett & Wanner 1993), which is an eight-order Runge–Kutta scheme with adaptive step size. It is well suited for a subsequent Fourier analysis of trajectory because of its ability to produce ‘dense output’, i.e. interpolated values of the function at arbitrary (in particular, equally spaced) moments of time. The energy of orbit is conserved to the relative accuracy typically better than $10^{-8}$ per 100 dynamical times for all smooth potentials except the Spline expansion, which is only twice continuously differentiable and hence demonstrates a lower (but still sufficient for most purposes) energy conservation accuracy ($\sim 10^{-5}$–$10^{-6}$) in the high-order integrator.

For the frozen N-body potential, we use a leap-frog integrator with adaptive timestep selection and optimal timestep symmetrization (Hut, Makino & McMillan 1995) which reduces secular energy drift. The reason for using a lower order integrator is that the potential of the tree-code is discontinuous: when a trajectory crosses a point at which a nearby tree cell is opened (i.e. decomposed into sub-elements), which occurs when the distance to the cell is smaller than the cell size divided by the opening angle $\theta$, the potentials of an unresolved and resolved cell do not match. Therefore, the energy of an orbit is not well conserved during integration, no matter how small timesteps are. The error in potential approximation rapidly decreases with decreasing $\theta$; however, computational cost also increases quickly. Overall, for $\theta < 0.5$ the accuracy of energy conservation is $\sim 10^{-3}$; for more discussion see Barnes & Hut (1989).

For a given value of energy the period of long(x)-axis orbit with the same energy is calculated, and it is used as a unit of time (hereafter $T_{dyn}$, dynamical time) and frequency in the following analysis. (This is different from most studies that use the period of closed loop orbit in $x$–$y$ plane for this purpose, but our definition has the advantage of being the longest possible period of any orbit with a given energy).

Visualization of orbits is quite an important tool; orbit may be rendered either in projection on one of the three principal planes or in 3D. Additionally, an algorithm for rendering orbit as a solid body is implemented, which is based on Delaunay tessellation of set of points comprising the orbit, and removal of hidden surfaces to leave out only the outer boundary of the volume that the orbit fills (Fig. 1).

4.2 Frequency analysis and orbit classification

Orbits are classified by their spectra using the following scheme, based on Carpintero & Aguilar (1998) (which, in turn, is an improvement of the method proposed by Binney & Spergel 1982). First, we obtain the complex spectra of each spatial coordinate $x_i(t)$ by Fourier transform. Then we extract the most prominent spectral lines $\omega_{i,j}$ for each coordinate: for each line, its amplitude and phase is determined using Hunter’s DFT method (Hunter 2002); other studies used similar techniques based on Laskar’s NAFF algorithm (Laskar 1993; Valluri & Merritt 1998) or FMFT (Šidlichovský & Nesvorný 1997). All these methods employ Hanning window filtering on input data, which enhances the accuracy of frequency determination relative to the simple Fourier transform, used in the pioneering work of Binney & Spergel (1982). The contribution of each detected line to the complex spectrum is subtracted and the process is repeated until we find 10 lines in each coordinate or the amplitude of a line drops below $10^{-3}$ times the amplitude of the first line. Finally, the orbit classification consists in analysing the relations between $\omega_{i,j}$.

A regular orbit in three dimensions should have no more than three fundamental frequencies $\Omega_i$, so that each spectral line may be expressed as a sum of harmonics of these fundamental frequencies with integer coefficients: $\omega_{i,j} = \sum_{k=1}^{3} a_{ijk} \Omega_k$. For some orbits the most prominent lines$^3$ $\omega_{i,1} \equiv \omega_i$ in each coordinate $c$ (called LFCCs – leading frequencies in Cartesian coordinates) coincide with the fundamental frequencies, these orbits are non-resonant boxes. For others, two or more LFCCs are in a resonant $m : n$ relation ($m \omega_{1,1} = n \omega_{2,1}$ with integer $m$, $n$). These orbits belong to the given resonant family, in which the parent orbit is closed in the $c_1$–$c_2$ plane. The additional fundamental frequency corresponds to the libration about this parent orbit, and is given by the difference between frequencies of the main and satellite lines. Besides this, in 3D systems there exist an important class of thin orbits with the three leading frequencies being linearly dependent (with integer coefficients): $l \omega_{1,1} + m \omega_{2,1} + n \omega_{3,1} = 0$. These are labelled as $(l, m, n)$ thin orbits (of the three numbers at least one is negative) and are easily

$^3$ Our choice of unit frequency makes it possible to put the LFCC in the correct order $1 \leq \omega_i \leq \omega_j \leq \omega_k$, even if these lines are not the largest in amplitude.
identified on a frequency map (see Section 4.4). In fact, resonant orbits are a sub-class of thin orbits: \((m, -l, 0)\) thin orbit may be alternatively termed as \(l : m : n^*\) resonance and \((mn, nl, -lm)\) orbit is named \(l : m : n\) resonance. The origin of term ‘thin’ lies in the fact that a parent orbit of such an orbit family is indeed confined to a 2D surface in the configuration space (possibly self-crossing). This parent orbit has only two fundamental frequencies, and in the case of a closed orbit there is only one frequency. All orbits belonging to the associated family of a particular thin orbit have additional fundamental frequencies which may be viewed as libration frequencies around the parent orbit. These thin orbit families are particularly important for triaxial models with cuspy density profiles, as they replace classical box orbits and provide the structure of the phase space (Merritt & Valluri 1999).

The most abundant sub-class of resonant orbits are tube orbits, which have a 1:1 relation between frequencies in two coordinates. They have a fixed sense of rotation around the remaining axis. Accordingly, \(*:1:1\) resonant orbits are called LAT (\(x_-,\) or long-axis tubes), and \(1:1:*\) orbits are SAT (\(z_,\) short-axis tubes); there are no stable tube orbits around the intermediate axis. However, some of chaotic orbits may also have a 1:1 correspondence between leading frequencies, but may not have a definite sense of rotation; in this case they are not labelled as tube orbits.

All other orbits which are not tubes, thin orbits or resonances, are called box orbits.\(^4\) Classification in two dimensions is similar but simpler: there exist only boxes and \(mn:n\) resonances (of which 1:1 are tubes), which also may be regular or chaotic.

Those orbits which display more complicated spectra (not all lines are expressed as linear combination of two/three frequencies) are additionally labelled chaotic (based on analysis of spectra), although this criterion for chaos is less strict than those discussed below. Note that we do not have a special class for chaotic orbits: they fall into the most appropriate basic class (usually box, but some weakly chaotic orbits are also found among tubes and other resonances).

There is an additional criterion for chaoticity of an orbit: if we take the difference between corresponding fundamental frequencies \(\Omega^{(1)}_c\) and \(\Omega^{(2)}_c\) calculated on the first and second halves of the orbit, then this ‘frequency diffusion rate’ (FDR) is large for chaotic orbits and small for regular ones (Laskar 1993). We use the LFCC diffusion rate, defined as the average relative change of three frequencies:

\[
\Delta \omega = \frac{1}{3} \sum_{c=1}^{3} \frac{\left| \omega^{(1)}_c - \omega^{(2)}_c \right|}{\left( \omega^{(1)}_c + \omega^{(2)}_c \right) / 2}.
\]  

(6)

To account for the possibility of misinterpretation of two lines with similar amplitudes, frequencies that have relative difference greater than 0.5 are excluded from this averaging. Fig. 2 (right-hand panel) shows an example of a weakly chaotic \((\Delta \omega \sim 10^{-2})\) orbit in the vicinity of the \(1 : 2 : -1\) resonance, for which the leading frequencies do not form isolated peaks but rather clusters of lines, demonstrating that the complex spectrum is not described by just one line in the vicinity of a peak. Consequently, lines in these clusters change erratically in amplitude and position between the first and last halves of the orbit, which contributes to the rather high value of \(\Delta \omega\).

The FDR is, unfortunately, not a strict measure of chaos, nor is it well defined by itself. The spectrum of values of FDR is typically continuous with no clear distinction between regular and chaotic orbits, which in part is due to the existence of ‘sticky’ chaotic orbits (e.g. Contopoulos & Harsoula 2010), which resemble regular ones for many periods, and consequently have low FDR. If one computes the FDR over a longer interval of time, these sticky orbits may become unstuck and demonstrate a higher value of FDR. On the other hand, a perfectly regular orbit may sometimes have a rather large FDR because of two very close spectral lines with comparable amplitudes, which both degrades the accuracy of frequency determination and increases the time required for an orbit to fill its invariant torus uniformly. Over a longer interval, these nearby lines would be better resolved and such an orbit would attain a substantially lower FDR, which for regular orbits is ultimately limited by the accuracy of energy conservation.

A more fundamental problem is that even for the same time span the FDR is not a strictly defined quantity itself, i.e. it may vary by a factor of few when measured for two successive time intervals. This can be understood from the following simplified argument: suppose the ‘instant’ value of frequency is a random quantity with a mean value \(\omega_0\) and a dispersion \(\delta \omega\), and the frequency measured over an interval is simply an average of this quantity. Then the FDR over any interval is a random quantity with dispersion \(\sim \sqrt{2} \delta \omega\), or, if we take the distribution of \(\log |\Delta \omega|\), it will be peaked around \(\log \delta \omega\) with a scatter of \(\sim 1/2\) dex. Similar uncertainty relates the values of FDR calculated for two different intervals of time. Indeed, for a particular case of a triaxial Dehnen potential we find that the correspondence between \(\log \Delta \omega\) measured for two different intervals of the same orbit is well described by the following probability distribution: \(P(\log |\Delta \omega^{(1)} - \log |\Delta \omega^{(2)}| = X) = \frac{1}{2} \exp(-3|X|)\) (Fig. 3).

To summarize, FDR is an approximate measure of chaos with uncertainty of 0.3—0.5 orders of magnitude. Yet it correlates with the other chaos indicator, the Lyapunov exponent.

4 In the vicinity of the central BH, box orbits are replaced by regular pyramid orbits (Merritt & Valluri 1999).

Figure 2. Spectral analysis of two orbits with slightly different initial conditions in \(y = 1\) Dehnen model. The left one is \(1 : 2 : -1\) regular thin tube orbit, the right is weakly chaotic, sticky orbit in the vicinity of the same resonance. Shown in red/green/blue are amplitudes of complex spectra of motion in \(x/y/z\) coordinate, and vertical lines denote identified lines. Although the spectra look similar at first sight (when looking at amplitudes only), clustering of lines at the same frequency in the right-hand panel hints at the chaotic nature of the orbit: after the subtraction of a line from the complex spectrum, its amplitude decreases only weakly, and the next line is found at almost the same frequency. By contrast, in the left-hand panel all satellite lines are clearly identified and distinct, and the whole spectrum is a sum of linear combinations of just three fundamental frequencies with integer coefficients.

4.3 Lyapunov exponent

The Lyapunov exponent (or, more precisely, the largest Lyapunov exponent) \(\Lambda\) is another measure of chaoticity of an orbit. Given a trajectory \(x(t)\) and another infinitely close trajectory \(x + w(t)\), we
follow the evolution of deviation vector $\mathbf{w}$. For a regular orbit, the magnitude of this vector, averaged over some time interval longer than the orbit period, grows at most linearly with $t$; for a chaotic one it grows exponentially, and

$$\Lambda \equiv \lim_{t \to \infty} (\ln |\mathbf{w}|)/t . \quad (7)$$

The usual method of computing $\Lambda$ is integration of the variational equation (e.g. Skokos 2010) along with the orbit, or simply integration of a nearby orbit. While the first method is more powerful in the sense that it may give not only the largest, but in principle the whole set of Lyapunov numbers (Udry & Pfenniger 1988), it requires the knowledge of the second derivatives of the potential. The second method is relatively straightforward – one needs to integrate the same equations of motions twice, and compute the deviation vector as the difference between the two orbits. The only issue is to keep $\mathbf{w}$ small, that is, the orbits must stay close despite the exponential divergence. Thus, $\mathbf{w}$ should be renormalized to a very small value each time it grows above certain threshold (still small enough, but orders of magnitude larger than initial separation); however, one should be careful to avoid false positive values of $\Lambda$ that may appear due to roundoff errors (Tancredi, Sánchez & Roig 2001), in particular, for orbits that come very close to the central BH.

