MODEST-1: Integrating Stellar Evolution and Stellar Dynamics

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We summarize the main results from MODEST-1, the first workshop on MOdeling DEnse STellar systems. Our goal is to go beyond traditional population synthesis models, by introducing dynamical interactions between single stars, binaries, and multiple systems. The challenge is to define and develop a software framework to enable us to combine in one simulation existing computer codes in stellar evolution, stellar dynamics, and stellar hydrodynamics. With this objective, the workshop brought together experts in these three fields, as well as other interested astrophysicists and computer scientists. We report here our main conclusions, questions and suggestions for further steps toward integrating stellar evolution and stellar (hydro)dynamics.

1. Introduction

Population synthesis models have been used successfully in comparisons with observations of the global properties of stars, star clusters, and galaxies. The simplest models are constructed from a weighted sum of individual stellar evolution tracks, while more detailed models incorporate some additional information about binary stellar evolution.

For some stellar environments such a synthesis approach is perfectly adequate, and there the main challenge is to deal with the considerable complexities of binary star evolution. However, the situation is very different for the class of dense stellar systems, defined as environments in which a typical star has a significant chance to interact and possibly collide with another star during its lifetime. In such an environment stars of different ages can exchange mass, disrupt each other

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or merge, and their merger products can get involved in similar interactions; binary stars can encounter single stars as well as other binaries, where one or more of the stars may already be a merger product; and so on. Examples of dense stellar systems are star-forming regions and the dense cores of open and globular clusters, as well as galactic nuclei.

It is clear that the possibilities are almost endless. While population synthesis based on single-star evolution can easily be exhaustive, and synthesis based on a mixture of single stars and binaries can at least aim to be reasonably complete, there is no way that one can anticipate and tabulate all possible multiple-star interactions in dense stellar systems. Detailed attempts at population synthesis for such systems by necessity have to be dynamical, taking into account the particular ways that stars encounter one another in a given simulation.

During the last few years, several dynamical population synthesis studies have appeared (cf. Portegies Zwart et al. 2001, Hurley et al. 2001). In these studies, the dynamics of a dense stellar system is modeled through direct N-body integration, while the stellar evolution is modeled through fitting formulae that have been obtained from large numbers of individual stellar evolution tracks. Binary stellar evolution is modeled through the use of semi-analytic and heuristic recipes (Hurley et al. 2002).

Astrophysically these results are novel and exciting, but their reliability is not so easy to assess. Validation is a core issue here, requiring not only detailed internal checks but also comparison between different codes run by different groups. This question was discussed at some length last year at IAU Symposium 208 in Tokyo, resulting in the specification of a well defined set of initial cluster and stellar parameters (Heggie 2002). Given the fact that the necessary codes are rather complex, requiring years of development, so far few groups have been able to confront this new challenge. This stands in contrast to the first collaborative experiment (Heggie et al. 1998), which was confined to stellar dynamics (without stellar evolution), and attracted "entries" from about 10 groups. We hope that our new MODEST initiative will stimulate more groups to engage also in the friendly competition of the second collaborative experiment.

Further improvement to the more comprehensive simulations referred to above will require the use of "live" stellar evolution models before too long, in order to deal with the unusual types of new stars that can be formed by mergers in dense stellar systems. However, the challenges of coupling existing stellar evolution codes and stellar dynamics codes are quite daunting. The first workshop specifically organized to address these challenges was held during July 17-21, 2002 at the American Museum of Natural History in New York City. The workshop brought together a group of experts in stellar evolution, stellar dynamics, stellar hydrodynamics and other fields of astrophysics, as well as computer scientists.

Originally, the workshop was announced to a small group of people who were known to work on the interface of dynamics and evolution, under the title "Integrating Stellar Evolution and Stellar Dynamics". We originally expected to see a handful of participants for an informal roundtable discussion. The fact that instead 34 attendants convened is a clear sign of the timeliness of the meeting, and the desirability to form a concerted effort to bridge the gap between the stellar evolution and dynamics communities.

This paper offers a summary of the week-long series of discussions held during the workshop, distilled by the organizers (Piet Hut and Mike Shara) and eight of the participants representing a cross section of expertise available during the meeting. In addition, we have created a web site\(^2\) where the name ‘modest’ reflects our renaming of the meeting during the last day to MODEST-1, the first workshop on MOdeling DEnse STellar systems. We plan to hold biannual follow-up meetings, MODEST-2 in Amsterdam in December 2002, and MODEST-3 in Australia in July 2003. In addition, we have started an email list to facilitate ongoing discussions about technical details of dynamical population synthesis simulations. Further information can be found on our web site.

\(^{2}\)http://www.manybody.org/modest.html
As a summary of our workshop, this paper contains the input of all of the participants, which are listed below under the acknowledgments. While many of the authors have contributed to various sections, each section has one or two main authors, as follows. §1 and §4 were written by Piet Hut, §2 by Michael Shara, §3 and §6 by Piet Hut and Jun Makino, §5 by Onno Pols and Ronald Webbink, §7 and §8 by James Lombardi, §9 by Sverre Aarseth and Ralf Klessen, §10 by Steve McMillan and Peter Teuben, and §11 by Steve McMillan.

In order to make the discussion concrete we have provided specific code fragments in §6 and §8 below. We see this paper as the start of a discussion that will ultimately result in the definition of clear standards for interfaces between stellar dynamics, evolution, and hydrodynamics. However, the current fragments are for illustration only, and are not necessarily intended to become part of any future standard.

2. Predictions

The successful marriage of $N$-body simulations with increasingly sophisticated stellar evolution codes of all flavors will yield progeny whose genetic characteristics should be designed now, to avoid petabytes of untestable output.

Essential ingredients of science are predictions and testability. Of course, we all look forward to detailed models of star clusters with self-consistent stellar evolution spanning aeons of time. But I want to emphasize how critical it is to generate those models with enough genetic markers to allow observers to tell us if our models have anything at all to do with physical reality.

A poster child for this kind of approach is the important paper by di Stefano and Rappaport (1994), where directly testable predictions of the cataclysmic binaries in a few selected globular clusters were made. Such predictions are dangerous for the egos of theorists (it’s not fun when observers find many orders of magnitude more or less than what you predicted) but it’s essential to the health of our science.

Modelers of star clusters are confronted with datasets rich in genetic markers from HST, Chandra and other observatories. Detailed sequences of blue stragglers, white dwarfs, X-ray binaries, millisecond pulsars and “missing” red giants are now available for significant numbers of globular clusters. While these often represent less than ten percent of the cluster (both in terms of numbers and in terms of mass), they must be reproduced in the correct numbers and positions in clusters if we are to have any confidence in the coming generations of MODEST models.

A slightly more subtle, but no less important set of predictions that should be made by combined $N$-body and stellar evolution codes concerns the lineages of tracer stars. It is just as informative to know how each blue straggler in a cluster got that way as it is to know how many blue stragglers are predicted in a cluster. The “synthetic history” of each star should not be taken literally because of the chaotic nature of the individual particle trajectories. However, the cumulative, statistical histories of entire classes of stars are important because these make testable predictions.

A concrete example comes from recent simulations of Shara and Hurley (2002) of M67-like star clusters. The life cycle of every white dwarf binary in every simulation was followed in detail, focusing on the systems that eventually merge. The key result is that the white dwarf merger rate is enhanced, relative to the field, by an order of magnitude. The life story of any particular binary white dwarf in this simulation isn’t important. However, the history of the entire class of objects is very important: it directly predicts that SNIa may be preferentially produced in star clusters. This is observable and hence testable.

In summary, theorists should consider providing not just the numbers, lifetimes, luminosities, colors and spatial distributions of every class of “tracer” star in a cluster. These will be indispensable in directly matching observed clusters to simulated clusters. But deeper insights into the evolution of star clusters can be gained by retaining statistical information about the histories of stellar populations from the $N$-body with stellar evolution simulations.
3. A MODEST Approach

3.1. Divide and Conquer

Conceptually, it would be easiest to start from scratch in order to model the gravitational, hydrodynamic, and internal interactions between stars. In such an approach, one could choose a particular computer language and style of programming, define the appropriate data structures and abstraction barriers, and write the various parts of the program accordingly. And indeed, such a project might be feasible, but would probably take a team of people years to accomplish. For the near future it makes more sense to work with existing computer codes that already can handle the dynamics or evolution or hydrodynamics that are needed to model dense stellar systems.

For one thing, many of these three types of programs already incorporates tens to hundreds of person-years of collective experience, and it will be far from easy to codify and reproduce that expertise, much of which has never been formalized, and some of which may never even have been commented properly. For another, we literally have no experience at this point in setting up large-scale attempts at integrating these various physical aspects in simulations of dense stellar systems. Given this situation, it would seem most prudent to start experimenting with existing codes, matching them with toy models first, and then with each other, in order to gain some initial experience concerning their collective behavior.

Our MODEST acronym lends itself very conveniently to express this aspect of our philosophy: our approach is one of MODifying Existing STEllar codes. We hope this reading will avoid the false impression that either we or our projects could possibly be considered modest.

The main price to pay for MODESTy is that working with black boxes as components allows a swapping of those black boxes, which will make validation of the final results much easier. If we can easily change the use of one stellar evolution code for another, for example, we can quickly get an impression of the relative accuracy of those codes (to the extent that they are truly independent). Such a divide-and-conquer approach is crucial in proving correctness of the outcome of highly complex large-scale simulations.

To sum up, the challenge is to construct a software framework that allows us to model a wide variety of astrophysical situations, using existing programs that encapsulate specialized astrophysical expertise. Where necessary, we will write wrappers, drivers, and other modules that will communicate and translate information between the already existing programs. What is needed first is to define a convenient and well-specified set of interfaces that allow us to mix and match the various unrelated programs, written in different languages and in different styles, in such a way that they can appear as black boxes to each other and to one or more driver programs.

3.2. Specification of Interfaces

A central task in setting up a software framework for any type of large-scale simulation is the specification of interfaces between different computational modules. On the one hand, we must be careful not to force any particular organization on the variables that are private to each module. On the other hand, we should maintain consistency across an interface.

In general, for each interface there should be an agreement about the particular names and types of a minimal set of variables that will be passed through the interface. This does not mean that the modules themselves will be forced to use those externally constrained names and types; it is straightforward to provide extra levels of data abstraction, for example by writing wrappers around existing modules that translate the information from the relevant variable within the module to the names and types specified in the interface.