In practice, one may calculate only the finite-time approximation for the true Lyapunov exponent for a given integration time. Equation (7) shows that such a finite-time estimate for a regular orbit decreases as $\sim r^{-1}$; therefore, the usually adopted approach is to find a threshold value for the finite-time estimate of $\Lambda$ that roughly separates regular and chaotic orbits. We use the following improved method that does not require to define such a threshold and gives either a non-zero estimate for $\Lambda$ in the case of a chaotic orbit, or zero if the orbit was not detected to be chaotic. For a regular orbit – or for some initial interval of a chaotic orbit – $|\mathbf{w}|/t$ fluctuates around some constant value (the period of fluctuations corresponds to characteristic orbital period; to eliminate these oscillations, we use a median value of $|\mathbf{w}|$ over the interval $2T_{\text{dyn}}$). When (and if) the exponential growth starts to dominate, $\Lambda$ may be estimated as the average value of $(\ln |\mathbf{w}|)/t$ over the period of exponential growth. If no such growth is detected, then $\Lambda$ is assumed to be zero (or, more precisely, an upper limit may be placed). The exponential growth regime is triggered when the current value of $|\mathbf{w}|/t$ is several times larger than the average over previous time. In addition, we normalize $\Lambda$ to the characteristic orbital frequency, so that its value is a relative measure of chaotic behaviour of an orbit independent of its period (so-called specific finite-time Lyapunov characteristic number, Voglis, Kalapotharakos & Stavroulopoulou 2002). Our scheme is summarized in Fig. 4. One should keep in mind that the ability of detecting chaotic orbits by their positive Lyapunov exponent depends on the interval of integration: for weakly chaotic orbits the exponential growth starts to manifest itself only after a long enough time. The finite-time estimate for $\Lambda$ may also depend on the integration time because the growth of $\mathbf{w}$ is not exactly exponential and exhibits fluctuations before reaching asymptotic regime.

The two chaos indicators – the FDR $\Delta \omega$ (6) and the Lyapunov exponent $\Lambda$ (7) – are based on different methods yet demonstrate a rather good agreement in the chaos detection (see e.g. Maffione et al. (2013) for a detailed comparison of various chaos indicators based on variational equation and on frequency analysis). Fig. 5 presents a comparison between $\Delta \omega$ and $\Lambda$ for 10$^4$ orbits in a particular triaxial Dehnen model. It demonstrates that orbits labelled as regular or chaotic, based on $\Delta \omega$, have quite well-separated distributions in $\Delta \omega$, with the overlap being comparable to the intrinsic uncertainty of FDR determination. Moreover, for orbits with $\Lambda > 0$ there is a clear correlation between the two chaos indicators. Meanwhile, the distribution in $\Delta \omega$ and $\Lambda$ is quite different for intervals of 100 and 500$T_{\text{dyn}}$, as explained above, and the threshold $\Delta \omega_{\text{th}}$ separating regular and chaotic orbits does depend on the integration time.

### 4.4 Frequency map

Frequency map is a convenient and illustrative tool for analysing orbital structure of a potential (Papaphillipou & Laskar 1998; Valluri...
identify the regions of phase space which contain mostly regular or chaotic orbits and highlights the most prominent resonances.

The Poincaré surface of section (e.g. Lichtenberg & Lieberman 1992) is another important tool for analysing orbital structure of a two-dimensional potential, as well as for studying orbits confined to one of the principal planes in a 3D potential. This tool is also implemented in SMILE.

5 SCHWARZSCHILD MODELLING

5.1 Spherical mass models

Before discussing the Schwarzschild method itself, we first outline the formalism used to deal with spherical isotropic mass models with an arbitrary density profile. These models are constructed from an array of \( \{r, M(r)\} \) pairs specifying the dependence of enclosed mass on radius. The mass, potential and other dynamical properties are represented as spline functions in radius (with logarithmic scaling and careful extrapolation to small and large radii beyond the definition region, in the way similar to the Spline potential approximation). The unique isotropic distribution function is given by the Eddington inversion formula (Binney & Tremaine 2008):

\[
f(E) = \frac{1}{\sqrt{8\pi^2}} \int_0^\infty \frac{d^2\Phi}{d\Phi^2} \frac{d\Phi}{\sqrt{\Phi - E}}.
\]

These spherical models are used throughout SMILE in various contexts, in particular, to generate the initial conditions for the
orbit library used to create Schwarzschild models: for the given triaxial potential, a spherically symmetric approximation model is created and the initial conditions are drawn from its distribution function. Of course, one may also use such a model to create a spherically symmetric isotropic $N$-body model with a given density profile (there is a separate tool, **SphSpherical**, that does just that). The approach based on the distribution function gives better results than using Jeans equations with a locally Maxwellian approximation to the velocity distribution function (Kazantzidis, Magorrian & Moore 2004). Of course, even the simplest variants of Jeans models can account for varying degree of velocity anisotropy, but there also exist methods to derive anisotropic distribution function for spherical models (e.g. Ossipkov 1979; Merritt 1985; Cuddeford 1991; Baes & van Hese 2007). As we use spherical models only as a "seed" to construct a more general Schwarzschild model, a simple isotropic distribution function is enough for our purposes.

Another application of these models is to study dynamical properties of an existing $N$-body snapshot, for instance, the dependence of dynamical and relaxation time on radius or energy. To create such a model from an $N$-body snapshot, we again use the penalized spline smoothing approach. That is, we first sort particles in radius and define a radial grid of roughly logarithmically spaced $N_{\text{grid}} \sim 10$–50 points so that each interval contains at least 10–50 of particles. Then the $M(r)$ profile is obtained by fitting a penalized spline to the scaled variable $\mu(\xi \equiv \log r) \equiv \log (M(r)/M_{\infty} - M(r))$. The degree of smoothing may be adjusted to obtain a distribution function that is not very much fluctuating (or at least does not become negative occasionally). An alternative approach would be, for instance, fitting a local power-law density profile whose parameters depend on radius and are computed from a maximum likelihood estimate taking into account nearby radial points (e.g. Churazov et al. 2009). The creation of spherical models from $N$-body snapshots is also implemented as a separate tool; in **SMILE** the same is done using the intermediate step of basis-set or Spline potential initialization.

### 5.2 Variants of Schwarzschild method

Each orbit in a given potential is a solution of CBE (2): the distribution function is constant over the region of phase space occupied by the orbit, provided that it was integrated for a sufficiently long time to sample this region uniformly. The essence of the Schwarzschild’s orbit superposition method is to obtain the self-consistent solution of both CBE and the Poisson equation (1) by combining these individual elements with certain weights to reproduce the density profile consistent with the potential used to integrate the orbits, possibly with some additional constraints (e.g. kinematical data). It is clear that such a superposition is sought only in the configuration space, i.e. involves only the density or the potential created by individual orbits; for instance, one may write the equation for the density

$$\rho(r) = \sum_{\omega=1}^{N_{\text{seg}}} w_{\omega} \rho_{\omega}(r),$$

where $w_{\omega} \geq 0$ are the orbit weights to be determined, and each orbit has a density profile $\rho_{\omega}(r)$. This equation can be approximately solved by discretizing both the original density profile and that of each individual orbit into a sum of certain basis functions, thus converting the continuous equation (9) into a finite linear system. The classical Schwarzschild method consists of splitting the configuration space into $N_{\text{e}}$ cells, computing the required mass $m_{\text{e}}$ in each cell from the original density profile, and recording the fraction of time $t_{\text{e}}$ that $o$-th orbit spends in $c$-th cell. Then the linear system of equations reads

$$m_{c} = \sum_{\omega=1}^{N_{\text{seg}}} w_{\omega} t_{\omega c}, \quad c = 1..N_{\text{e}}.$$  \hspace{1cm} (10)

We may generalize the above definition to replace cells with some arbitrary constraints to be satisfied exactly or as closely as possible. Two such alternative formulations naturally arise from the definitions of BSE and Spline potential expansions. Namely, we may use the coefficients of potential expansion of the original model as target constraints $m_{\text{e}}$, compute the expansion coefficients from the mass distribution of each orbit as $t_{\omega c}$ and then find the orbit weights $w_{\omega}$ so that the weighted sum of these coefficients reproduces the total potential. The linear character of orbit superposition is preserved in the potential expansion formalism. Accordingly, we introduce two additional variants of Schwarzschild models, named after the potential expansions: the BSE model and the Spline model; the original, grid-based formulation is called the Classic model. The BSE model is analogous to the SCF method of Hernquist & Ostriker (1992), with the difference that the coefficients are built up by summing not over individual particles, but over entire orbits.

An important difference between the Classic and the two new variants of the Schwarzschild method is that in the latter case, the basis functions of density expansion and the elements of the matrix $t_{\omega c}$ and the vector $m_{\text{e}}$ in (10) are not necessarily non-negative. In other words, a single orbit with rather sharp edges, when represented by a small number of expansion coefficients, does not necessarily have a positive density everywhere; however, when adding up contributions from many orbits the resulting density is typically well behaved (as long as the target expansion had a positive density everywhere). On the other hand, the classical Schwarzschild method ensures only that the average density within each cell is equal to the required value, and does not address the issue of continuous variation of density across grid boundaries. The basis functions in the classical formulation are $\Box$-shaped functions with finite support and sharp edges, while in the proposed new variants these are smooth functions. Recently Jalali & Tremaine (2011) suggested another generalization of the Schwarzschild method (although presently only for the spherical geometry), in which density, velocity dispersion and other quantities are represented as expansions over smooth functions with finite support, and additional Jeans equation constraints are used to improve the quality of solution.

The partitioning of configuration space in the Classic model is done in the similar way as in Siopis (1999). We define $n_{\text{shell}} + 1$ concentric spherical shells at radii $r_i$ (the last shell’s outer boundary goes to infinity, and this shell is not used in modelling). By default, shells are spaced in radii to contain approximately the same mass each, but this requirement is not necessary, and one may build a grid with a refinement near the centre. Shells are further divided by the following angular grid. The sphere is split into three sectors by planes $x_i = x_j$; then each sector is divided into $n_{\text{seg}} \times n_{\text{seg}}$ rectangles by lines $x_i/x_j = \tan(\pi/2 n_{\text{seg}})$ ($n = 1..n_{\text{seg}}$). This way we get $3 n_{\text{seg}}^2 n_{\text{shell}}$ cells (Fig. 7). The time $t_{\omega c}$ that an orbit spends in each cell is calculated with great precision thanks to the dense output feature: if two subsequent points on a trajectory fall into different cells, we find the exact moment of cell boundary crossing by nested binary divisions of the interval of time (this may in turn reveal a third cell in between, etc.). This approach is more straightforward and precise than used in Siopis (1999); Terzić (2002).
In the BSE and Spline models, we simply compute coefficients of expansion for each orbit, so that there are \( n_{\text{rad}} \times (l_{\text{max}}/2 + 1) \times (l_{\text{max}}/2 + 2)/2 \) constraints in the model, where \( n_{\text{rad}} \) is the number of radial basis functions in the BSE model or the number of radial points in the Spline models (which are chosen in the same way as the concentric shells in the Classic model), and \( l_{\text{max}} \) is the order of angular expansion (the factor of 1/2 comes from using only the even harmonics). Unlike the initialization of the Spline potential from a set of discrete points, we do not perform penalized spline smoothing for orbit coefficients, to keep the problem linear.

In all three variants, we also constrain the total mass of the model and may have additional (e.g. kinematic) constraints; at present, one fairly simple variant of kinematic data modelling is implemented, which sets the velocity anisotropy profile as a function of radius. The configuration space is split into shells and the mean squared radial and transversal velocity \( v_{r,\text{os}}^2 \) and \( v_{\theta,\text{os}}^2 \) of each orbit in each shell is recorded. Then the following quantity is used as a constraint required to be zero:

\[
\sum_{o=1}^{N_c} w_o (v_{r,\text{os}}^2 - 2(1 - \beta) v_{\theta,\text{os}}^2). \tag{11}
\]

Here \( \beta \) is the required value of the velocity anisotropy coefficient \( \beta \equiv 1 - \sigma_r^2/2\sigma_{\theta}^2 \) (Binney & Tremaine 2008) in the given radial shell. In the practical application of the Schwarzschild method to the modelling of individual galaxies, the projected velocity distribution function is usually constrained; this is easy to add in the general formalism implemented in SMILE.

Traditional approach to the construction of the orbit library is the following: choose the number of energy levels (typically corresponding to potential at the intersection of \( N \)-th radial shell with \( x \)-axis) and assign initial conditions at each energy shell similarly to that of frequency map (stationary and principal-plane start-spaces). The outer shell may be taken as either equipotential or equidensity surface, whichever is rounder, so that all finite spatial cells are assured to be threaded by some orbits. However, we found that such discrete distribution in energy and in initial positions is not welcome when translating the Schwarzschild model to its \( N \)-body representation. Step-like distribution in energy tends to relax to continuous one, which introduces systematic evolution (Vasiliev & Athanassoula 2012); in addition, the coarse radial resolution \( (n_{\text{shell}} \sim 20-50) \) does not allow us to sample the innermost particles well enough. Therefore, we draw the initial conditions for the orbit library from the spherical mass model discussed in the previous section, constructed for the given potential.

### 5.3 Solving the optimization problem

The linear system (10) may be reformulated as an optimization problem, introducing auxiliary variables \( \delta_c \), which are deviations between the required and the actual constraint values, normalized to some scaling constants \( \eta_c \):

\[
\delta_c = \left( \sum_{o=1}^{N_o} w_o l_{\text{loc}} - m_c \right) / \eta_c, \quad c = 1..N_c. \tag{12}
\]

Obviously, we seek to minimize \( |\delta_c| \), ideally making them zero, but in addition one may require that some other relations be satisfied as closely as possible, or to within a predefined tolerance, or that a certain functional of orbit weights be minimized (e.g. the sum of weights of chaotic orbits). A rather general formulation of this problem is to introduce an objective (penalty) function \( F \) and find

\[
\min F(w_o), \quad \text{subject to the constraints } w_o \geq 0.
\]

Various studies have adopted different forms for the penalty function and different methods to find the minimum.

One could incorporate the requirement of exact match between required and calculated mass in each cell (set \( \delta_c = 0 \) as linear constraints); however, this is not physically justified – given a number of other approximations used in modelling, sampling orbits etc., it is unreasonable to require strict equality. Instead, one may penalize the deviation of \( \delta_c \) from zero and search for the solution that minimizes this penalty. (The scaling coefficients \( \eta_c \) may be taken as some ‘typical’ values of \( m_c \), to give roughly similar significance to the deviations \( \delta_c \).) This may be done in various ways: for instance, one may take

\[
F = \alpha \sum_{c=1}^{N_c} \delta_c^2 + F_{\text{additional}}, \tag{13}
\]

(this is called non-negative least-square method, used in Merritt & Fridman 1996; Zhao 1996b), or introduce additional non-negative variables and use

\[
F = \alpha \sum_{c=1}^{N_c} (\mu_c + \nu_c) + F_{\text{additional}}; \quad \mu_c, \nu_c \geq 0, \quad \delta_c = \mu_c - \nu_c, \tag{14}
\]

this effectively reduces to \( F = \alpha \sum |\delta_c| \); the latter approach was used in Siopis (1999); Terzić (2002). Here \( \alpha \) is the penalty coefficient discussed below. In principle, the two formulations do the same job – if possible, reach the exact solution, if not, attain the ‘nearest possible’ one. Another option is to drop the condition that \( \delta_c = 0 \) and instead require that \( |\delta_c| \leq \beta |m_c| \), where \( \beta \) is the allowed fractional deviation from the exact constraint value (for example, 1 per cent); this approach was adopted in van den Bosch et al. (2008). It may be reformulated as the standard non-negative optimization problem by introducing additional variables \( \mu_c, \nu_c \geq 0 \) and doubling the number of equations:

\[
\delta_c = \mu_c - \nu_c, \quad \mu_c + \nu_c = \beta |m_c|, \quad c = 1, \ldots, N_c. \tag{15}
\]
We have implemented the last two approaches – either a tolerance range defined by fractional constraint deviation $\beta$, or a linear term in the penalty function proportional to $\alpha$. (In principle, a combination of both variants is trivial to implement). The first approach is also easily formulated in terms of the quadratic optimization problem, but it involves a dense matrix of quadratic coefficients with size $N_o \times N_o$, and hence is less practical from the computational point of view. However, such quadratic problems with a specific form of the penalty function are efficiently solved with the non-negative least-squares method (Lawson & Hanson 1974).

When Schwarzschild modelling is used to construct representations of observed galaxies, one usually computes the observable quantities (surface density and line-of-sight velocity distribution as functions of projected position) in the model ($Q_{i, \text{mod}}$) and minimize their deviation from the actual observations $Q_{i, \text{obs}}$, normalized by the measurement uncertainties $\Delta Q_{i}$, via:

$$\chi^2 = \sum_{i=1}^{N_o} \left( \frac{Q_{i, \text{mod}} - Q_{i, \text{obs}}}{\Delta Q_{i, \text{obs}}} \right)^2 + F_{\text{additional}}. \quad (16)$$

Then one seeks to minimize $\chi^2$ and derive the confidence intervals of the model parameters based on standard statistical $\Delta \chi^2$ criteria. In this approach, the self-consistency constraints for the density model (10) may be either included in the same way as the observational constraints, with some artificially assigned uncertainty $\Delta Q_i$ (e.g. Valluri et al. 2004), or as tolerance intervals via (15) (e.g. van den Bosch et al. 2008).

An important feature of Schwarzschild modelling is that a solution to the optimization problem, if exists, is typically highly non-unique. In principle, the number of orbits that are assigned non-zero weights may be as low as the number of constraints (which is typically $\geq 10$ times smaller than the total number of orbits $N_o$); however, in some studies the opposite inequality is true, e.g. Verolme et al. (2002) had four times more constraints than orbits. While this is a solution to the problem in the mathematical sense, it is often unacceptable from the physical point of view: large fluctuations in weights of nearby orbits in phase space are almost always unwelcome. For this reason, many studies employ additional means of ‘regularization’ of orbit weight distribution, effectively adding a functional of orbit weights $F_{\text{additional}}$ to the penalty function $\chi$ or $\chi^2$. There are two conceptually distinct methods of regularization (which are not mutually exclusive): ‘local’ try to achieve smoothness by penalizing large variations in weight for orbits which are close in the phase space, according to some metric; ‘global’ intend to minimize deviations of orbit weights from some pre-determined prior values (most commonly, uniform or flat priors).

The first approach (e.g. Zhao 1996b; Cretton et al. 1999; Verolme et al. 2002; van den Bosch et al. 2008) minimizes the second derivatives of orbit weight as a function of initial conditions assigned on a regular grid in the start-space, in which the proximity of orbits is determined by indices of the grid nodes. Since in our code we do not use regularly spaced initial conditions, this method is not applicable in our case. The second kind of regularization assumes some prior values $\tilde{w}_o$ for orbit weights (most commonly, uniform values $\tilde{w}_o = M_{\text{total}}/N_o$, although Morganti & Gerhard (2012) argue for on-the-fly adjustment of the weight priors in the context of made-to-measure modelling). Then a penalty term is added to the cost function which minimizes the deviations of the actual weights from these priors. This could be done in various ways. Richstone & Tremaine (1988) introduced the maximum entropy method, in which $F_{\text{additional}} = -\lambda S$, where $\lambda$ is the regularization parameter, and $S$ is the (normalized) entropy, defined as:

$$S = -\frac{1}{M_{\text{total}}} \sum_{o=1}^{N_o} w_o \ln(w_o/\tilde{w}_o), \quad M_{\text{total}} = \sum_{o=1}^{N_o} w_o. \quad (17)$$

This method was used in Gebhardt et al. (2000); Thomas et al. (2004); Siopis et al. (2009). Another possibility is to use a quadratic regularization term in the penalty function:

$$F_{\text{additional}} = \frac{\lambda}{N_o} \sum_{o=1}^{N_o} (w_o/\tilde{w}_o)^2. \quad (18)$$

This approach, used in Merritt & Fridman (1996), Siopis (1999), Valluri et al. (2004), gives very similar results to the maximum entropy method, but is simpler from computational point of view, since the regularization term is quadratic in $w_o$ and not a non-linear function as in (17). We adopted this second variant with uniform weight priors $\tilde{w}_o$, which is not a bad assumption given that our method of assigning initial conditions populates the phase space according to the isotropic distribution function for a given density profile.

In the context of Schwarzschild modelling of observational data, the regularization is considered a necessary ingredient because the number of observational constraints is typically much smaller than the number of free parameters (orbit weights), so that some smoothing is desirable to prevent overfitting (fitting the noise instead of actual physical properties of the system). The amount of smoothing is then controlled by a regularization parameter $\lambda$ which scales the contribution of the penalty term to the cost function, and there are standard statistical methods of determining the optimal value of this parameter (e.g. cross-validation technique, Wahba 1990). Most commonly, the regularization coefficient is chosen as to achieve maximal smoothing which still does not deteriorate the quality of the fit more than by some acceptable value of $\Delta \chi^2$. In our application of Schwarzschild method to the construction of models with pre-defined, noise-free properties, the necessity of smoothing is not obvious a priori. The linear system (10) either has no solutions or infinitely many solutions, and unless some non-linear objective function is used, the solution will tend to have only $\sim N_o$ orbits with non-zero weights. Vasiliev & Athanassoula (2012) have shown that a solution with a larger number of orbits (lower average orbit weight) is more stable in the $N$-body simulation (in addition to be smoother and more aesthetically pleasant), therefore it is preferable to use regularization to select a solution with $O(N_o)$ rather than $O(N_o)$ effective orbits from all possible set of solutions that satisfy all constraints. Since we do not have any trade-off between fit quality and smoothness, the value of regularization parameter $\lambda$ itself is not important, only insofar as it should not outweigh the penalties for constraint violation in (14), parametrized by the coefficient $\alpha$.

An important issue in modelling is whether (and how) to include chaotic orbits into the orbit library. The generalized Jeans’ theorem (e.g. Kandrup 1998) states that to satisfy CBE, the distribution function must be constant in every ‘well-connected’ region of phase space, whether it is an invariant torus defined by three integrals of motion for a regular orbit or a hypersurface of higher dimensionality for a stochastic orbit. The difficulty is that in many cases the distinction between regular and chaotic orbits is very blurry.