It also does not mean that interface specifica-
tions will be put in stone. On the contrary, an essential aspect of good interface design is to leave open the possibility of significant future extensions of what will be passed through an interface, perhaps totally unforeseen at present. The only requirement will be compatibility with older specifications of the interface.

In the concrete case of simulations of dense stellar systems we have three broad classes of existing programs that already model aspects of astrophysical phenomena. These are stellar dynamics, stellar evolution and stellar hydrodynamics. In the future, we may want to write a special driver/scheduler/manager program, but in existing stellar dynamics programs, more than 90% of the lines of code are already dedicated to such orchestration details. Therefore, initially at least it will be simplest to consider the evolution and hydrodynamics programs as black boxes that are invoked by the dynamics program when needed. Later implementations may grant a more active role to the evolution and hydrodynamics programs, if that would reduce complexity and dependencies.

Given that current codes are written in totally different styles and in different languages, our first task is to specify interfaces and to develop wrappers around existing programs that are compliant with those interfaces. Since we all have different backgrounds, we can help reach this goal in different ways, according to what we enjoy doing and what we’re already good at. It would be counterproductive to require a specialist in stellar evolution to suddenly learn new computational science tools he or she is not comfortable with; similarly it would be counterproductive to require a stellar dynamicist to become familiar with the inner details of how a stellar evolution code is set up. If someone has programmed in Fortran for thirty years, there is absolutely no reason to require this person to learn and use other languages (although it might be fun). There is even no need for that person to do any work on writing the wrapper around his or her program for the interface; the minimum collaboration needed is a clear specification of which variables in his or her program correspond to those specified for the interface. Providing those in a Fortran common block, say, would be fine if that is the style this person is used to program in.

4. Stellar Dynamics

The earliest published $N$-body simulations are the 10-body runs by von Hoerner (von Hoerner 1960). By the early seventies, larger systems could be modeled, up to $N = 500$. Key ingredients in making it possible to integrate these larger systems were the use of individual time steps (Aarseth 1963), as well as special treatments of binaries through various ways of analytical and other forms of regularization (Aarseth 1985). The two leading families of $N$-body codes tailored to simulations of dense stellar systems are NBODYx$^3$ (Aarseth 2002, Spurzem and Baumgardt 2002), and the \texttt{kiro} integrator distributed with the Starlab$^4$ software suite (Portegies Zwart \textit{et al.} 2001; Hut 2002). For a general treatment of dense stellar systems, and especially of rich star clusters, see Heggie & Hut (2002).

Hardware improvements were important as well, in reaching the goal of simulating whole star clusters. The GRAPE project of constructing special-purpose computers, initiated at Tokyo University in 1989, has led to the installment of dozens of such computers world wide. An example of calculations made possible by the GRAPE was the first demonstration of the occurrence of core oscillations in direct $N$-body systems by Makino (1996), using the GRAPE-4 to perform a 32,000-body calculation. The acronym “GRAPE” stands for GRAvity PipE; more information can be found on the GRAPE web site,$^5$ in the book by Makino & Taiji (1998), and the review articles by Hut & Makino (1999) and Makino (2002).

4.1. The Physical Role of Stellar Dynamics

Stellar dynamics is perfectly adequate in modeling the motions of stars as point masses moving under the influence of gravity, even in dense stel-

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$^3$NBODY4 is optimized for use on the GRAPE special-purpose hardware; other members include NBODY6 for general-purpose single processor computers, and NBODY6++ for parallel computers

$^4$http://www.manybody.org/starlab.html

$^5$http://www.astrogrape.org
Figure 1. Three aspects of simulations of dense stellar systems, and three ways to classify them into two categories.

lar systems, unless individual stars approach each other to within a few stellar radii. When that happens, the internal structure of the stars has to be taken into account, and we have to switch to a hydrodynamics module to follow the encounter, which may lead to mass transfer and even to the merging of two or more stars. After the dust has settled, we then have to update the stellar evolution models for the stars involved, and in case of mergers we will have to construct new models from scratch, often with highly unusual chemical compositions. All of this has to happen automatically, which means that the individual modules have to be robust, and that the interfaces should be well-defined.

The three types of physics involved in stellar interactions are sketched in Fig. 1. Each type plays a unique role in terms of type of degrees of freedom, time scale, and duration. For example, stellar dynamics is concerned with external degrees of freedom, on a dynamical time scale, and for the duration of the whole history of the star system.

To start with the first distinction: the hydrodynamics and evolution codes are only concerned with the internal degrees of freedom of the stars, whereas the stellar dynamics module orchestrates the evolution of the external degrees of freedom: positions and velocities and higher derivatives thereof. The dynamics needs to know the masses and radii of the stars, the masses to compute gravitational forces and the radii to warn for possible collisions, but it only actively updates the positions (and velocities, accelerations, jerks, etc.) of the system.

A second distinction is given by the time scales on which the different processes evolve the stars. Stellar dynamics and hydrodynamics both use explicit integration schemes in order to follow the stars on a dynamical time scale. Stellar evolution codes, in contrast, use implicit integration schemes to follow the changes in internal structure of a star on thermal and nuclear time scales. The physical reason is that dynamical equilibrium can be assumed to be accurately preserved during almost all stages of stellar evolution. In contrast, it is exactly the deviation from dynamical equilibrium that drives the hydrodynamical phenomena.

The situation is intermediate in the case of stellar dynamics: a Fokker-Planck code, for example, follows a star system on a “thermal” (two-body relaxation) time scale, but direct N-body codes follow all stars on a dynamical time scale, which is necessary to accurately model phenomena involving binaries and multiple star systems.

The third distinction concerns the duration of the relevance of each physical process. Each star in the system will always be represented as a point mass in the stellar dynamics part of the code, and as a star with internal structure in the stellar evolution part of the code. While these two representations persist throughout the full history of a simulation, the third type of representation, offered by a hydro code, is temporary. Only during a close encounter do hydrodynamical models for a few stars spring to life, and they are again discarded after they have done their duty, after a period comparable to a few crossing times of the system (a day or so for normal stars, a year at most in the case of giant stars).

Note that this description only applies after

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6This simple distinction may become blurred when more complex dynamical processes, such as tidal interactions and possibly tidal capture, are considered.
the stars have been formed. During the earlier stages, when a star system is born through the collapse of molecular clouds, hydrodynamics also plays a more global role. Like stellar dynamics, the hydrodynamics describes the external degrees of freedom of the gas clouds, and it is a persistent element in the computer code for the simulation, as long as gas remains present in the system. See §9 for more details.

### 4.2. The Computational Role of Stellar Dynamics

When we compare the complexity of the three physical processes, it is clear that stellar dynamics is by far the simplest, conceptually. The only computational task is the integration of Newton’s classical gravitational equations of motion. What could be simpler? In comparison, the dynamical fluid equations of hydrodynamics are far more subtle, largely because they are partial differential equations rather than ordinary differential equations. The possible occurrence of shock waves and turbulence has no analogy in the simple world of stellar dynamics. And the intricacies of stellar evolution are even more subtle, with the interplay between radiative transfer, nuclear energy generation, convection, the largely still unknown roles of rotation and magnetic fields, and so on.

Given this situation, why are state-of-the-art stellar dynamics codes so complex, and why are they still being improved, after forty years of collective experience in writing them? They answer lies in the fact that what we call a stellar dynamics code is in fact mostly a complex scheduling manager where almost all the logic is used to make sure that the integrations retain accuracy. In the thousands of lines of computer code in a modern stellar dynamics program, only a few hundred lines contain Newton’s force calculation and the integration thereof. All the rest of the code involves special forms of treatment for each star.

For instance, unlike almost all text book examples of the integration of differential equations, stars in N-body systems are integrated with individual time steps. In addition, close encounters between stars are treated in special ways, by constructing local coordinate systems to represent their positions in order to avoid round-off errors. Not only does the proper creation and destruction of these coordinate patches require quite a bit of intelligence in a dynamics code, the real fun starts when two or more such coordinate patches meet, and have to merge or split. And on top of all that, specific code is often written to avoid the numerical singularities involved in close encounters of stars, for example by adopting special treatments of unperturbed motion, or mapping the three-dimensional Kepler motion onto that of a four-dimensional harmonic oscillator through the Kustaanheimo-Stiefel transformation (cf. Aarseth 2002).

For all these reasons, the structure of a computer program that can model stellar dynamics, stellar evolution, and hydrodynamics is not well described by the schematic diagram in Fig. 1, that focuses only on physical processes. Instead, we can discriminate between three different aspects of a typical stellar dynamics program for dense stellar systems. The most straightforward part of the program governs the integration of the global objects in the system. These objects can be single stars, isolated binaries, triples or higher multiples, as well as temporarily interacting groups of stars. Each non-trivial object (anything that is not a single star) has additional internal gravitational degrees of freedom. For example, an isolated binary might be represented through an analytic expression in the form of a Kepler orbit, which can be used to predict the position of the stars when they are needed, during a relatively close encounter. And the dynamics of an interacting group of stars will be computed using its own local coordinate system, possibly using regularization methods.

Besides this division of labor between global and local gravitational interactions, each stellar dynamics code contains a third segment in the form of a piece of code that takes care of the overall scheduling of all events that occur. This scheduler acts as a system clock that tells each particle when it has to move (remember that different particles have different time steps), and in addition it issues the orders for the creation and destruction of local coordinate patches, as well as their
Figure 2. The stellar dynamics part of a combined simulation code contains three different parts: one modeling the global dynamics, one for the local dynamics, and an overall scheduler. It might be most natural to let the scheduler do the synchronization for the hydrodynamics and stellar evolution modules as well.

merging and splitting. Therefore, from a computational point of view, the neat division into three different physical processes translates into the five different computational processes sketched in Fig. 2.

The same logic that is in place in current stellar dynamics codes already effectively contains an interface between the local and global part of the gravitational calculations, as well as a mechanism for the partly asynchronous evolution of the various components. It would be natural to use these features, in our philosophy of trying to make only minor modifications to existing stellar codes, as stressed in §3. Such a strategy would lead to the following requirements, at least for initial progress in realizing the physics of Fig. 1:

1. make a clear and clean separation between the local and global gravitational components of current stellar dynamics codes.

2. make a clear and clean separation of the synchronization part of such a code from the rest of the dynamics.

3. specify interfaces between these three parts of a dynamics code, in order to allow a homogeneous treatment between those interfaces (currently internal in dynamics codes) and interfaces with the external modules that govern hydrodynamics and evolution.