\footnote{Equation 44 in Thomas et al. (2004) has a sign error in the definition of entropy.}
and some weakly chaotic orbits retain a quasi-regular character for many periods – much longer than any time-scale of the model – before jumping into another part of their reachable region of phase space, thereby violating the assumption of time-invariance of each building block of the model. To combat this, Pfenniger (1984) adaptively increased integration time for those orbits (presumably chaotic ones) that exhibited substantial variation of time spent in each cell during the integration; however, for practical purposes there should be an upper limit to this time, and it may not guarantee an adequate phase space coverage of sticky orbits. Alternatively, van den Bosch et al. (2008) proposed to use ‘dithered’ orbits, starting a bunch of orbits with slightly different initial conditions and combining their density into one block to be used in the optimization routine, which increases the coverage of phase space available for a given (slightly perturbed) orbit. We have experimented with this approach but did not find it to be superior to just using equivalently larger number of separate orbits in the solution. Another interesting option is suggested by Siopis & Kandrup (2000), who found that adding a weak noise term to the equations of motion substantially increases the rate of chaotic diffusion and helps to reduce negative effects of stickiness. Apparently, this approach has not yet found an application in the context of Schwarzschild modelling. If the initial conditions for orbit library are sampled at a small number of energy levels (which we do not encourage), one could average the contributions of all chaotic orbits with a given energy into one ‘super-orbit’ which is then treated in modelling as any of the regular orbits (Merritt & Fridman 1996). The reason for this is that in 3D systems, all such orbits are parts of one interconnected (although not necessarily well-connected) region in phase space, so-called Arnold web (Lichtenberg & Lieberman 1992).

If necessary, one may enhance or reduce the relative fraction of chaotic orbits (or, in principle, any orbit family) by including an additional term in objective function, penalizing the use of such orbits (e.g. Siopis 1999). However, Vasiliev & Athanassoulas (2012) demonstrated that decreasing the number of chaotic orbits in Schwarzschild model does not enhance the stability of the corresponding N-body model.

In our implementation, we have a linear or quadratic optimization problem, defined by a set of linear equations (12) and (14) or (15), optionally with additional quadratic penalty terms (18) and/or penalties for given orbit families (e.g. tubes, chaotic orbits, etc.). This optimization problem is solved by one of the available solvers, using a unified interface. Presently, three options are implemented: the BIPME SOLVER (Mészáros 1999), the CVXOPT library9 (Andersen et al. 2012) or the GLPK package10 (only for linear problems). The typical time required to handle a model with \( N_o = \mathcal{O}(10^5) \), \( N_c = \mathcal{O}(10^3) \) is within few minutes on a typical workstation, much less than the time needed to build the orbit library.

A Schwarzschild model may be converted to an N-body model by sampling each orbit in the solution by a number of points proportional to its weight in the model. For a collisionless simulation, the sampling scheme may be improved by using unequal mass particles (Zemp et al. 2008; Zhang & Magorrian 2008), achieving better mass resolution and reducing two-body relaxation effects in the central parts of the model. The criteria for mass refinement may be based on energy, pericentre distance or any other orbit parameter; the option for mass refinement is implemented in the code but has not been much explored.

### 6 TESTS FOR ACCURACY OF POTENTIAL APPROXIMATIONS

In this section, we test the accuracy of three general-purpose potential approximations introduced above (basis-set, spline and N-body), from several different aspects. The first two potential expansions are smooth and should be able to represent analytically defined density profiles quite well, given a sufficient number of terms. All three are capable of approximating an arbitrary density model represented by a set of point masses. However, in this case there is a fundamental limit on the accuracy of approximation, set by discrete nature of underlying potential model; moreover, the optimal representation is achieved at some particular choice of parameters (order of expansion or N-body softening length), which needs to be determined: increasing the accuracy actually would only increase noise and not improve the approximation.

The accuracy of potential approximation is usually examined with the help of integral indicators such as integral square error or average square error (ISE or ASE; Merritt 1996; Athanassoula et al. 2000). We compare the accuracy of representation of not only force, but also potential and density, and replace the absolute error with the relative one, since it better reflects the concept of accuracy, and allows us to compare models with different underlying density profiles. Therefore, our measure of accuracy is defined by

\[
\text{ISE} = \int \rho(r) \left| 1 - \frac{F_{\text{approx}}(r)}{F_{\text{exact}}(r)} \right|^2 \, dr.
\] (19)

For all comparisons in this section, we use a set of five Dehnen models with different cusp slopes \( (\gamma = 0, 0.5, 1, 1.5, 2) \) and axis ratio of \( 1 : q : p = 1 : 0.8 : 0.5 \). The density profile is given by equation (3). For \( \gamma = 1 \) and \( p = q = 1 \) this reduces to the spherical Hernquist profile, and for \( \gamma = 2 \) – to the Jaffe profile.

We examine not only the integral error over the entire model, but also its variation with radius, to check which range of radii is well represented by the approximation, and what radii contribute the most to the integral error. For instance, a \( \gamma = 0 \) model may have large relative errors in force approximation at small radii, because the true force tends to zero for \( r \to 0 \) while the approximated one does not, but since the fraction of total mass at these small radii is negligible, it will not contribute to the total ISE. Nevertheless, one may argue that such a different behaviour of force at origin may substantially change the nature of orbits in the potential, for example, inducing more chaos as an orbit passes near the centre.

Therefore, the next step is to compare the orbits in the true and approximated potential, which is done as follows. A set of 5000 initial conditions, drawn uniformly from the corresponding density model at all radii (in the same way as for Schwarzschild modelling), is integrated in both the exact and approximate potentials for 100 dynamical times, and we compare several quantities on both per-orbit basis and on average. The properties of orbits to examine include the leading frequencies, LFCC diffusion rate \( \Delta \omega \), Lyapunov exponent \( \Lambda \) and the minimum squared angular momentum \( L_{\min}^2 \) (which distinguishes between centrophilic and

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9 For models with figure rotation, this chaotic region is spatially unbounded for the values of Jacobi constant (which replaces energy as the classical integral of motion) greater than the saddle point of the effective potential at Lagrange points. For this reason, the model of Hafner et al. (2000) did not have any irregular orbits beyond corotation radius.

10 http://cvxopt.org/

http://www.gnu.org/software/glpk/
centrophobic orbits). As explained in Section 4.2, $\Delta\omega$ is not a strictly defined quantity, and so we could not expect it to match perfectly at the level of individual orbits; however, the scatter in $\log(\Delta\omega_{\text{approx}}/\Delta\omega_{\text{exact}})$ should not be much larger than the expected 0.3–0.5 dex for a smooth potential, and there should not be any systematic shift towards higher (or lower) $\Delta\omega$. Similar considerations apply to $\Lambda$.

### 6.1 Approximating a smooth potential model with basis set or splines

First, we examine the accuracy of basis-set representation of a smooth potential. The basis sets used in the literature are completely in the sense that they may approximate any well-behaved potential–density pair, given a sufficient number of terms. However, to be effective with a small number of terms, their lowest order term should resemble the underlying profile as close as possible. For example, the Clutton-Brock basis set is not very suitable for representing density profiles with central cusps.

The Zhao (1996a) basis set implemented in SMILE is based on the two-power density profile with the inner and outer slopes equal to $-2 + 1/\alpha$ and $-3 - 1/\alpha$ correspondingly (Section A2), where the parameter $\alpha \geq 1/2$ may be chosen to give the highest possible accuracy for a given density profile. One might think that, for example, a better approximation to a model with finite central density is obtained with a value of $\alpha = 1/2$ (corresponding to the cored Plummer profile as the zero-order term), but it turns out that matching the cusp slope is not necessarily the best idea. More important is the range of radii in which the basis-set approximation is reasonably good, which depends both on the maximal order of expansion $n_{\text{rad}}$ and on $\alpha$: higher $\alpha$ gives a greater range because the break in density basis functions is more extended, and because their zeroes cover larger range of radii (Fig. 8).

For each value of $\gamma$, we constructed a series of BSE approximations with the number of radial terms $n_{\text{rad}}$ varying from 5 to 20 (in steps of 5) and the angular expansion order $l_{\text{max}}$ from 2 to 10 (even values only). Fig. 9, top panel, shows the integrated relative squared errors in potential, force and density approximations. A general trend is that increasing $n_{\text{rad}}$ always makes errors smaller, and increasing $l_{\text{max}}$ improves the approximation up to $l_{\text{max}} = 6$, after which there is no appreciable difference in most cases. This may be understood as ‘saturation’ of the approximation accuracy in the range of radii which contributes the most to the integral quantity.

An interesting result is that for weak-cusp models, increasing $\alpha$ above the value corresponding to the inner cusp slope actually makes the approximation much better. The reason is just a greater useful radial range of higher-$\alpha$ basis sets, as exemplified in Fig. 10, top, for $\gamma = 0$ and $\alpha = 1/2$ (Clutton-Brock) versus $\alpha = 1$ (Hernquist–Ostriker) basis sets. The latter clearly performs better at larger range of radii, and the improvement of error saturates at larger values of $l_{\text{max}}$ (for the former, there is no practical difference beyond $l_{\text{max}} = 4$). Similarly, even for $\gamma = 1$ Dehnen model which is traditionally represented with $\alpha = 1$ Hernquist–Ostriker basis set, the $\alpha = 2$ approximation actually works much better, both in the integral sense and in the range of radii for which relative error is small and improves with increasing $l_{\text{max}}$ (‘bottom of the trough’ depicted on the above figure). However, for even greater $\gamma$ it does not make sense to increase $\alpha$ beyond the value corresponding to the inner cusp slope; that is, for $\gamma = 1.5$ model $\alpha = 2$ is the best choice. The case $\gamma = 2$ is particularly difficult, since the potential diverges at origin, and formally $\alpha \rightarrow \infty$; we restrict this parameter to be $\leq 4$ for the reason that the magnitude of coefficients rapidly increases with $\alpha$ and $l$, and roundoff errors become intolerable.

In the case of Spline expansion, there is an additional freedom of choice of grid nodes, either to get a higher resolution (more frequently spaced nodes) at the intermediate radii where the bulk of integrated error comes from, or to achieve a better approximation at small or large radii. We find that exponentially spaced nodes are a good way to afford a large dynamic range in radius with relatively few nodes ($n_{\text{grid}} \sim 10–20$ for $r_{\text{rad}}/r_{\text{in}} \gtrsim 10^3$), so that the adjacent nodes differ by a factor of 1.5–3 in radius. Under these conditions, the accuracy at intermediate radii is mostly limited by order of angular expansion, at least for $n_{\text{grid}} \geq 10$ and $l_{\text{max}} \lesssim 8$; only the steepest cusp slopes require more than 20 nodes to achieve really small errors. Overall, the spline expansion outperforms BSE for comparable number of coefficients, both in terms of the radial range in which errors are small, and in integral characteristics such as ISE (Figs 9 and 10, bottom panels). In terms of computational efficiency of orbit integration, spline expansion is also faster than BSE and its performance is almost independent of the number of radial nodes (at least up to $n_{\text{grid}} = 40$). Both are substantially, by a factor of few, faster than the exact Dehnen potential (Fig. 11).

Finally, we compare the properties of orbits integrated in exact and approximated potentials, to address the question how much the relative errors in potential and especially force affect the dynamics. In particular, many of the approximate models have asymptotic behaviour of force at small radii which is different from the exact
model (in particular, BSE with the parameter $\alpha$ not matching the inner cusp slope). It is not obvious to which extent these deviations actually matter, without comparing the actual orbits. Even if properties of individual orbits do not strictly match between exact and approximate potentials, we want the ensemble of orbits to exhibit similar characteristics (e.g. distribution in $\Delta\omega$, $\Lambda$, number of centrophilic orbits, etc.).

These studies basically confirm the conclusions of the above discussion. For BSE approximations, almost any model with $n_{\text{rad}} \geq 10$ and $I_{\text{max}} \geq 6$ is close enough to the exact potential. For given $\gamma$, models with higher $\alpha$ are better approximations (that is, for $\gamma = 0$ the case $\alpha = 1/2$ performs much worse than any other model, and $\alpha = 1$ is already good; for $1/2 \leq \gamma \leq 3/2$ the models with $\alpha = 2$ are preferred). Comparing properties of individual orbits, we find that frequencies $\omega$, orbit shape (measured by diagonal values of the inertia tensor), and values of minimum squared angular momentum $L_{\text{min}}^2$ are recovered to within few percents, and log $\Delta\omega$ typically has scatter of $0.5 - 1$ (this is the only quantity which improves steadily with increasing $n_{\text{rad}}$). For the entire orbit ensembles, distribution of orbits in $\Delta\omega$ and in $\Lambda$ is very close to the one from exact models for all cases with $n_{\text{rad}} \geq 10$, $I_{\text{max}} \geq 4$ (with the above-mentioned exception of $\gamma = 0, \alpha = 1/2$ model). The fraction of centrophilic orbits is determined with $\sim 5$ per cent accuracy (and does not further improve with increasing the precision), and the number of orbits with $\Lambda = 0$ matches to within 1 per cent. For spline expansion, conclusions are similar; $n_{\text{grid}} = 20$ was sufficient for all models except $\gamma = 2$, for which 40 nodes did show improvement over 20; and there is no substantial change after $I_{\text{max}} = 6 - 8$.

Another test of the same kind is a study of orbital properties of BSE/Spline approximations of the Perfect Ellipsoid (Kuzmin 1956; de Zeeuw 1985), which is a fully integrable triaxial potential corresponding to the density profile $\rho(R) = (\pi^2 pq)^{-1}(1 + R^2)^{-2}$. Since all orbits in the exact potential should be regular, we may easily assess the quality of approximation by counting the number of chaotic orbits. Fig. 12 shows that indeed BSE approximations with higher $\alpha$ parameters are much better at representing the potential, despite that their asymptotic behaviour of force at origin is different from the exact potential.

Overall conclusion from this section is that BSE and spline expansions with a sufficient number of terms are good approximations to Dehnen models with all values of $\gamma \in [0, \ldots, 2]$. For BSE, the parameter $\alpha = 2$ gives best results for $1/2 \leq \gamma \leq 3/2$, with $\gamma = 0$ and 2 requiring $\alpha = 1$ and 4, correspondingly; $n_{\text{rad}} = 10 - 15$ is sufficient. The order of angular expansion $I_{\text{max}} = 6 - 8$ is enough for moderately flattened systems considered in this section, but may need to be increased for highly flattened, discy models.

Figure 9. Integrated square relative errors for potential (left), force (centre) and density (right) approximations of Dehnen models with basis-set (top) and Spline spherical-harmonic expansion (bottom). Different symbols/colours represent models with different cusp slopes and $\alpha$ parameters in BSE. Four groups of points correspond to varying the number of radial coefficients $n_{\text{rad}}$ from 5 to 20 for BSE, and radial grid points for spline interpolation coefficients $n_{\text{grid}}$ from 5 to 40; each group has 5 points for different number of angular terms varying from $I_{\text{max}} = 2$ to 10 (even numbers only). Top: BSE models for $\gamma = 0, 0.5$ and 1 are constructed using two different $\alpha$ parameters: first matches the inner cusp slope ($\alpha = 1/2 - \gamma$), second is a higher value of $\alpha$, which, as seen from the figure, performs substantially better (open versus filled triangles). It is also clear that increasing $I_{\text{max}}$ beyond 6 only makes a marginal improvement and only for some models, and $n_{\text{rad}} = 15$ is, in general, sufficiently accurate. (The reason for non-monotonic behaviour of density error with $I_{\text{max}}$ for the $\gamma = 2$ model is a large relative deviation at $r \gtrsim 100$; at smaller radii approximations with larger number of terms are more accurate). Bottom: Dehnen models with $\gamma \leq 1$ are already well represented by spline approximations with $n_{\text{grid}} = 10$ radial points and $I_{\text{max}} = 6$ angular terms; steeper cusps slopes require somewhat larger $n_{\text{grid}}$, with only $\gamma = 2$ model benefiting from increasing this number to 40 (because of logarithmic divergence of potential at origin it requires considerably smaller radius of the inner grid point). At large $n_{\text{grid}}$, the number of angular terms is actually the factor that limits accuracy. For almost all applications, $n_{\text{grid}} = 15 - 20$ and $I_{\text{max}} = 6 - 8$ will suffice and outperform BSE approximation.
Figure 10. Relative squared errors for potential (left), force (centre) and density (right) approximations of $\gamma = 0$ Dehnen model, as functions of radius. Top: BSE with $\alpha = 1/2$ (blue dotted line for $n_{rad} = 10$, blue dot–dashed for $n_{rad} = 20$, $l_{max} = 10$, purple dot–dashed for $n_{rad} = 20$, $l_{max} = 4$), and with $\alpha = 1$ (green dashed – $n_{rad} = 10$, $l_{max} = 10$, green solid – $n_{rad} = 20$, $l_{max} = 10$, red solid – $n_{rad} = 20$, $l_{max} = 4$). This panel illustrates that BSE with larger $\alpha$ have larger useful radial range, even if the inner cusp slope does not match that of the underlying model. Increasing the number of radial functions does extend the range of radii in which approximation works well (trough-like shape of the bottom curve at intermediate radii, $r \sim 10^{-1}$–$10^{-2}$); for a fixed $n_{rad}$, increasing the order of angular expansion improves approximation at these intermediate radii (compare solid curves for $l_{max} = 4$ and 10), but only until the ‘bottom of the trough’ is reached. At small or large radii increasing $l_{max}$ has no effect. It is the integral over these intermediate radii which mainly contributes to the ISE values of Fig. 9, but large relative errors at small or large radii may be hidden in that integral characteristic. Bottom: Spline approximations with $n_{grid} = 10$ (dashed) and $n_{grid} = 20$ (solid), top (red) is for $l_{max} = 4$, bottom (green) – for $l_{max} = 10$. Clearly this expansion performs much better overall than BSE, and continues to improve with increasing $l_{max}$ for given $n_{grid}$, saturating at smaller errors and in larger radial range.

Figure 11. Wall-clock integration time of a single orbit on a typical workstation (for 100 orbital periods), depending on the number of terms in BSE and spline approximations. Horizontal axis is the order of angular expansion, with the number of terms being $\frac{1}{2}(l_{max} + 1)(l_{max} + 2)$. Solid line is for Spline with $n_{grid} = 40$ (it almost does not depend on number of radial points), other lines from bottom to top are for BSE with $n_{rad} = 10$, 20 and 30 radial terms. For comparison, integration time for an exact triaxial $\gamma = 1$ Dehnen potential is ~1.5 sec/orb.

Figure 12. Left: Fraction of chaotic orbits in BSE and Spline approximations of the Perfect Ellipsoid potential (with $q = 0.8$, $p = 0.5$), detected by computing Lyapunov exponent over 100 dynamical times. Right: Mean value of FDR $\Delta \omega$. Three groups correspond to $n_{rad} = 10$, 15, 20 for BSE and $n_{grid} = 10$, 20, 40 for Spline; points in each group have $l_{max} = 4, 6, 8, 10$. The exact potential is integrable, therefore the lower is the number of orbits with $\Lambda > 0$ or the value of $\Delta \omega$, the better is the approximation. It is clear that while all approximations do perform better with increasing number of terms, some do it much faster: generally, $\alpha = 2$ BSE expansion wins the race even though its behaviour at origin is very different from the flat core of the Perfect Ellipsoid.

6.2 Basis-set and spline representation of a discrete particle set

A rather different case is when the coefficients of potential expansion are evaluated from a set of point masses, for example to study orbital properties of an $N$-body system. As is general for this kind of problems, the approximation error is composed of two terms. The bias is the deviation of approximated potential from the presumable ‘intrinsic’ smooth potential, which is the continuum limit of the $N$-body system, and was explored in the previous section: increasing number of terms never makes it worse, although may not improve substantially after a certain threshold is reached. The variance is the...
discreteness noise associated with finite number of particles, and it actually increases with the order of expansion. Therefore, a balance between these two terms is achieved at some optimal choice of the number of coefficients (e.g. Weinberg 1996). This conclusion is well known in the context of choice of optimal softening length in collisionless N-body simulations (Merritt 1996; Athanassoula et al. 2000; Dehnen 2001), and will also be reiterated in the following section in application to the tree-code potential. Here, we show that a similar effect arises in the smooth BSE and Spline representations of a discrete particle set.

An illustration of the effect of variance is provided by the following exercise. We generate several realizations of the same triaxial density profile, compute expansion coefficients for each one and calculate their average values and dispersions, comparing to the coefficients evaluated from the analytical density model. Fig. 13 demonstrates that all BSE coefficients with sufficiently high indices $n$, $l$, $m$ are dominated by noise. Somewhat surprising is the small number of ‘usable’ terms – just a few dozen even for an $N = 10^5$ particle model. The range of significant terms depends on the number of particles and on the details of density distribution and $\alpha$ parameter in BSE, but in general, angular terms beyond $l = 6$ are unreliable, with only a few first $n$ and $m$ terms at that value of $l$ are significant.

For Spline potential, the situation is similar, but instead of variation of coefficients with $n$, we follow their variation in radius for a given $l$, $m$. Noise limits the useful order of angular expansion especially at small radii, where the interior mass is represented by just a small number of points; for intermediate to larger radii the coefficients are reliable for a somewhat higher angular order (e.g. up to $l = 8, m \leq 2$ for $N = 10^5$ model). Fig. 14 shows the radial variation of the $l = 4$, $m = 4$ spherical-harmonic coefficient for a particular $10^5$ particle model, together with spline approximations with $n_{\text{grid}} = 10$ and as many as 40 points, compared to the coefficient from exact density profile. It is clear that in this case, increasing number of nodes may make the spline approximation match the radial dependence of this coefficient almost perfectly, but it turns out to fit mostly discreteness noise rather than true behaviour of this harmonic. While some regularization techniques may be applied to achieve balance between approximation accuracy and spline smoothness (e.g. Green & Silverman 1994), it may be easier just to keep the number of radial grid nodes small enough (10−15 in this case).

We explore the accuracy of approximation for the same five Dehnen profiles as in the previous section, but now initializing coefficients from $N = 10^5$ and $10^6$ particle realizations of corresponding models. Again we compare the ISE indicator and the range of radii for which the error is tolerable, as well as the orbital properties.