4. construct interfaces between the scheduler and the hydrodynamics and stellar evolution modules along similar lines as was done for stellar dynamics.

5. finally define the interface between stellar dynamics and hydrodynamics, as well as between hydrodynamics and stellar evolution, in such a way that the internal stellar properties can be modeled with a similar predictor-corrector structure as is currently done for the external variables in stellar dynamics codes.

To clarify point 4): for the foreseeable future, it is probably most efficient to represent a hydrodynamics module for a star through a Smooth Particle Hydrodynamics (SPH) module, a form of an $N$-body code where each particle is given an entropy in addition to a mass and position and velocity. Computationally, both hydrodynamical and stellar dynamical degrees of freedom are then modeled as external degrees of freedom, while the stellar dynamics is still modeled as a black box with internal degrees of freedom. When two or more stars come close, their gravitational point mass external information, together with their stellar evolution internal information, are used together to construct a temporary hydrodynamic representation. After the encounter, the hydrodynamic information is translated back left and right into the stellar dynamics and evolution modules. With the hydrodynamic module as a go-between, there may be no need for the stellar dynamics and stellar evolution modules ever to talk to each other directly.

To clarify point 5): note that an $N$-body code is not really a code that follows $N$ point masses. Rather, its internal representation deals with $N$ orbit segments. Each star has a position, velocity and higher derivatives that have last been calculated at a give time. Based on that information,
the future orbit of the particle can be predicted up to a particular later time, a type of ‘latest sales date’ for which the accuracy is guaranteed to stay within the required bounds. As soon as the system time exceeds this later time, that particular star will be updated, so that its ‘latest sales date’ again is pushed into the future, beyond the current time. Until that new time is reached, all other particles can once again rely on the newly computed orbit segment to provide information about the given star when needed, at times other than the time at which this star was updated.

This elaborate mechanism that makes it possible to advance stars at individual time steps can be extended to the treatment of hydrodynamics and stellar evolution as well. What is needed in this case is a type of interface that can ask the stellar evolution module, for example, to provide an estimate of its near-future behavior, and a ‘latest sales date’ until which this information can be considered to be accurate. For a user at the stellar dynamics side of the interface, it is irrelevant whether such a prediction is a true prediction, or simply a reading of an entry in a table, or an actual calculation by evolving a stellar evolution model for some duration into the future. We will come back to these issues in §6.

5. Stellar Evolution

5.1. Background and motivation

We require a code that models the evolution of any star, either single or binary, from an arbitrary initial condition up to the end of its nuclear and thermal evolution. Such a code will have a wide range of applications, but the main application we consider here is for modeling dense stellar systems such as globular clusters, galactic nuclei or starbursts, where many stars (of order $10^5$ or more) interact with one another and would have to be modeled simultaneously. The requirements of such a code are therefore: (1) it should be able to run autonomously and without outside interference given a sufficient set of initial conditions for the star(s); (2) it is robust and gives a – hopefully meaningful – result under any conceivable circumstance; (3) it is sufficiently fast that an entire simulation of $10^5$ stars or more takes a reasonable amount of time (days at most); and (4) it should be able to interact with its surroundings at any time, i.e. yield information about its current status and also receive information that can modify its status. These requirements are by no means trivial! At present no full-scale stellar evolution code exists that satisfies all of these requirements, especially points (1) to (3). Every stellar evolution code of which we are aware is prone to break down and needs to be nursed at some point between the pre-main sequence and the white dwarf stage under the vast majority of circumstances, and is certain to break down irretrievably under many circumstances! Besides the codes are still too slow, taking of the order of a few minutes at least per star on the fastest processors available today, so that a full simulation would take months.

Nevertheless, it should be pointed out that at the lowest level, at least two codes that satisfy all four requirements are already in existence. These are not full-scale evolution codes but rather parametrize stellar evolution, using detailed evolution models as a basis wherever possible, and making educated guesses otherwise. These codes (i.e. bse, developed by Tout et al. [1997] and Hurley, Tout & Pols [2002]; and seba, developed by Portegies Zwart & Verbunt [1996]) perform well and have been successfully integrated into N-body codes (Hurley et al. 2001; Portegies Zwart et al. 2001). They appear to give reasonable results under many circumstances. However there are circumstances, the most common of which – in a dense stellar environment – are probably the occurrence of mergers and collisions, where the result of this approach probably has very little to do with reality (cf. point 2). It is especially with these collision products in mind, as well as the fact that with 100,000 stars interacting something unexpected and unparametrized is almost certain to happen, that we would like to improve on these codes and make them more generally applicable.

It should be noted that a full-scale stellar evo-
olution code, named TYCHO, is freely available online\(^8\), courtesy of D. Arnett. TYCHO is an open-source, community code written in Fortran. For more details on the code, see Young et al. (2001) and references therein.

In what follows we will discuss what has been done so far, what needs to be improved, and how this can best be achieved. We first discuss the situation for single stars and then the more complex situation for binary stars.

### 5.2. Single stars

The theory of single-star evolution is rather well-developed, although major uncertainties remain. In particular convection can only be modeled in a very crude way, while other internal mixing processes, e.g. induced by rotation, have only begun to be explored. The possible effect of internal magnetic fields has hardly been studied at all. Furthermore, mass loss is a major uncertainty, especially for very massive post-MS stars, Wolf-Rayet stars and AGB stars. Nevertheless we are confident that single stars can be modeled in a satisfactory way, and of all the uncertainties only mass loss directly affects the dynamical evolution of a star cluster.

Single stars as they occur in dense stellar systems can be divided into primordial stars, which should evolve no different than single field stars, and merged stars, the products of collisions or mergers, which may evolve quite differently from primordial stars.

#### 5.2.1. Current status and shortcomings

In both codes mentioned above, BSE and SEBA, single-star evolution is modeled using a set of analytic formulae that have been fitted to detailed stellar evolution tracks. The BSE code uses the formulae constructed by Hurley, Pols and Tout (2000) that give several global stellar quantities, such as luminosity \(L\), radius \(R\), mass \(M\) and core mass \(M_c\), as a function of initial mass \(M_0\), metallicity \(Z\) and age \(t\). Some of the formulae are rather ad hoc fits that reproduce the shape of an evolutionary track in certain phases like the main sequence, while others represent (in a simple way) actual physics underlying the evolution, such as the core-mass luminosity relation that drives the evolution of low-mass giants and AGB stars. The fits also allow very fast evaluation of certain important evolutionary timescales. Mass loss is not included in the fits but parametrized separately, so that different mass loss prescriptions can be used in conjunction with the formulae. The formulae also provide other global quantities like the moment of inertia and the depth of the convective envelope, so that the rotational evolution can be modeled (if assumed rigid) as well as magnetic braking.

Although this approach has been applied successfully both in ordinary (binary) population synthesis and in dynamical studies, it has several shortcomings. First and foremost, since the formulae have been fitted to standard stellar models, they can only be expected to represent the evolution of primordial stars. Merged stars, on the other hand, are expected to have rather different internal structure (i.e. composition profiles) and so are probably not very well represented by the formulae. As recent hydrodynamical studies have shown (see §7), the structure of collision products is neither homogeneous nor resembles that of a primordial star with the same total mass. Furthermore, the collision products rotate very rapidly and are initially strongly out of thermal equilibrium. At present, however, the formulae are being used to represent merger products and ordinary stars alike.

Also very massive primordial stars, whose evolution is determined to a large extent by mass loss, are represented rather poorly because the formulae are based on constant-mass models even though mass loss is taken care of when applying the formulae. Although such very massive stars only form a tiny fraction of the initial population of a globular cluster, their evolution and mass loss (which is poorly constrained in the first place) is crucial for the early dynamical evolution of a cluster. On the other hand, for the other much more common types of star with very strong mass loss, low- and intermediate-mass AGB stars, a formulaic approach is arguably the best way of representing their evolution. Their evolution is driven by a core-mass luminosity relation, which itself is not or only weakly dependent on mass loss. Mass

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\(^8\)\url{http://chandra.as.arizona.edu/~dave/tycho-intro.html}
loss is however crucial in determining the lifetime of the AGB phase.

Another shortcoming, that becomes serious when we start modeling collision and merger products in any detail, is that the formulae give no information on the internal composition and entropy profiles. Although it is possible – and useful for some purposes – to represent and follow surface compositions in a formulaic approach, deriving fitting formulae for entire composition and entropy profiles is an extremely daunting task, and given our experience with fitting even simple quantities like radius in a satisfactory manner, a task that no one can realistically be expected to carry out. For the same reason, it is unlikely that the formulae will be updated or replaced when a newer generation or extended set of stellar evolution models becomes available.

5.2.2. The best way forward

Given these shortcomings, a different approach will need to be taken in the future. Here we must make a distinction between primordial stars and merged stars.

Primordial stars, as argued above, all evolve alike for a given mass and metallicity, if we neglect for the moment the possible effect of (differential) rotation on the internal mixing processes. Therefore the most feasible approach, given the problems with speed and robustness of current evolution codes, is to interpolate in a library of stellar models. Such a library only needs two dimensions, \( M \) and \( Z \) (with a time-sequence for each entry), and so is of manageable size. For massive stars, perhaps several libraries should be computed/compiled for different mass-loss prescriptions. On the other hand, for stars with a clear core-envelope structure that follow a core-mass luminosity relation, i.e. AGB stars and low-mass giants, parametrizing the evolution with analytic formulae probably remains superior to table interpolation. For these stars the envelopes are homogeneous and (nearly) isentropic so it is sufficient to follow the surface composition and entropy. Hence a combination of table interpolation and analytic formulae seems the best approach for the near future.

It should be noted that interpolation between stellar models is a non-trivial task! It is of the utmost importance that interpolation is done between models in corresponding stages of evolution. Hence these evolution stages, at any rate the main critical turning points (e.g. terminal-age main sequence, base of the giant branch, etc.), should be identified on each track. Furthermore internal composition and entropy profiles need to be interpolated. Constructing an interpolation routine that can do all this automatically will be a difficult task, but the advantage is that once it is available, it can handle any library so that model libraries can be exchanged or updated at will. An alternative approach that circumvents the difficulties of interpolation is to use discrete models from the library to represent a range of stellar masses. For this to work the library has to be sufficiently densely spaced in mass (and metallicity), i.e. masses not differing by more than a few per cent. This may be sufficient for modeling the dynamics, but if we want to compare e.g. a color-magnitude diagram with observations we may still wish to interpolate in order to prevent a discrete appearance.

As for merger products, these have been shown to have internal structures quite unlike primordial stars. The resulting structures from hydro simulations rotate rapidly and are strongly out of thermal equilibrium, and in order to relax they need to shed a large amount of angular momentum. It is not clear from the hydro calculations how this is achieved, nor can stellar evolution codes answer this question, and this transition is likely to remain a grey area for quite some time. In any case it has to be supposed that somehow the merged star manages to get rid of its excess angular momentum, perhaps by shedding a small amount of mass in a disk.

It is also conceivable, but has yet to be verified by detailed calculations, that the strong differential rotation leads to additional mixing that significantly changes the chemical profile by the time the star has relaxed to thermal equilibrium. However, it seems unlikely that the merger products will be completely homogenized (§7). If the latter were the case, it would be conceivable to construct an extended library of stellar models, with an additional dimension namely the helium.
content, so that all of stellar evolution could be done by table interpolation. However, with arbitrary initial composition profiles this clearly becomes impossible.

It seems therefore necessary to be able to do on-demand stellar evolution calculations, for arbitrary initial entropy and composition profiles, during a cluster simulation. As discussed in Section 1, currently available codes do not satisfy the demands that make integration into an N-body code feasible. Speed is one problem, although this will become less and less important as processors get faster. Nevertheless some effort will have to go into making existing codes as fast as possible, by simplifying much of the input physics like the equation of state, employing a minimal nuclear network, and taking as few zones as is necessary to still achieve reasonable accuracy. In this way it should be possible, with current processors, to evolve a star from the zero-age main sequence to the start of double-shell burning in under one minute. If only merger products are calculated this way, this may not slow down a full N-body simulation too drastically, particularly if many stars are computed in parallel on separate processors.

A much more daunting problem is robustness. Although some codes can now evolve unaided through the helium core flash and through many thermal pulses along the AGB, this cannot be expected to be reliable under all circumstances (in particular the unusual circumstances that we are interested in). Nor is it desirable, because each thermal pulse cycle takes as much computing time as the entire evolution up to the first pulse. So even if codes can be made faster and more robust, we may still want to parametrize the AGB phase, and perhaps skip over the He core flash by using a set of pre-calculated zero-age horizontal branch models.

5.3. Binary stars

Binary star evolution presents a number of major problems of long standing that have yet to be satisfactorily resolved (see, for example, Shore, Livio & van den Heuvel 1994). A comprehensive description of how these problems are dealt with (or circumvented) in one particular recipe-based binary evolution code is given by Hurley, Pols & Tout (2002). We enumerate here only some of the more prominent ones.

Contact binaries, which are characterized by large-scale energy exchange between components in their common envelopes, account for nearly 1% of solar-type stars in the solar neighborhood. No established model exists for the physics of that energy exchange, even though in extreme cases it must account for as much as 99% of the energy radiated by the less massive star. Evolutionary models of contact binaries, using heuristic models of energy exchange, predict long phases of semi-detached evolution which are not observed, or else demand such rapid angular momentum loss in order to suppress the semi-detached state that they cannot account for the abundance of contact binaries. Not even heuristic models exist for early-type contact configurations, which nevertheless arise with startling frequency in models of massive binary evolution.

Evolutionary analyses of individual Algol-type binaries (longer-period semi-detached binaries with low-mass subgiant donors) almost invariably demand that they have lost a significant fraction of their initial orbital angular momenta (and possibly also significant fractions of the mass being transferred between components) to have arrived at their present evolutionary state. It is widely believed that magnetic stellar winds are responsible for these angular momentum losses, and also those which drive unevolved binaries into contact and cataclysmic binaries into mass transfer; yet the magnetic stellar wind braking rates adopted in evolutionary calculations are almost invariably gross extrapolations from the much weaker rates deduced empirically from rotation velocities of main sequence stars in young star clusters.

The affinity of other close, but not yet mass-transferring, binaries (the RS CVn binaries) for nearly equal or slightly reversed mass ratios, strongly suggests that normal stellar wind mass loss rates can be amplified significantly as stars approach their Roche lobes, but no physical model exists to quantify this amplification.

Field cataclysmic binaries (as also low-mass X-ray binaries and close double white dwarfs) are clearly products of dissipative evolution with a
more slowly rotating common envelope.\textsuperscript{9} Realistic three-dimensional models of common envelope evolution, including radiative transfer, from onset to completion remain an unrealized dream. Rather, population synthesis models (for the distribution of properties of an ensemble of coeval binaries) rely upon simple energy arguments plus an efficiency parameter to estimate the outcomes of common envelope evolution. Even so, evolutionary analyses of the origins of known close double white dwarfs frequently demand efficiencies greater than unity, an unmistakable sign of deficiencies in even simple accounts of the energy budgets of these binaries.

Certain types of binaries (for example, massive X-ray binaries, symbiotic stars, and barium stars) are fueled or created through wind accretion. Here as well, three-dimensional radiation-hydrodynamical models are needed for realistic treatments of mass and angular momentum accretion.

Finally, while the broad effects of supernova kicks on the dynamics of the binaries in which they occur have been explored, the dependence of the magnitude and direction of those kicks on the history, mass, and orbit of the exploding star remains unknown, but potentially of critical importance to the survival or disruption of those binaries.

5.3.1. New issues

To this litany of unsolved problems in close binary evolution must now be added several which are unique to a dense stellar environment, or at any rate assume far greater significance there.

Close tidal encounters or mergers can impart to the component stars or merger product rotational angular momenta which would have been impossibly large for a main sequence star to support.\textsuperscript{10} Theoretical models are not yet capable of answering how a thermally-distended merger product may be capable of shedding its excess angular momentum and relaxing to a normal, pressure-supported (if still rapidly rotating) state.

Mergers and binary mass transfer can also give rise to non-canonical stars, by which we mean stars with chemical profiles which never occur among single stars – mass-losing stars with anomalously-large cores for their masses, or mass-gaining stars with anomalously-small cores. Such stars appear regularly in binary mass transfer calculations, but no systematic survey of their evolution exists, nor may one even be practical. These are stars for which it will likely become necessary to integrate on-demand stellar evolutionary calculations into evolving cluster dynamical models.\textsuperscript{11}

Another issue arising uniquely in dense stellar clusters is the role of small perturbations from passing stars – energy exchanges too small to be of consequence for cluster evolution – which may nevertheless exert a profound influence on binary evolution. For example, mass transfer rates in cataclysmic binaries scale exponentially with the ratio of the difference between stellar and Roche lobe radii to the stellar atmospheric pressure scale height, a ratio which is typically of order $10^{-4}$. Fractional perturbations to the orbital separation of this magnitude or higher could profoundly effect the outburst behavior of those variables, or even conceivably drive the white dwarf to evolve back toward the giant branch, triggering a new common envelope phase of evolution. Existing detailed models of close binary mass transfer rest on the stability of mass transfer to small-amplitude perturbations, but the stability of those models to large-amplitude perturbations remains unexplored territory.

Among binaries evolving in isolation, theory generally predicts that (post-supernova binaries excepted) any orbital eccentricity will be tidally damped to insignificance before Roche lobe over-

\textsuperscript{9}This phenomenon is widely labeled ‘common envelope evolution’, but should not be confused with the evolution of contact binaries within a quasistatic common envelope.

\textsuperscript{10}This circumstance can arise in ordinary binary mass transfer as well, although an argument can be made from the survival of Algol-type binaries that tidal torques must in those cases be capable of mitigating the concentration of angular momentum in the accreting star.

\textsuperscript{11}An efficient implementation of this strategy will, however, require the capability of interpolating detailed interior models from a library of single star models for a given mass and age as predicate for constructing the desired non-canonical star; and likewise the ability to identify circumstances in which the interior of a non-canonical star has converged so closely with that of a canonical star that it no longer need be followed in detail.
flow actually occurs; but that circumstance cer-
tainly cannot hold among the perturbed systems
just described, in which the general problem of
mass transfer in eccentric binaries must be revis-
ited. Physically sound solutions to all of these
problems will require a more fundamental under-
standing of the sources and properties of stellar
viscosity than is now at hand.

5.3.2. Prospectus
The added dimensionality of binary and merger
evolutionary problems effectively precludes the
practicality of a library look-up approach as ad-
vocated above for single star evolution. The un-
certainties in binary evolution, particularly in the
all-important mass and angular momentum loss
rates, are so great as to vitiate any attempt at
present to build such a library. Rather, the most
practical approach seems clearly a recipe-based
formalism, as is commonly used in population
synthesis studies of binary evolution, e.g. in the
BSE and SEBA codes mentioned in Section 5.1
above. This approach uses relatively simple, ap-
proximately formulae to describe the outcome of
mass transfer in a given situation. These recipes
have been built up piecemeal from a qualita-
tive understanding of the criteria which dictate
which evolutionary path a given binary will fol-
low. With no systematic survey of close binary
evolution in all three major dimensions (mass,
mass ratio, and orbital separation) yet practical,
major gaps inevitably remain in these prescrip-
tions, but they have the virtues of extreme speed;
and for any single binary, the uncertainties in its
outcome state are in most circumstances domi-
nated by the parametrization of mass and angular
momentum losses, which afflict detailed models
and recipes alike.

Despite the bleak perspective offered above on
the current state and capabilities of binary or
merger evolutionary models, one should not lose
sight of the fact that the duration of intense inter-
action in those models is typically extremely brief,
compared with cluster dynamical time scales.
Where that is not the case (mass transfer driven
on a nuclear or angular momentum loss – mag-
netic stellar wind or gravitational radiation – time
scale), the donor star is in thermal equilibrium,
and can (excepting non-canonical stars) be well-
approximated by an appropriate model from a
single-star library of evolutionary models. Mass
transfer rates and evolutionary pathways can be
derived implicitly from that library. One expects
that, at a given instant in time, most stars in
most binaries, whether interacting or not, can be
represented by members of a single star library.

6. Stellar Dynamics and Stellar Evolution
Interface

6.1. Single Stars, without Hydrodynamics
During the workshop, we discussed the devel-
opment of specifications for the interfaces be-
tween different modules in simulations of dense
stellar systems. As a concrete example, we fo-
cused first on the simplest interface, that between
stellar dynamics and stellar evolution, without
using hydrodynamics, and without allowing any
binaries. Such an interface could be used in an
$N$-body program where single stars can collide
and merge, while signaling the stellar evolution
counterparts of two merging stars to construct a
new model for the merger product.