Not surprisingly, it turns out that the ISE in potential, force and density approximations cannot be reduced below a certain level, which is roughly $10^{-6}$, $10^{-4}$ and $10^{-2}$ correspondingly, for a $10^5$ particle model, and somewhat lower for $N = 10^6$ model. There is almost no improvement in integrated error after $l = 4$, $n = 10$ for $N = 10^5$. In terms of orbit properties, however, higher $l_{\text{max}}$ and $n_{\text{rad}}$ actually make things worse, in the sense that orbits become more irregular compared to the exact potential. If the number of coefficients is too low ($l_{\text{max}} = 2$ or $n_{\text{rad}} = 5$), the model is too far from the exact one, i.e. the difference in orbit properties is dominated by bias: there are fewer chaotic orbits and their properties are noticeably different from the exact potential. $n_{\text{rad}} = 10$, $l_{\text{max}} = 4$ seems to be the optimal choice for an $N = 10^5$ model and $n_{\text{rad}} = 15$, $l_{\text{max}} = 6$ is comparably good as the previous choice for an $N = 10^6$ case. Taking the expansion to higher orders introduces substantial bias in orbit properties: number of chaotic ($\Lambda > 0$) and centrophilic orbits is increased, and $\Delta \omega$ is noticeably shifted to higher values. Other properties of orbits are recovered quite well and are insensitive to the choice of expansion order: the mean-square difference in orbital frequencies is $\lesssim 5$ per cent, and the orbit shape is accurate within 2 per cent. Also worth mentioning is that here the results from ISE and orbit studies disagree: while the former indicator is still improving or at least constant with increasing $l_{\text{max}}$ and $n_{\text{rad}}$, the divergence in chaotic properties of orbits is noticeably increasing beyond the optimal choice of these parameters. Actually, the integral error is often accumulated at small radii, where the enclosed number of particles is small ($\mathcal{O}(10)$, with relative force and density errors $\gtrsim \mathcal{O}(10^{-1})$), but apparently the influence of these deviations on the gross dynamics is not very strong; more important are the high-frequency fluctuations at all radii, caused by higher harmonics whose amplitudes are pumped up by noise.

To summarize, representing a discrete N-body system by a smooth potential approximation requires a judicious choice of order of expansion, which balances the contribution of bias and variance to the approximation error. A rough estimate of how many terms should be retained can be obtained by evaluating expansion.

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**Figure 13.** Coefficients of an $\alpha = 1$ BSE approximation of a triaxial $\gamma = 0.5$ Dehnen model. Groups of 15 points represent radial coefficients at a given angular harmonic $l$, $m$; values of $l$ increase from 0 to 8 (even only) and $m$ from 0 to $l$ between groups. Points are the coefficients from analytical density profile, error bars are average over 10 realizations of $10^5$ points; error bar extending to zero means that value of this coefficient is noise dominated in the discrete realizations.

**Figure 14.** Spline approximation of the radial variation of $l = 4$, $m = 4$ spherical-harmonic coefficient (normalized to $l = 0$ coefficient) for a $10^5$ particle $\gamma = 1$ Dehnen model. The green solid line – coefficients evaluated at particle positions; red dash–dotted and blue dashed lines – spline approximations with $n_{\text{grid}} = 10$ and 40, correspondingly; grid nodes are marked with beads (square for $n_{\text{grid}} = 10$, round for 40). The purple dashed line – coefficients computed from exact density profile. It is clear that increasing the number of grid points may lead to an almost exact representation of $A_{\text{max}}(r)$ for a given N-body snapshot, but it does not converge to the ‘true’ coefficient for the exact underlying density model and actually deviates from it more as we increase $n_{\text{grid}}$. 
coefficients for several snapshots of the system, and retaining only those coefficients which do not have scatter greater than their magnitude. (Alternatively, one may wish to average over several snapshots to get a lower effective discreteness limit, but still the truncation order should be compatible with the data.) If multiple snapshots are not available, a rule of thumb would be to leave terms up to $l_{\text{max}} = 4(6)$ and $n_{\text{rad}} \sim 10-15$ for $N = 10^5 (10^6)$ particle model.

**6.3 Optimal softening for an N-body potential**

In this section, we explore the accuracy of the tree-code potential solver, which evaluates the potential directly from a set of $N$ point masses, without approximating it as a basis-set or spline spherical-harmonic expansion. There are two tunable parameters in this potential solver, the softening length $\epsilon$ and the opening angle $\theta$ of the tree-code.

Since our intention is to use the $N$-body snapshot as a discrete representation of a smooth mass distribution, it makes sense to introduce some sort of softening, for the same reasons as for the construction of smoothing splines from a discrete point mass set: the frozen $N$-body potential typically does not mean to represent this particular collection of particles, but rather samples a given smooth potential with a discrete set of points. Therefore, the error in this representation arises from two constituents: *variance* coming from fluctuations in particle positions, and *bias* from replacing Newtonian potential with a softened one. These two errors depend in the opposite way on $\epsilon$, so there exists a formally optimal value of softening length associated with a given mass distribution and $N$ (Merritt 1996; Athanassoula et al. 2000). Introduction of a softening length $\epsilon$ and modification of the Newtonian potential of particles at radii $\leq \epsilon$ is essentially equivalent to considering the gravitational field created by a smoothed density distribution (see Barnes 2012 for a discussion), therefore the problem of determining the optimal softening length is reduced to the problem of density estimation from a discrete point set. It is reasonable that the choice of softening length should depend on the local density (Dehnen 2001), although the optimal value of $\epsilon$ for the potential, acceleration and density estimation may differ by a factor of few. Collisionless $N$-body codes with spatially variable and time-adaptive $\epsilon$ are not widespread due to additional complications arising from the necessity of modifying the equations of motion to account for time-varying softening (Iannuzzi & Dolag 2011 describe one such implementation), but there is no reason why such spatially adaptive softening should not be used in a frozen $N$-body potential solver.

Another source of error in force computation in the tree-code comes from the tree algorithm itself, which replaces many particles with single tree cells. The additional error in force coming from this approximation depends on the cell opening angle $\theta$ and is generally rather low compared to the noise due to the discreteness of mass distribution (Hernquist, Hut & Makino 1993). Indeed, the tests below demonstrate that the error in the force and, to a lesser extent, potential approximation is rather insensitive to the accuracy of force calculation itself (opening angle $\theta$ and the use of quadrupole versus monopole terms), and is limited by discreteness noise. However, the error in calculation of an orbit does depend on the accuracy of tree-code (and on the integration timestep). To keep the errors in energy conservation at acceptable level, one should therefore appropriately choose the parameters of the tree-code.

The first set of tests is therefore dedicated to the determination of optimal softening parameters for a static approximation of a known potential by its discrete point mass representation. There are many choices for the functional form of softening. We define the softening kernel $\hat{\rho}(x)$ so that the density associated with a single particle of mass $m_i$ at location $r_i$ with a softening length $\epsilon_i$ is given by $m_i \epsilon_i^{-3} \hat{\rho}(|r-r_i|/\epsilon_i)$. The simplest form, widely used in $N$-body simulations, is the Plummer softening, given by $\hat{\rho}(x) \equiv (3/4\pi)(1 + x^2)^{-5/2}$; another choice, often employed in the non-parametric density estimation, is the Ferrers kernel: $\hat{\rho}(x) \equiv (15/8\pi)(1 - x^2)$ for $x < 1$ and zero for $x \geq 1$. The adaptive softening length for each particle of mass $m_i$ is taken to be

$$\epsilon_i = \epsilon (m_i/\overline{m})^{1/2} n_i^{-1/3},$$

where $\overline{m} \equiv M_{\text{total}}/N$ is the average particle mass, $n_i$ is the estimate of the local number density and $\epsilon$ is the global softening parameter. This local variation of softening length results in substantially lower bias than a single $\epsilon$ for all particles (Dehnen 2001). The tests below are done for equal values of $m_i$ and seek to determine the optimal value of $\epsilon$, which in this case is the coefficient of proportionality between the softening length and the local mean interparticle distance.

We consider the same five Dehnen models as in the previous sections. Each model is sampled with $10^5$ or $10^6$ equal point masses, and the relative square error in potential and force approximation is plotted as a function of radius and the integrated value (ISE) is computed, as a function of the softening parameter $\epsilon$. Fig. 15 shows that for the Plummer softening there is a well-defined minimum in ISE of force approximation at $\epsilon \sim 0.5-1$, while for the potential approximation the no-softening case gives formally the best result. This is because larger values of $\epsilon$ substantially increase the bias, since the Plummer softening modifies the potential at all radii and that dominates the approximation error. For the compact softening

![Figure 15](https://academic.oup.com/mnras/article-abstract/434/4/3174/960645)

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Orbit analysis and Schwarzschild modelling

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by guest on 30 July 2018
kernel, the dependence of ISE on $\varepsilon$ is much weaker; values of $\varepsilon \sim 1-2$ give near-optimal results in most cases, with steeper cusps requiring smaller $\varepsilon$. This is in agreement with Dehnen (2001) who demonstrated that Plummer softening is inferior to compact kernels for various reasons. There is almost no dependence on the accuracy of the tree-code algorithm for the force estimation, while for the potential estimation the inclusion of quadrupole moments and a smaller opening angle $\theta$ gives a better result.

Interestingly, but perhaps not surprisingly, approximation of an $N$-body snapshot with a BSE or Spline potential (Section 6.2) gives a substantially better result in terms of ISE, not even speaking of efficiency of orbit integration. This suggests that if one wants to analyse the orbital structure of an $N$-body system, such expansion methods (e.g. Hoffman et al. 2010) are preferred over a frozen $N$-body potential representation (e.g. Valluri et al. 2010). Not only they allow for a better accuracy of the force approximation, but also enable to compute Lyapunov exponents and even the FDR with a greater precision.

Secondly, to study how the granularity of the potential affects the chaotic properties of orbits, and to determine the parameters of tree-code that do not compromise the accuracy of orbit integration, we take an $N$-body realization of a triaxial Perfect Ellipsoid (with axis ratio $1:0.8:0.5$) which is known to be an integrable potential, and calculate $10^5$ orbits covering the entire range of energies. The chaoticity of orbits is measured by the mean value of FDR $\Delta \omega$, which should be as low as possible (Fig. 12 shows that it is around $10^{-5}$ for a BSE or Spline potential constructed from an analytic density profile). The force computed by the tree algorithm is not a continuous function of position, it has jumps when a cell opening criterion is triggered for any group of particles. As a result, energy of an orbit is not conserved exactly, no matter how small timesteps are taken, and this limits the accuracy of frequency determination.

Fig. 7 in Vasiliev & Athanassoula (2012) suggests that as long as the energy conservation error is kept below some threshold, it does not affect the values of FDR for orbits in the $N$-body potential. To estimate this threshold and the necessary parameters of tree-code, we run the orbit integration for various values of opening angle $\theta = 1, 0.7, 0.5, 0.4$ and softening parameters $\varepsilon$. We found that an opening angle $\theta \leq 0.5$ is necessary to achieve a reasonable energy conservation error of $\sim 10^{-8}$, which is well below the average value of $\Delta \omega = 10^{-3}(10^{-2.4})$ for an $N = 10^5(10^4)$ particle model. Better results are obtained with larger softening parameter (up to $\varepsilon = 3$). For comparison, orbits integrated in BSE and Spline potentials initialized from the same $N$-body snapshots had $\Delta \omega \sim 10^{-4}$, and the lowest value was obtained for $l_{\text{max}} = 4(6)$ angular terms for $N = 10^5(10^4)$, with little dependence on the number of radial terms $n_{\text{rad}}$. This is still higher than for an analytic density profile, but way better than for the discrete potential, and allows us to perform a meaningful detection of chaotic orbits using a conservative threshold of $\Delta \omega_{\text{ch}} = 10^{-3}$.

7 TESTS OF VARIANTS OF SCHWARZSCHILD MODELS

Here, we present comparison of the three variants of Schwarzschild modelling presented in Section 5.2. We use two test cases: the first one is a triaxial $\gamma = 1$ Dehnen model with a central BH, and the second one is a mildly cusped strongly triaxial model created by a cold collapse, which does not have an a priori known analytical density profile.

7.1 Triaxial Dehnen model with a central black hole

The first test case is the $\gamma = 1$ Dehnen model with axes ratio $x:y:z = 1:0.8:0.5$, with a total mass of unity, and a central point mass $M_\bullet = 0.01$ representing a galactic supermassive BH. It is well established (e.g. Valluri & Merritt 1998) that such a model with a relatively large central point mass has a large fraction of chaotic orbits at radii larger than a few BH influence radius $r_\bullet$ (defined as the radius containing stellar mass equal to $2M_\bullet$).