In specifying such an interface, we do not want
to make any assumption about the computational
processes that may take place at either side of
the interface. The stellar evolution information
may be provided from look-up tables or fitting
formulae based on sets of evolutionary tracks that
have been computed earlier, or it may be provided
by an actual stellar evolution code running in real
time. The stellar dynamics information can come
from an actual $N$-body code, or it can come from
a simple toy model that effectively produces a
population synthesis of distributions of colliding
stars without any ‘live’ dynamics involved. In all
these and other cases, the interface should not
care what is happening at either side, as long as
the specifications for the interface are obeyed.

There are at least three quite different ways
to specify an interface between evolution and dy-
namics:

1. Minimal Interface. The stellar dynamics
code can ask the stellar evolution code to
take one step forward. This type of hand-
shaking places the least demand on the stel-
lar evolution side of things, since everything is driven from the stellar dynamics side.

2. **Multi-Criterion Interface.** The stellar dynamics code can ask the stellar evolution code to proceed for a specified increase in time, as long as a number of criteria are met (no unacceptably large changes in important physical variables). This gives the stellar dynamics more control over the situation. It requires some additional code to be written to steer the stellar evolution code, but nothing very complicated.

3. **Maximal Interface.** The other extreme would be for the stellar evolution part of the code to compute the complete future evolution of every new star created. This may involve reading a precomputed table in the case of a star that starts on the main sequence, or it may entail the production of a new table by running a stellar evolution code, in the case of a merger product.

While the first approach may be convenient, in that it requires the least amount of changes to existing stellar evolution codes, such a specification violates our requirement of a black-box approach. It is not clear what it would mean to ask a table look-up implementation of stellar evolution to ‘take a next step’; nor can one ask a fitting function to take a step.

In the second choice of interface there is a danger that the time steps requested are unnaturally small from a stellar evolution point of view, leading perhaps to unacceptable round-off error. However, such problems can be easily anticipated, for example by letting a star take an evolutionary step only when the accumulated time increments become comparable to its own natural integration time step.

In case 3), the stellar evolution module will provide a complete future history of each new star, and will make it available to the stellar dynamics side. This will make it possible for the dynamics to request the value of any physical quantity at any current time or any predicted time in the future. The main drawback is that too much computer time may be used in the complicated late phases of unusual stars, when those stars themselves may well merge again with another star before reaching those stages. This may not be a problem as long as the majority of such stars do end their life without further significant perturbations.

6.2. **Interface Function Specifications: an Example**

Below we describe in some detail what a multi-criterion interface could look like. Note that the specifications given there are only a first step toward the simplest possible case. A real specification should include a treatment of binaries as separate computational objects, which are far more complicated than those corresponding to single stars. And in any case, in practice we will wait to formalize the interface until we have developed some experience with both the single-star and the binary cases. We may want to start with an alpha specification, then a beta specification, and then freeze the specification in a public version, in the sense that future interfaces will have to at least respect the requirements listed in the public version.

Here is a wish list from the stellar dynamics point of view. We would like to give an $N$-body program access to the following functions, which can be considered as a type of library function call:

1) a star creation function
2) a star evolution function
3) a star destruction function
4) a function providing the mass of a star
5) a function providing the radius of a star
6) a function providing the current time for a star
7) a function providing the total helium fraction of a star
8) a function providing the total metallicity of a star
For concreteness, we will write the specifications for these functions in the form of desiderata for functions written in Fortran. Similar specifications can be prescribed for other languages, such as C or C++, and in many cases the interface will be used to connect two modules that may be written, say, in Fortran and C++.

The function

\[
\text{integer function CreateStar}(M, Y, Z) \\
\text{accepts as arguments real*8 variables for the initial mass, the helium abundance and metallicity of a star created at the zero age main sequence. The return value is a unique integer that acts as the identifier for the particular star that has been created. A negative return value will signal an error (e.g. not enough storage left; unreasonable initial conditions provided; or some other internal error in the stellar evolution module). The units for the variables are:} \\
M \text{ in solar masses} \\
Y \text{ helium abundance fraction by weight; } 0 \leq Y \leq 1 \\
Z \text{ metallicity abundance by weight; } 0 \leq Y + Z \leq 1 \\
The function
\]

\[
\text{real*8 function EvolveStar(id, dtmax, dMmax, dRmax, dYmax, dZmax) \\
accepts as first argument an integer variable for the identifier id, followed by five real*8 variables that determine halting criteria. The stellar evolution code will start evolving the star, from the current time } t_{\text{now}}, \text{ at which the mass, radius, and compositions are } M_{\text{now}}, R_{\text{now}}, Y_{\text{now}}, Z_{\text{now}}. \text{ The code will stop as soon as one of the following halting criteria is satisfied:} \\
\text{if the time } t \geq t_{\text{now}} + dt \\
\text{if the mass } M \text{ obeys } |M - M_{\text{now}}| > dM_{\text{max}} \\
\text{if the radius } R \text{ obeys } |R - R_{\text{now}}| > dR_{\text{max}} \\
\text{if the helium fraction } Y \text{ obeys } |Y - Y_{\text{now}}| > dY_{\text{max}} \\
\text{if the metallicity } Z \text{ obeys } |Z - Z_{\text{now}}| > dZ_{\text{max}} \\
The function returns the new time } t_{\text{now}}; \text{ a negative value for } t \text{ indicates an error condition. The additional units used here are:} \\
t \text{ in millions of years} \\
R \text{ in solar radii} \\
The function
\]

\[
\text{integer function DestroyStar}(id) \\
\text{accepts an integer id, the identifier for the star that should be destroyed. The function will remove that star, freeing up the memory assigned to it. Successful completion will be indicated by returning a positive or zero integer; a negative integer will indicate an error condition.} \\
The function
\]

\[
\text{real*8 getMass(id) \\
accepts an integer id, and returns the value for the mass of the corresponding star. Similarly the functions} \\
\text{real*8 getRadius(id) \\
real*8 getTime(id) \\
real*8 getY(id) \\
real*8 getZ(id)} \\
\text{accept an integer id, and return the values for the stellar radius, current, helium fraction, and metallicity, respectively.} \\
\]
real*8 m1,m2,newmass,newY,newZ
integer newstar
...
m1 = getMass(id1)
m2 = getMass(id2)
newmass = m1+m2
newY = (getY(id1)*m1 + getY(id2)*m2)/newmass
newZ = (getZ(id1)*m1 + getZ(id2)*m2)/newmass
newstar = CreateStar(newmass,newY,newZ)

This code fragment creates a homogeneous ZAMS star from the matter obtained by adding the previous two stars. This procedure assumes complete mixing and ignores transient effects that will die out during a thermal time scale, such as an increase in radius due to shock heating. Other complications, such as a possibly rapid rotation after the merger are neglected as well.

With a black box approach, it is vital to test for possible errors, since you have no idea what is going on internally. The right defensive programming approach would be to let the lines above be followed by error checks, with appropriate actions (here indicated by ... for each particular type of error):

if (newstar .lt. 0) then
  ...
endif
if (DestroyStar(id1) .lt. 0) then
  ...
endif
if (DestroyStar(id2) .lt. 0) then
  ...
endif

For any serious production runs, the above treatment will need to be extended to model binaries as well. Such an extension may well imply modifications of the above simple treatment. Therefore, we do not intend our presentation here to be definitive in any way. Indeed, even for single stars, we may want to extend the above interface, for example by including the possibility of a star receiving a kick-velocity at the time of a supernova explosion. The only requirement will be that an agreed-upon future version of the above specification, when adapted as a standard, should remain valid in later versions that will be upwards compatible with that standard.

7. Stellar Hydrodynamics

7.1. Smoothed Particle Hydrodynamics

Hydrodynamic interactions such as collisions and mergers can strongly affect the overall energy budget of a cluster and even alter the timing of important dynamical phases such as core collapse. Furthermore, stellar collisions and close encounters are believed to produce a number of non-canonical objects, including blue stragglers, low-mass X-ray binaries, recycled pulsars, double neutron star systems, cataclysmic variables and contact binaries. As discussed in §5, these stars and systems are among the most challenging to model, and they are also among the most interesting observational markers. Predicting their numbers, distributions and other observable characteristics is essential for detailed comparisons with observations.

In galactic nuclei, collisions are very energetic events that typically result in two unbound stars that have suffered mass loss (a kind of “fly-by”). Mergers are actually a rare outcome, as collisions with small impact parameters often result in complete destruction of the parent stars (Freitag & Benz 2002a). In globular clusters, the velocity dispersion is less than the escape velocity from the surface of a parent main sequence star, and therefore mergers are much more likely. In any case, the structure and chemical composition profiles of a collision product are clearly of central importance, because they determine its observable properties and evolutionary track in a color magnitude diagram (e.g. Sills & Bailyn 1999).

Three-dimensional hydrodynamic simulation is one means to study stellar collisions and to determine the trajectories and interior profiles of the resulting products. Mostly using the Smoothed Particle Hydrodynamics (SPH) method, numerous scenarios of stellar collisions and interactions have been simulated in recent years, including, for example, collisions between two main sequence stars (Benz & Hills 1987, Lai, Rasio & Shapiro...

SPH is a Lagrangian technique in which the system is broken up into a large number of fluid particles whose positions and velocities are integrated forward in time according to hydrodynamic and self-gravitational forces. Local densities and hydrodynamic forces at each particle position are calculated by a kernel estimation that involves summing over nearest neighbors. For an overview of the basic SPH equations, see, for example, Monaghan(1992) or Rasio and Lombardi (1999).

The so-called entropic variable \( A \equiv P/\rho^{\Gamma} \) turns out to be critical for understanding the physics of mergers and therefore the results of SPH simulations. Here \( P \) is pressure, \( \rho \) is density, and \( \Gamma \) is the adiabatic index of the gas (assumed here to be constant). Given the importance of \( A \), we will first discuss this quantity in the context of single, isolated stars. It is straightforward to show analytically that the condition \( dA/dr > 0 \) is equivalent to the usual Ledoux criterion for convective stability of a non-rotating star (Lombardi et al. 1996). The basic idea can be seen by considering a small fluid element inside a star in dynamical equilibrium. If this element is perturbed outward adiabatically (that is, with constant \( A \)), then it will sink back toward equilibrium only if its new density is larger than that of its new environment. Because pressure equilibrium between the element and its immediate environment is established nearly instantaneously, the ratio of densities satisfies

\[
\frac{\rho_{\text{ele}}}{\rho_{\text{env}}} = \left(\frac{A_{\text{ele}}}{A_{\text{env}}}\right)^{-1/\Gamma}.
\]

Therefore, a fluid element with a lower \( A \) than its new surroundings will sink back down toward the equilibrium position. Likewise, if an inwardly perturbed fluid element has a larger \( A \) than its new environment, buoyancy will push the element outwards, back toward equilibrium. As a result, a stable stratification of fluid requires that the entropic variable \( A \) increase outward: \( dA/dr > 0 \). In such a star, a perturbed element will experience restoring forces that cause it to oscillate about its equilibrium position. For a detailed discussion of the stability conditions within rotating stars, see §7.3 of Tassoul (1978) or Tassoul (2000).

In practice, SPH calculations show that, even in rapidly rotating stars, fluid distributes itself in such a way that the entropic variable \( A \) increases outwards.

For a merger of stars, SPH simulations reveal that fluid elements with low values of \( A \) do indeed sink to the bottom of a gravitational potential well, and the \( A \) profile of a merger product in stable dynamical equilibrium increases radially outwards. Because this \( A \) profile is typically steep, especially in the outermost layers, collision products, in contrast to normal pre-main sequence stars, do not develop convective envelopes. An additional consequence of having the fluid stratify itself according to \( A \) is that parent stars are not thoroughly mixed during collisions; instead, strong chemical composition gradients are present even in the final configuration. The stellar evolution of collision products therefore can depart significantly from that of normal stars that begin their lives as chemically homogeneous, “zero-age” main sequence stars.

Because the quantity \( A \) depends directly upon the chemical composition and the entropy, it is conserved for each fluid particle during gentle, adiabatic processes. During a collision, the entropic variable \( A \) of a fluid element can increase due to shock heating. However, at least in open, globular and young compact star clusters, the relative impact speed of two stars is comparable to the speed of sound in these parents: both speeds are of order \((GM/R)^{1/2}\), where \( G \) is Newton’s gravitational constant, and \( M \) and \( R \) are respectively the mass and length scales of a parent star. Consequently, the resulting shocks have Mach numbers of order unity and shock heating is relatively weak. Therefore, to a reasonable approximation, a fluid element maintains a constant \( A \) throughout a collision. Hyperbolic collisions,
appropriate in galactic nuclei, do result in significantly more shock heating; however, the brunt of the shocks are absorbed by what becomes the ejected mass, thereby shielding the cores of the parent stars at least in less extreme cases.

7.2. Generating Collision Product Models, Quickly

A substantial fraction of the stars in a cluster will experience a collision sometime during their lifetimes. The direct integration of low resolution SPH calculations into a cluster evolution code may allow the modeling of such events in the not very far future. However, a single high resolution hydrodynamic simulation can typically take hundreds or even thousands of hours to complete. Therefore, the excessive computing time required of hydrodynamics simulations makes it extremely impractical, if not currently impossible, to couple them directly to stellar dynamics calculations.

One solution, taken by Freitag & Benz (2002a), is to calculate first an extensive set of hydrodynamics simulations, varying the parent stars, as well as the eccentricity and periastron separation of their initial orbit. The SPH database of Freitag & Benz treats all types of hyperbolic collisions between main sequence stars: mergers, fly-bys and cases of complete destruction. The tremendous amount of parameter space surveyed precludes having high enough resolution to determine in detail the structure and composition profiles of the collision products for all cases; however, critical quantities such as mass loss and final orbital elements are indeed determined accurately. By interpolating among these hydrodynamics results, Freitag & Benz (2002b) have successfully integrated collisions into a Monte Carlo star cluster code, yielding the most realistic treatment ever of stellar collisions in a stellar dynamics code.

It should be noted that, even without relying directly on SPH results, certain aspects of collisions could be modeled in a cluster simulation code using existing techniques. For example, Fig. 3 of Freitag & Benz (2002a) shows that the mass loss in high velocity collisions with relatively large impact parameters \( |d_{\text{min}}/(R_1 + R_2) | > 0.5 \) is surprisingly well predicted by a simple method devised by Spitzer & Saslaw (1966) and based on conservation of momentum and energy. Such collisions are likely to occur in a galactic nucleus near a massive black hole, as high velocities quench focusing and make collisions with small impact parameters rare.

As an approach for generating merger product models without running hydrodynamics calculations, Lombardi et al. (2002) have developed a method that calculates the structure and composition profiles from simple algorithms based on conservation laws and a basic qualitative understanding of the hydrodynamics. The thermodynamic and chemical composition profiles of the simple models, as well as their subsequent stellar evolution, agree very well with those from the SPH models. Because the method takes only a few seconds to generate a model on a typical workstation, it becomes feasible to incorporate the effects of mergers in dynamics simulations of globular clusters. The algorithms have been implemented in an easy to use software package dubbed “Make Me A Star.”

The underlying principle behind this method exploits the two special properties of \( A \) discussed in §7.1: Namely, the entropic variable \( A \) will (1) increase outward in a stable star and (2) be approximately conserved during a collision. Therefore, to a good approximation, the distribution of fluid in a merger product can be determined simply by sorting the fluid from both parent stars in order of increasing \( A \): the lowest \( A \) fluid from the parent stars is placed at the core of the merger product and is surrounded by shells with increasingly higher \( A \). This treatment is further improved upon by modeling the shock heating, hydrodynamic mixing, mass ejection, and angular momentum distribution with physically motivated fitting formulas calibrated from the results of SPH simulations. Although the algorithms currently are capable of treating only parabolic collisions between stars obeying an ideal gas equation of state, the method and code have been developed with the intent of ultimately generalizing to other collision scenarios.

\[ \text{http://vassun.vassar.edu/~lombardi/mmas} \]
8. Stellar Dynamics, Hydrodynamics and Stellar Evolution Interfaces

8.1. Including Hydrodynamics

Here we discuss how the communication between a stellar dynamics (SD) and stellar evolution (SE) module, presented in §6, can be extended to include an additional interface with a stellar hydrodynamics (SH) module. The SD module will continue to be the scheduler and manager, passing only the minimum amount of data necessary to any SE or SH routine. However, for hydrodynamic processes to be modeled, we must now also allow for the storage and retrieval of stellar structure and composition profiles. The general purpose of the SH module is to take such profiles for parent stars, and return profiles for newly created collision product(s). Rotation of collision products is neglected in this simple interface, but could be treated by also including a profile for the specific angular momentum.

SH is likely to play an important role in single-binary and binary-binary interactions. In such cases, a merger product created in an initial collision will have a greatly enhanced collisional cross-section due to the shock heating of fluid. A second (and even a third) collision can likely result before the first merger product significantly contracts as it thermally relaxes. Communication among the various modules is therefore high: the SD module tracks the stellar trajectories, the SH module generates collision product models, and the SE module evolves the structure of these models. The simple SH module that we discuss below would treat each successive collision separately, neglecting tidal forces from nearby stars and the slight possibility that a third star could collide or strongly interact while the first two parent stars are in the process of colliding.

8.2. Routines Provided by the Stellar Hydrodynamics Module

There are two main routines supplied by the SH module: (1) a stellar collision function that determines what happens during a collision, and (2) a subroutine that returns the structure and composition profiles, as well as the position and velocity, of any collision products. For concreteness, we will write the specifications for these functions in Fortran (taking some liberties with indentation and continuation lines).

The function

```fortran
integer function CollideStars(r,v, 
  mProfile1,rProfile1, 
  PProfile1,rhoProfile1, 
  chemicalProfiles1, 
  mProfile2,rProfile2, 
  PProfile2,rhoProfile2, 
  chemicalProfiles2, 
  numShells1,numShells2, 
  numChemicals)
```

accepts input arguments declared as follows.

```fortran
real*8 r(3), v(3) 
integer numShells1,numShells2, 
  numChemicals 
real*8 mProfile1(numShells1), 
  rProfile1(numShells1), 
  PProfile1(numShells1), 
  rhoProfile1(numShells1), 
  mProfile1(numShells2), 
  rProfile2(numShells2), 
  PProfile2(numShells2), 
  rhoProfile2(numShells2), 
  chemicalProfiles1(numShells1, 
                  numChemicals), 
  chemicalProfiles2(numShells2, 
                  numChemicals)
```

The arrays `r` and `v` specify, respectively, the relative position and velocity of parent star 2 with respect to parent star 1, in Cartesian coordinates. The integers `numShells1` and `numShells2` give the number of shells in which the structure and chemical composition profiles are stored for stars 1 and 2, respectively. The integer `numChemicals` gives the number of different chemical species that are being considered, presumed to be the same for each parent star.

The arrays `mProfile1`, `rProfile1`, `PProfile1` and `rhoProfile1` specify the structure of parent star 1 by specifying its enclosed mass, radial, pressure and density profiles, respectively. That is, `mProfile1(i)`, `PProfile1(i)`
and \( \rho_{\text{Profile1}}(i) \) give the enclosed mass, pressure and density, respectively, in star 1 at a spherical shell of radius \( r_{\text{Profile1}}(i) \), for any integer \( i \) in the range from 1 to \( \text{numShells1} \). These arrays contain redundant information that could also have been obtained by integrating \( dm = 4\pi r^2 \rho dr \) or the equation of hydrostatic equilibrium (if appropriate); nevertheless, as a matter of convenience it is useful to have all four arrays available.

The array element \( \text{chemicalProfiles1}(i,j) \) gives the fractional composition, by weight, of chemical species number \( j \) in shell \( i \). Similarly, the arrays \( m_{\text{Profile2}}, r_{\text{Profile2}}, P_{\text{Profile2}}, \rho_{\text{Profile2}} \) and \( \text{chemicalProfiles2} \) specify the structure and composition profiles of parent star 2. The last element of the radial profile arrays, \( r_{\text{Profile1}}(\text{numShells1}) \) and \( r_{\text{Profile2}}(\text{numShells2}) \), are taken as the stellar radii of the parents. Likewise, \( m_{\text{Profile1}}(\text{numShells1}) \) and \( m_{\text{Profile2}}(\text{numShells2}) \) are the masses of the parent stars. All values are stored in cgs units. Chemical composition profiles are dimensionless, as they represent the fractional abundance by mass.

The function \( \text{CollideStars} \) returns the number of collision products generated. For parabolic and weakly hyperbolic encounters the returned value will often be 1, but it could also be 2 (with the two stars having new mass and internal profiles). For strongly hyperbolic encounters, the returned integer is 0 when the stars are destroyed by the collision, and possibly, at least in principle, larger than 2 if multiple stars collapse out of a remnant gas cloud after a catastrophic collision. A returned value that is negative will signal an error condition (e.g., the input relative separation, velocity and stellar radii not being consistent with a close interaction).