Earlier studies (Gerhard & Binney 1985; Merritt & Quinlan 1998) suggested that when $M_\bullet \gtrsim 2$ per cent of the total mass of the galaxy, its effect on the centrophilic orbits is strong enough to destroy triaxiality in just a few crossing times. Subsequent works, however, presented examples of a long-lived triaxial models with comparably large central masses: for instance, Holley-Bockelman et al. (2002) considered a moderately triaxial $\gamma = 1$ Dehnen model with $M_\bullet = 0.01$ in which the loss of triaxiality was confined to the inner 2 per cent of particles, while Kalapotharakos, Voglis & Contopoulos (2004) found that such a BH mass could drive a model towards axisymmetry in a time comparable to Hubble time, and Poon & Merritt (2004) demonstrated that truncated power-law models of inner parts of a galaxy (enclosing total mass of a few times $M_\bullet$) can remain triaxial even with a large fraction ($\gtrsim 50$ per cent) of chaotic orbits.

The goal of this test case is not to present a comprehensive study of evolution of triaxial cuspy models with BHs, but to show that at least in some cases the Schwarzschild method can be used to construct a relatively stable model. We use the three variants of Schwarzschild modelling techniques considered above (classical, basis-set and Spline spherical-harmonic expansion, hereafter labelled as C, B and S), with $N_r = 30$ radial shells or radial basis functions and 27 (28) angular constraints for C (B, S) models, giving $N_c \approx 800$ total constraints for $N_r = 4 \times 10^2$ orbits. All three models were solved with quadratic optimization routine and were required to keep velocity isotropy at all radii. The fraction of chaotic orbits was around 30 per cent. Then an $N = 10^5$ $N$-body model was created from each Schwarzschild model and evolved for $T = 100$ time units with the direct, hardware-accelerated $N$-body integration code $\phi{\text{Grape}\text{Pech}}$ (Hafst et al. 2008), which uses algorithmic regularization to improve the accuracy of integration of trajectories that come close to the BH. The softening length was set to zero and the accuracy parameter of the Hermite integrator was set to $\eta = 0.01$; total energy was conserved to a relative accuracy better than $10^{-4}$.

To check how close the models are to equilibrium, we used a number of indicators. First the density profiles of the $N$-body models were compared to that of the analytical mass profile used to create Schwarzschild model. Fig. 16, top panel, shows the ratio of densities of the $N$-body snapshots over the expected density, which is quite close to the expected unity. The three variants performed similarly well over most of the radial range, while there are some minor differences at small and large radii. Next, we compute the rate of energy diffusion as a function of particle energy, which is defined as the slope of the mean-square relative change of energy ($\Delta E/E$) as a function of time and compare it to the theoretically predicted diffusion coefficients (see Theuns 1996 for another example of such comparison). If the model is in a stable equilibrium, the particle energy changes only due to two-body relaxation, with a rate that can be computed using standard expressions for diffusion coefficients if one knows the distribution function (e.g. Merritt 2013, equations 5.55, 5.125). The only adjustable parameter in this computation is the value of Coulomb logarithm, and the usual practice is to take it to be roughly $\ln N$ for the entire model, or $\ln M_\bullet/m_\bullet$ for the
Three variants of Schwarzschild models for a triaxial at the radius of influence and particle model (in which $\Lambda_1 \times \Lambda_2 / \Lambda_z / 10$ ($2N_E = \\gamma_0.08$, independent of radius, so that the $\sim \ln \rho (0.01, \text{unit radius and unit mass, with } \gamma_0.01$. Top panel: density normalized to the expected value, as a function of radius. Shaded region shows typical uncertainty region for a random realization of a $10^5$ particle model (in which particle positions are distributed according to the analytical profile). Bottom panel: energy diffusion rate (relative mean-square change of particle energy per unit time, as a function of initial energy). Solid line shows theoretically computed value, points with error bars are the results from the simulation. Particles are binned into 100 bins of unequal size, with the most bound bins each containing $\sim 10$ particles; error bars represent the uncertainties in the estimate of the slope of energy growth with time. Different colours are for the three models, but there is no systematic difference between them. A spherically symmetric model with the same azimuthally averaged density profile, created by Eddington inversion formula, is also plotted for comparison; it demonstrates the same energy diffusion rate as the triaxial Schwarzschild models.

region around the BH; we interpolated between these asymptotic regimes by taking $\ln \Lambda = \ln \rho (M_e + M(E))/m_e$, where $M(E)$ is the mass in stars with binding energies higher than $E$. Fig. 16, bottom panel, demonstrates that the energy diffusion rate is again very similar between different variants and close to the theoretical prediction (which is a non-trivial fact: in an earlier, somewhat buggy implementation one of the models displayed substantially larger initial diffusion rate before settling into a new equilibrium, while looking reasonable in other aspects). This plot also shows that the relaxation time, computed as the inverse of relative energy diffusion rate, is a few times longer than the integration time of the runs, therefore we do not expect the two-body effects to play a substantial role, at least in the initial evolution. For comparison, the dynamical time is $\sim 10^{-1}$ at the radius of influence and $\sim 1$ at the half-mass radius.

Finally, most interesting is the evolution of model shapes (axis ratios). We used an iterative procedure (see Zemp et al. 2011 for an extended discussion) to determine the axes of inertia tensor of particles within a certain ellipsoidal radius, with the scaling of this ellipsoid’s axes being iteratively updated from the inertia tensor until both converge to within a defined tolerance. Plotted in Fig. 17 are the axis ratios ($y/x$ and $z/x$) as functions of time, for radii containing 2, 10 and 50 per cent of total mass (not including the BH). Here again the three variants performed almost identically. Not surprisingly, the evolution was strongest in the inner parts, although even there the triaxiality remained substantial. A more detailed exploration of the effects of resolution, initial shape and other parameters is left for a future study.

### 7.2 Cold collapse model

Instead of creating models for a known, analytical mass distribution such as the Dehnen profile, we use a triaxial $N$-body model obtained by cold collapse as the target density profile. Namely, we take $N = 0.5 \times 10^8$ equal-mass particles distributed according to $\rho_{\text{in}}(r) = (2\pi r)^{-1}$ in the sphere of unit radius and unit mass, with velocities assigned from isotropic Gaussian distribution with 1D velocity dispersion $\sigma_{1D} \approx 0.08$, independent of radius, so that the initial virial ratio of the system is $3 \times 10^{-2}$. The collapse of this sphere results in a strongly triaxial system due to the development of radial-orbit instability (Polyachenko & Shukhman 1981; Merritt & Aguilar 1985); the density in the inner part increases by a factor of few while retaining the slope, and axis ratios vary with radius, but roughly equal to $1 : 0.6 : 0.4$. Such cold collapse models are rather well studied (e.g. Aguilar & Merritt 1990; Cannizzo & Hollister 1992; Boily & Athanassoula 2006); a similar approach was used by Voglis et al. (2002) and Muzzio, Carpintero & Wachlin (2005) for the same goal of creating triaxial models in equilibrium, although the properties of these models are related to the initial conditions in a quite complicated and not always predictable way. Unlike these
studies, we follow the collapse and subsequent relaxation of the model by a less approximate N-body code.

First, we use the direct, hardware-accelerated N-body code \( \PhiGRAPES \) (Hafst et al. 2007) to follow the initial collapse and formation of triaxial bar up to \( t = 10 \) (the peak of the density and depth of the central potential is reached around \( t = 1 \)); at this first stage, we set a very small softening length \( \epsilon = 10^{-4} \) (to prevent the formation of binaries), which is nevertheless smaller than the mean interparticle separation at the peak of collapse. The energy is conserved to \( 10^{-4} \) during this first stage. Then we followed the evolution for another 100 time units by the efficient tree-code GYRFALCONS (Dehnen 2000), with \( \epsilon = 10^{-3} \), to create a well-mixed system (the traces of initial cold streams in phase space disappeared); the energy was conserved to the relative accuracy of \( 3 \times 10^{-4} \). The density profile did not change appreciably at this stage, apart from the expansion of outermost layers composed of unbound particles and some flattening at small radii. This flattening is not unexpected, since the relaxation time in the inner parts of the model is quite short: the locally evaluated \( T_{\text{rel}} = 0.34 \epsilon^2 \sigma^2 / (m_\star \rho(r) \ln \Lambda) \) (Binney & Tremaine 2008) is shorter than 100 time units for \( r < 0.04 \), while the dynamical time at this radius is \( \sim 0.1 \). The density profile can be roughly described by a \( \gamma = 0 \) Dehnen model with the scale radius \( r_0 = 0.06 \), but with moderate deviations and a non-constant axis ratio. Our chosen value for the softening length is just slightly larger than the optimal value for a \( \gamma = 0 \) Dehnen sphere, estimated in Athanassoula et al. (2000) to be \( \sim 0.01 \) at this number of particles and for the scale radius of unity. At the end of the second stage, we eliminated the unbound particles (which comprised 12 per cent of mass) and rotated the snapshot to be aligned with principal axes. In addition, the model demonstrated a slow figure rotation, similar to what has been found in analogous experiments (e.g. Aquilano et al. 2007; Muzzio, Navone & Zorzi 2009), which we neutralized by flipping the positions and velocities of a fraction of particles about one or more principal planes in such a way as to make the total linear and angular momentum as small as possible. Thus, we obtained the initial model for the third stage, which is expected to be an equilibrium configuration.

A stacked-up combination of last five snapshots from the second stage, taken at moments of time from 96 to 100 and joined to decrease shot noise, served as the input to the potential initialization (BSE and Spline potential expansions) used to generate three variants of Schwarzschild models (C, B and S), each having 10\(^5\) orbits and \( \sim 500 \) constraints. (Models C and S used the Spline potential expansion and model B used the BSE potential with the shape parameter \( \alpha = 1 \), i.e. the Hernquist-Ostriker basis set). The orbital structure of these models is quite rich, with roughly half of orbits being tubes (short- and long-axis tubes in comparable quantities) and the rest are boxes, of which about a third belong to various resonant families. Most of box orbits are chaotic while most tubes are not; the percentage of chaotic orbits depends on the number of terms in potential expansion (adding more high-frequency components makes more orbits chaotic), but in general about a half of all orbits are chaotic.

The original model, composed of \( 4.4 \times 10^5 \) particles, and the N-body models generated from each variant of Schwarzschild model with the same number of particles, were evolved for 200 time units using GYRFALCON with the same softening length as for the second stage. Fig. 18 shows that by the end of simulation, all three variants of Schwarzschild model have become somewhat less triaxial; the evolution of axis ratios was virtually the same for all models and for the original collapse simulation. Therefore, we have confirmed the ability of the Schwarzschild method to create strongly triaxial models which are as stable as the models created with other methods (in this example, cold collapse), which was recently called in question by Zorzi & Muzzio (2012).

8 CONCLUSIONS

We reviewed the methods for studying non-spherical galactic models, analysing properties and regularity of orbits and the overall structure of phase space in a given potential, and constructing self-consistent equilibrium models by the Schwarzschild method.

We developed a new software named SMILE, which is intended to be a convenient tool for performing orbit analysis and Schwarzschild modelling for a number of standard potential models, as well as for an arbitrary density profile (including a discrete N-body model) approximated by a flexible potential expansion. The new and improved methods implemented in SMILE include:

(i) two general-purpose potential expansions for arbitrary density models, in which the angular dependence of density and potential is decomposed into spherical harmonics and the radial dependence of the expansion coefficients is represented either as a sum of basis functions (using Zhao (1996a) basis set, which is a generalization of the commonly employed Hernquist & Ostriker (1992) set), or as a spline. Another variant of potential representation is via a set of fixed point masses, using a tree-code approach and a spatially adaptive softening length.