Internally this routine will generate structure and chemical composition profiles for the collision product(s). This could be done in any number of ways, for example by actually running an SPH simulation, by interpolating SPH results, or with simple recipes or fitting formula (see §7). These profiles will be stored in memory until the next call to \( \text{CollideStars} \), and can be retrieved in the meantime through the subroutine \( \text{getProduct} \).

When called after the function \( \text{CollideStars} \), the subroutine

\[
\text{getProduct}(r_{\text{Product}}, v_{\text{Product}}, m_{\text{Profile}}, r_{\text{Profile}}, P_{\text{Profile}}, \rho_{\text{Profile}}, \text{chemicalProfiles}, \text{numShells})
\]

returns the position and velocity, as well as the structure and composition profiles, of the collision product(s) generated in the most recent call to \( \text{CollideStars} \). The returned arrays \( r_{\text{Product}} \) and \( v_{\text{Product}} \) specify, respectively, the relative position and velocity of a collision product with respect to the input (pre-collision) position and velocity of parent star 1, in Cartesian coordinates. The enclosed mass, radius, pressure, density and chemical composition profiles are returned as the \( \text{real*8} \) arrays \( m_{\text{Profile}}, r_{\text{Profile}}, P_{\text{Profile}}, \rho_{\text{Profile}} \) and \( \text{chemicalProfiles} \), respectively.

The first call to \( \text{getProduct} \) yields the profiles for the first collision product; the second call is for the second collision product, etc. Also returned, as the final argument to the subroutine, is the integer \( \text{numShells} \) specifying the number of shells in the structure and composition arrays. The \( \text{chemicalProfiles} \) array is two dimensional with the second argument running from 1 up to \( \text{numChemicals} \), automatically set to the same number of chemical species being considered as in the parent stars.

The declaration of the arguments for \( \text{getProduct} \) is therefore as follows.

\[
\begin{align*}
\text{integer numShells} \\
\text{real*8 rProduct(3), vProduct(3)} \\
\text{real*8 mProfile(numShells),} \\
\text{rProfile(numShells),} \\
\text{PProfile(numShells),} \\
\text{rhoProfile(numShells),} \\
\text{chemicalProfiles(numShells, numChemicals)}
\end{align*}
\]

8.3. Routines Provided by the Stellar Evolution Module

Even with the inclusion of a SH module, the \( \text{EvolveStar}, \text{DestroyStar}, \text{getMass}, \text{getRadius} \)
and `getTime` functions would not need to be modified from their versions described in §6. However, the `CreateStar` function does need to be generalized to allow for the creation of stars with arbitrary structure and composition profiles.

The function

```plaintext
def integer CreateStarFromProfiles(  mProfile, rProfile, PProfile,  rhoProfile, chemicalProfiles,  nucleonNum,  numShells, numChemicals)
```

fills this role, accepting as arguments real*8 arrays for the enclosed mass, radius, pressure, density and chemical composition profiles, in that order. The intent is that these arrays will have been generated through calls to `CollideStars` and `getProduct`. The two dimensional real*8 array `nucleonNum` specifies the chemical species being considered, with `nucleonNum(1,j)` and `nucleonNum(2,j)` giving the number of protons and neutrons, respectively, for species $j$.

The final two input arguments are the integers `numShells` and `numChemicals` that, respectively, specify the number of shells and chemical species represented by these arrays. As in the `CreateStar` function of §6, the return value is a unique integer that acts as the identifier for the particular star model that has been created, with a negative return value signaling an error.

With the introduction of an SH module, another new routine required from the SE module is a subroutine providing the current profiles of a star:

```plaintext
def getProfiles(id, mProfile, rProfile, PProfile, rhoProfile, chemicalProfiles, numShells, numChemicals)
```

This subroutine accepts an integer `id` identifying a particular star model. The output arrays contain the same type of information as the arrays returned by `getProduct` (see §8.2). The chemical profiles returned in `chemicalProfiles` are for the same species, in the same order, as when the star was created.

### 8.4. Implementation in the Stellar Dynamics Code

The following code fragment will collide two parent star models with id numbers `id1` and `id2`. When no errors result, the two parent stars will be destroyed, and any collision product models will be created. Error conditions would be handled in the portions of the code represented by “...”

```plaintext
def CollideStars(r,v, mProfile1, rProfile1, PProfile1, rhoProfile1, chemicalProfiles1, mProfile2, rProfile2, PProfile2, rhoProfile2, chemicalProfiles2, numShells1, numShells2, numChemicals)
```

```plaintext
numproducts =
def getProduct(rProduct(1,i), vProduct(1,i), mProfile, rProfile, PProfile, rhoProfile, chemicalProfiles, numShells)
def CreateStar( mProfile, rProfile, PProfile, rhoProfile, chemicalProfiles, nucleonNum, numShells, numChemicals)
```

Even if one wanted to make the perhaps crude approximation that a merger product some-
how became chemically homogeneous on a short timescale, the composition information provided by \texttt{getProduct} can still be nontrivial: the ejected mass in collisions comes preferentially from the outermost layers of the stars, and therefore the total composition fractions in a collision product are not exactly a simple mass average of the total fractions in the parent stars.

9. Formation of Stars and Stellar Systems

To be able to follow the entire life cycle of stellar systems, we need to understand how stars form, and in particular, how stars form in dense aggregates and clusters. This knowledge allows us to define astrophysically relevant initial conditions for the in-detail investigation of the subsequent dynamical evolution of the cluster – the main aim of the MODEST collaboration. We briefly review here scope and limitations of the current numerical models of molecular cloud fragmentation and star cluster formation and then add some further more general considerations.

9.1. Turbulent Fragmentation and the Formation of Stellar Clusters

Careful stellar population analysis indicates that most stars in the Milky Way (of order of 90%) form in open clusters with a few hundred member stars. Rich stellar clusters with several thousands to ten thousands of stars account for most of the remaining stars (Adams & Myers 2001). Very rich stellar clusters with several hundred thousand or millions of stars (e.g. globular clusters) are extremely rare, and contribute only a very small fraction of the entire stellar population of a galaxy.

To our current understanding, all stars are born in turbulent interstellar clouds of molecular hydrogen. The location and the mass growth of young stars are hereby intimately coupled to the dynamical cloud environment. Supersonic molecular cloud turbulence establishes a complex network of interacting shocks, where converging shock fronts generate clumps of high density. The density enhancement may be large enough for the fluctuations to become gravitationally unstable and collapse. This happens when the local Jeans length becomes smaller than the size of the fluctuation. However, the fluctuations in turbulent velocity fields are highly transient. The random flow that creates local density enhancements can disperse them again. For local collapse to actually result in the formation of stars, individual gravitationally unstable shock-generated density fluctuations must collapse to sufficiently high densities on time scales shorter than the typical time interval between two successive shock passages. Only then are they able to ‘decouple’ from the ambient flow and survive subsequent shock interactions. The shorter the time between shock passages, the less likely these fluctuations are to survive. Hence, the efficiency of protostellar core formation, the growth rates and final masses of the protostars, essentially all properties of the nascent star cluster strongly depend on the intricate interplay between gravity on the one hand and the turbulent velocity field in the cloud on the other (e.g. Klessen et al. 2000, or Heitsch et al. 2001).

Clusters of stars build up in molecular cloud regions where self-gravity overwhelms turbulence. Once individual gas clumps become gravitationally unstable within the star forming region, they begin to collapse. The gas density increases and a hydrostatic protostellar object forms in the center of the collapsing core. In dense clusters, collapsing gas clumps may merge, producing new clumps that then contain multiple protostars. Dynamical interactions are common, close encounters occur frequently and will drastically alter the trajectories, thus changing the accretion rates. This has important consequences for the final stellar mass distribution (Klessen 2001a,b). Already in their infancy, i.e. already in the deeply embedded phase, stellar clusters are strongly influenced by collisional dynamics. Turbulent molecular cloud fragmentation, competitive accretion, and protostellar interaction, all are highly stochastic processes. In essence, a comprehensive theory of star formation thus needs to be a statistical theory.

In sufficiently populous clusters O stars may form. Their intense UV radiation photoionizes the surrounding molecular cloud region, leading to the rapid expulsion of the residual gas on timescales typically faster than the dynamical
time from the cluster. This leads to rapid dispersal of a large fraction of the embedded cluster population. The velocity dispersion of the expanding population is a function of the binding energy of the embedded cluster, and in massive clusters it may reach a few tens km/s. Such kinematically hot components may lead to the thickening of thin galactic disks in those instances when a disk galaxy goes through a star-burst phase. The thick disk of the Milky Way galaxy, as well as the hitherto not understood steep rise of the age–velocity dispersion relation of solar-neighbourhood stars, may be a direct consequence of such processes. Star-cluster birth may therefore be an essential ingredient if we are to understand the structural and kinematical properties of galaxies (Kroupa 2002).

The following is a list of results of star cluster formation calculation that are of relevance in the context of MODEST, i.e. for combining star cluster formation with star cluster evolution:

1. Star clusters form fast, on timescales of order of a few free-fall times.

2. Star clusters form with a considerable degree of substructure.

3. Star clusters form with a very high initial binary fraction (larger than 60%). This is consistent with inverse population synthesis models (Kroupa 1995).

4. The stellar mass spectrum predicted by turbulent cloud fragmentation in cluster forming regions is consistent with observational determinations of the IMF (Klessen 2001b; IMF see Kroupa 2002) and extends down into the brown dwarf regime (Bate et al. 2002).

5. Massive stars begin to form first and are able to maintain a high accretion rate. This is because massive stars form from the most massive and densest gas clumps in the star forming region. As these clumps are dense, collapse progresses fast. And because they are very massive, they constitute a local minimum of the cluster potential and are able to attract the inflow of further gas. Massive protostars therefore experience high accretion rates over an extended period of time. Low mass stars form from low mass gas clumps and only benefit from a short period of peak accretion.

6. Star clusters are expected to form mass segregated. Massive stars form close to the cluster center, low mass stars are likely to form at large cluster radii. This is because massive clumps constitute the central region of the nascent cluster and low mass clumps are predominantly found at the outskirts. Altogether, one expects a radial dependence of the cluster IMF. The current star forming models, however, give only first hints of this effect. Small number statistics do not allow for detailed predictions yet.