(ii) An improved method of chaos detection via Lyapunov exponent \( \Lambda \), which can distinguish between orbits having \( \Lambda > 0 \) and those that did not demonstrate any chaotic behaviour over the integration time.

(iii) The method of generating initial conditions for a Schwarzschild model, based on sampling of an isotropic distribution.
function of an equivalent spherical model. It eliminates some artefacts that appear in the traditional method of assigning initial conditions on a regular grid, and allows us to sample the phase space more uniformly. As a by-product, these spherical mass models may be used to efficiently analyse dynamical properties of arbitrary density profiles, including the ones from N-body snapshots.

(iv) Two new variants of Schwarzschild models, in which density of the target model and of the orbits is represented by one of the potential expansions mentioned above.

The program comes in two versions: an integrated graphical interface allows us to connect various related tasks and instantly analyse the results (e.g. select an orbit from the frequency map or the Poincaré section, visualize its shape and spectrum, compare orbit distributions in different models, etc.), while the console version is more suitable for performing scriptable tasks. It is written in C++ and available for download at http://td.lpi.ru/~eugvas/smile/. Modular design allows us to use parts of the code (e.g. orbit integration or analysis) in other programs, for example for analysing orbits from an N-body simulation; also provided is integration with NEMO N-body simulation framework (Teuben 1995).

We have performed accuracy tests of the potential expansions and found that they may be efficiently used for approximating various smooth density profiles with sufficient accuracy, using a rather modest number of terms; the Spline potential is generally more flexible and efficient. For a discrete mass set, the accuracy of these smooth representations is limited by the shot noise; even if the latter uses an appropriately chosen adaptive softening. We also tested our implementations of the classic and two new variants of Schwarzschild models on two test cases, and found them to perform with sufficient efficiency and fully capable of creating triaxial models in dynamic equilibrium.

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APPENDIX A: POTENTIAL EXPANSIONS

Two general-purpose potential representations used in SMILE are the basis-set and Spline expansions. They share the method of dealing with angular dependence of density and potential by expanding it in terms of spherical harmonics $Y^m_l(\theta, \phi)$, but differ in the description of radial variation of expansion coefficients. In addition, the same angular expansion may also be used as an approximation to the scale-free (single power-law) density profile. The angular dependence of density and potential is given by

$$\rho(r, \theta, \phi) = \sum_{l=0}^{l_{\text{max}}} \sum_{m=-l}^{l} A_{lm}(r) \sqrt{4\pi} P_l^m(\cos \theta) \text{trig} m \phi,$$

$$\Phi(r, \theta, \phi) = \sum_{l=0}^{l_{\text{max}}} \sum_{m=-l}^{l} C_{lm}(r) \sqrt{4\pi} P_l^m(\cos \theta) \text{trig} m \phi,$$

where $\text{trig} m \phi \equiv \begin{cases} 1, & m = 0 \\ \sqrt{2} \cos m \phi, & m > 0 \\ \sqrt{2} \sin |m| \phi, & m < 0. \end{cases}$

Here, we expanded the spherical harmonics: $Y_l^m(\theta, \phi) = (-1)^m P_l^m(\cos \theta) e^{im\phi}$, where $P_l^m(x)$ are normalized associated Legendre polynomials, and combined the exponents to obtain real coefficients with sines and cosines. The factors of $\sqrt{2}$ for $m \neq 0$ are introduced to keep $\sum_{m=-l}^{l} A_{lm}^2$ rotationally invariant. We introduced the convention that $m < 0$ corresponds to odd (sine) part of expansion. For triaxial systems only terms with non-negative even $l$ and $m$ are non-zero; however, we keep the more general form of expansion, while using only the necessary number of non-zero terms in the actual computations (e.g. only $l = 0$ terms in the axisymmetric case).

A1 BSE for a scale-free potential

To start with, consider a simpler case of a scale-free potential, where $A_{lm}(r) = A_{lm} r^{-\gamma}$, $C_{lm}(r) = C_{lm} r^{\gamma}$. The expansion coefficients for density and potential are related by the formula

$$C_{lm} = \frac{4\pi}{l(l+3-\gamma)(2l-\gamma)} A_{lm}.$$

We note that this representation is much more accurate for a given $l_{\text{max}}$ than cosine expansion used in Terzić (2002).
A2 BSE for a generic triaxial potential

Next we come to the representation of generic 3D potential–density pair in terms of basis function series, defining

\[ A_{nm}(r) = \sum_{n=0}^{n_{rad}} A_{nlm} \rho_n(r), \quad C_{nm}(r) = \sum_{n=0}^{n_{rad}} A_{nlm} \mathcal{C}_n(r). \]

We use the \( \alpha \)-model basis set described by Zhao (1996a). In the zeroth order it represents the following potential–density pair:

\[ \Phi_{00} = -\frac{1}{1 + r^{1/\alpha}}, \quad \rho_{00} = \frac{1 + \alpha}{4\pi \alpha} r^{2 - 1/\alpha} \left( 1 + r^{1/\alpha} \right)^{2 + \alpha}. \]

Setting \( \alpha = 1/2 \) gives the Plummer potential and associated Clutton-Brock (1973) basis set; value of \( \alpha = 1 \) corresponds to the Hernquist (1990) model and the Hernquist & Ostriker (1992) basis set. The higher order terms are given by

\[ \Phi_{nl} = -\frac{r^l}{(1 + r^{1/\alpha})^{2l + 1 + \alpha} G_{n}^{(1\alpha)}(\xi)}, \]

\[ \rho_{nl} = \frac{K_{nl}}{2\pi} \frac{r^{l-2 + 1/\alpha}}{(1 + r^{1/\alpha})^{2l+1+2\alpha}} G_{n}^{(1\alpha)}(\xi), \]

\[ K_{nl} = \frac{4(n + w)^2 - 1}{8\alpha^2}, \quad w = (2l + 1)\alpha + 1/2, \quad \xi = \frac{r^{1/\alpha} - 1}{r^{1/\alpha} + 1}, \]

where \( G_{n}^{(1\alpha)}(\xi) \) are Gegenbauer (ultraspherical) polynomials.

A3 Spline spherical-harmonic expansion

Another option for approximating an arbitrary potential is to represent its angular part in spherical harmonics, but retain an explicit radial dependence of the expansion coefficients \( C_{nm}(r) \), with forces and density given by derivatives of these functions. Accordingly, \( C_{nm}(r) \) is represented as a cubic spline interpolating between a grid of nodes. A straightforward approach faces some difficulties: it is hard to accommodate a large dynamic range in radii and, more importantly, ensure that the second derivative of \( C_{nm}(r) \) is reasonably close to that of the underlying model.

We use the following modification to the base scheme: first, the lowest order term \( C_{00} \) is represented as a double logarithmic function:

\[ \tilde{C}_{00}(\xi \equiv \log r) \equiv -\log(C_{00}(0)^{-1} - C_{00}(r)^{-1}). \]

The advantage of such scaling is that it can accommodate a power-law behaviour at small \( r \): if \( C_{00}(r) \approx C_{00}(0) \times (1 -Kr^{\gamma}) \), then \( \tilde{C}_{00}(\xi) \approx \log(-C_{00}(0)/K) - (2 - \gamma)\xi \) as \( \xi \to -\infty \). This modified function is approximated by a cubic spline in \( \log r \) and extrapolated at both small and large \( r \) for \( \log r \to -\infty \), we use the linear extrapolation described above, which corresponds to power-law density behaviour at \( r \to 0 \). For \( r \to \infty \) the extrapolation is mimicking a power-law density profile \( \rho \propto r^{-\sigma} \) with \( \gamma_{out} > 3 \); in this case

\[ C_{00}(r) \approx C_{00}(r_{out}) (r_{out}/r)^{\gamma_{out}/3} \left[ 1 - \mu [(r_{out}/r)^{\gamma_{out}/3} - 1] \right], \]

where \( \gamma_{out} \) is calculated from the last three grid nodes and \( \mu - \) from the last two.

Higher order coefficients cannot be represented in such a logarithmic way because they need not have the same sign over all \( r \); instead, they are normalized by \( C_{00}(r) \) and interpolated in \( \xi \equiv \log(1 + r) \): the spline is constructed for

\[ \tilde{C}_{lm}(\xi) \equiv C_{lm}(r)/C_{00}(r). \]

The evaluation of ‘real’ (not tilded) coefficients and their derivatives is just a number of simple algebraic transformations of the spline values and derivatives up to second order, which are trivially obtained for a cubic spline. Inside the first and beyond the last grid nodes, coefficients are extrapolated as power-law functions of radius.

The radial grid for representing the coefficients is taken to be exponentially spaced: \( r_i = r_1 \times [\exp(Zk) - 1]/[\exp(Z) - 1] \), where \( r_1 \) is the radius of the first node, and \( Z \) is assigned a value which gives some pre-defined radius \( r_G \) for the last node \( n_{grid} \). Typically, it makes sense that \( r_1 \) encloses some small fraction of mass (\( \delta \sim 10^{-3} \) or less), and \( r_G \) encloses \( 1 - \delta \) of total mass; logarithmic scaling makes a very good approximation with as small as \( n_{grid} = 10 - 20 \) nodes even for \( r_G/r_1 \gtrsim 10^4 \).

If the potential is initialized from a set of \( n_{pt} \) discrete point masses, the splines are constructed using an adjustable amount of smoothing, according to the following procedure. First, the inner and outer grid radii \( r_i, r_G \) are assigned so that there are only a few particles inside \( r_1 \) or beyond \( r_G \), and the radial grid is constructed using exponentially spaced nodes. Then the expansion coefficients \( C_{lm}(r) \) are computed at each particle’s radius \( r_i, i = 1, \ldots, n_{pt} \) and the penalized least-square fitting method is used to construct a smoothing spline \( \tilde{C}_{lm}(x) \) which minimizes the following functional:

\[ \mathcal{L} = \sum_{i=1}^{n_{pt}} \left( C_{lm,i} - \tilde{C}_{lm}(x_i) \right)^2 + \lambda \int \left\{ \tilde{C}_{lm}''(x) \right\}^2 dx . \]

Here, \( x \) are the scaled radial variables (\( \xi \) or \( \zeta \)), and \( \lambda \) is the smoothing parameter. As discussed in Merritt & Tremblay (1994), increasing this parameter to infinity is equivalent to fitting the data with a pre-specified functional form of regression; in general, unlike kernel-based smoothing approaches, minimization of \( \mathcal{L} \) tends to produce power-law regressions, in this case the \( l = 0 \) term corresponds to the Hernquist profile (for comparison, smoothing the mass profile as described in Section 5.1 would produce a two-power-law model \( \rho(r) \propto r^{-\gamma} (r_0 + r^2)^{-\gamma} \) in the limit of infinite \( \lambda \)). The value of the smoothing parameter \( \lambda \) may be chosen at will; while there are standard techniques such as generalized cross-validation method (Wahba 1990), they tend to produce very little smoothing if \( n_{pt} \gg n_{grid} \). A practical recipe implemented in SMILE uses the Akaike information criterion (AIC, e.g. Burnham & Anderson 2002) in such a way that the difference \( \Delta \text{AIC} \) between the fit with no smoothing and the fit taken to be ‘optimal’ does not exceed some pre-defined (adjustable) value. Typically, this results in little extra smoothing for low \( l, m \) where coefficients do indeed contain useful data, and a substantial reduction of noise for higher \( l, m \).

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