7. During its first few million years, a star cluster will contain a mixture of stars on the main sequence and stars still in the pre-main sequence phase. Very massive stars enter the main sequence already during the main accretion phase, while the pre main sequence contraction phase of low mass protostars may last for several $10^7$ years (e.g. Palla & Stahler 1999).

8. Star formation likely is a self-regulated process. Bipolar outflows from young stars stir the gas in star forming regions, thus modulating the accretion efficiency. Radiation from young stars will heat the gas. If O or B stars form, they will ionize the gas in their surrounding and prevent further mass growth and star formation. The same holds for supernovae explosions, which will also blow away the cluster gas. Regardless of the mechanism the removal of the remaining cluster gas terminates the star formation process and determines the efficiency of star formation.

9.2. Numerical Models of Clustered Star Formation

Most numerical calculations to describe molecular cloud fragmentation and star cluster forma-
tion use SPH to solve the equations of hydrodynamics that govern the dynamical evolution of gaseous clouds. Owing to the stochastic nature of supersonic turbulence, it is not known in advance where and when local collapse occurs. SPH is the method of choice because it is fully Lagrangian. The fluid is represented by an ensemble of particles and flow quantities are obtained by averaging over an appropriate subset of the SPH particles (Benz 1990, Monaghan 1992). The method is able to resolve large density contrasts as particle are free to move and so naturally the particle concentration increases in high-density regions. In addition, one can introduce ‘sink’ particles into SPH (e.g. Bate et al. 1995), which have the ability to accrete gas from their surroundings, while keeping track of mass and linear and angular momentum. By adequately replacing high-density protostars in the centers of collapsing gas clumps with sink particles, one is able to follow the dynamical evolution of the system over many free-fall times. This is an essential ingredient for following the formation of stellar clusters.

The first attempts to numerically model the formation of star clusters date back to the late 1970’s (e.g. Larson 1978). With the rapid increase of computer power in recent years, more realistic calculations became possible. Whitworth et al. (1995) and Bhattacharjee et al. (1998) investigated in detail the fragmentation of shock-tube interfaces of colliding molecular clouds into small stellar systems. Klessen et al. (1998), and Klessen & Burkert (2000, 2001) studied the formation of stellar clusters from random Gaussian density fluctuations in molecular clouds. Models that include molecular cloud turbulence and consistently follow clustered star formation from turbulent fragmentation have been presented by Klessen et al. (2000), Klessen (2001a,b) and Bate et al. (2002). Focusing on the role of competitive accretion and neglecting the processes that lead to the formation of protostellar cores, Bonnell et al. (2001a,b) study the mass growth of randomly placed accretion particles in simplified model clouds. It should be mentioned, that the majority of numerical studies of interstellar turbulence and molecular cloud fragmentation are based on grid-based methods. These models are conceptually more difficult to combine with star cluster evolution calculations than the SPH models discussed here.

Star formation is an enormously complex process. It spans 20 orders of magnitude in density (from molecular cloud cores to the stellar interior) and 7 – 8 decades in spatial scale (compare the pc-size scales of molecular clouds with typical stellar radii of $\sim 10^{11}$ cm). And it involves a large number of physical processes. An adequate treatment of star formation must not only take into account gas dynamics and self-gravity, as most models do, but also include heating and cooling processes, radiation transfer, magnetic fields, chemical phase transitions and reactions, and feedback processes from star formation itself. Star formation very likely is a feedback regulated process. Bi-polar outflows from young protostars deposit energy and momentum in star forming regions, stellar winds heat the remaining gas, UV radiation from massive stars may completely ionize the cluster gas thus preventing further gas accretion and terminating star formation. Supernovae explosions (e.g. from very massive young stars, or from a nearby OB association) finally may disrupt molecular clouds altogether, preventing further star formation on scales of cloud as a whole. One needs to keep in mind, that most of these processes are not included in the models of star cluster formation. Some first attempts to model the effects of gas expulsion on the subsequent dynamical evolution of young stellar clusters are reported by Geyer & Burkert (2002) and also by Kroupa et al. (2001).

9.3. Initial Conditions from Star Formation Calculations

In principle, it appears straightforward to adopt the results of molecular cloud fragmentation and star forming calculations as starting conditions of star cluster evolution models. First, one needs to identify all protostars in the star formation model, then second, determine their masses $m$, positions $\vec{r}$, and velocities $\vec{v}$, maybe also their angular momentum (spin $\vec{j}$), and finally, supply this list to the stellar dynamics code. However, in practice the situation is not that simple. Before we realistically apply results from star formation models to star cluster evolution
we have to address several inconsistencies of the methods. The following gives a list of assumptions that need to be introduced to be able to combine both methods.

1. **Gas removal:** Because star forming calculations typically do not treat protostellar feedback and gas removal, the overall star formation efficiency is a *free parameter*. The physically motivated range roughly lies between 20% to 60%. For smaller values feedback processes are likely to be still too weak to significantly alter or halt star formation, and for larger values the collective effects of protostellar outflows, winds, and UV radiation from massive stars (in the case of massive clusters) will have modified the star forming cloud so dramatically that the simple gas laws adopted in most cluster forming calculations break down.

When using coordinates and velocities of protostars from cluster forming calculation as input for the subsequent cluster evolution, one usually makes the further assumption of instantaneous gas removal, because the stellar dynamics calculations typically neglect any contributions of gas to the cluster potential.

2. **Close binaries:** Most cluster formation calculations can only describe the formation of very wide binaries which essentially form by a capturing process when two gas clumps each containing one protostar merge together. Close binaries are likely to form from gravitational instabilities in protostellar accretion disks, which are not resolved by typical cloud fragmentation calculations.

A notable exception is the very high resolution calculation by Bate *et al.* (2002). If accretion disks are not resolved in a calculation, one needs to *assume* a close binary fraction and assign mass ratios and orbital eccentricities to each core in the simulation before forwarding this information to the cluster evolution code.

3. **Small $N$:** The current molecular cloud fragmentation calculations are at best able to describe the formation of clusters with about a hundred stars. This is insufficient for most stellar dynamics purposes. However, with further advances in computer technology and with improved parallel algorithms modeling of the formation of star clusters with some thousand member will become possible in the near future. One will be able to follow the evolution of star clusters like the Trapezium Cluster in Orion, or of the Pleiades, or Hyades fully ab initio. For larger stellar clusters there is no hope to consistently include star formation into the stellar dynamic calculations. One needs to resort to theoretical considerations as discussed in Section 9.4.

4. **Pre-main sequence evolution:** Very massive stars may enter the stellar main sequence while still accreting being deeply embedded in their parental gas cocoon. Low-mass protostars, on the contrary, spend a long time in the classical pre-main sequence contraction phase (e.g. Palla & Stahler 1999). For the MODEST approach this means that even long after gas expulsion, during first several million of star cluster evolution, stellar dynamics not only needs to be combined with stellar evolution modules for main sequence stars, but also pre-main sequence modeling needs to be included. This is the more important as pre-main sequence stars have considerably larger stellar radii than stars on the main sequence and therefore are more susceptible to collisional processes.

### 9.4. Initial Conditions from Theoretical Considerations

Whereas the initial conditions for star clusters with a small number of members $N$ can be motivated by star forming calculations as advocated before, this is not true for very massive star clusters. It is not possible to simply ‘scale up’ the properties of small-$N$ clusters into the large number regime. Therefore, initial conditions for large-$N$ clusters mostly will be obtained by searching for an appropriate distribution function $f$.

In the absence of any information from computer-generated models, we may distinguish
between theoretical equilibrium configurations and more realistic cluster models based on observations of star-forming regions.

At the simplest level, Plummer models are often used. However, these have no direct connection with dynamics and should therefore be considered as convenient models for test purposes.

Theoreticians often like to investigate families of well-defined models. A wide variety of equilibrium models can be described using King-Michie distributions. Adopting a distribution function of the type \( f(E, J_z) \), we can generate a sequence of models both in terms of the central concentration parameter and the amount of rotation. In addition, velocity anisotropy can also be considered.

Initial subclustering forms a good alternative to idealized distributions and may actually be more useful for realistic cluster simulations. The origin of runaway stars gives rise to one interesting set of problems posed by this scenario.

Given the initial coordinates and velocities, we also need to specify an initial mass function and there is a wide choice for the latter. For practical purposes, a piecewise fitting function based on observational data may be adopted. This still leaves the question of the upper mass limit which plays an important role. Initial mass segregation presents a further uncertainty, although there are some observational constraints.

Primordial binaries represent another important ingredient of star cluster simulations. The main parameters here can be summarized as \( f(a, e, m_1/m_2) \), where the semi-major axis \( a \) distribution needs to span many decades. The distribution of eccentricities \( e \) may be of secondary importance but the mass ratios \( m_1/m_2 \) have a direct bearing on the end-point of binary evolution. Likewise the upper mass limit affects the production of degenerate objects which are known to reside in clusters. It is also worth emphasizing that cluster evolution is speeded up significantly by the presence of a mass spectrum.

Finally, a non-equilibrium value for the initial virial ratio leads to violent relaxation and core-halo formation on a short time scale. Such models may be relevant in connection with removal of the remaining gas.

10. Data Structures and Formats

10.1. Exchanging Data

Communication among the various independent modules of a running program may be accomplished via simple functional (subroutine) interfaces which define and strictly control how much one module needs to know about the workings of another. It is equally desirable for separate programs to communicate in some standard way. One can easily imagine situations where we wish to compare the operations of, or simply share data between, two programs implementing alternative treatments of the same underlying physics. Examples might be Monte-Carlo and \( N \)-body treatments of stellar dynamics, SPH and recipe formulations of fluid mechanics, or different sets of recipes for stellar and binary evolution. For purposes of comparison and communication, it is essential that these programs all be able to interpret and manage the same input data sets.

Such a requirement immediately raises several significant technical problems. Simply put, different programs may generate very different kinds of data, organized internally in unique, even conflicting, ways, and possibly sampled inhomogeneously in space or time. For example, a 2-D stellar evolution code might produce as output a series of two-dimensional arrays representing various thermodynamic quantities at uniformly spaced sampling times, or the sampling intervals may be chosen so that the data tend to cluster around interesting evolutionary stages. A simple SPH simulation with shared time steps would generate identical data for all particles, sampled uniformly in time but non-uniformly in space. An \( N \)-body simulation with variable time steps naturally generates inhomogeneous data (different data for different particles), sampled non-uniformly in both space and time, often with some sort of hierarchical (tree) structure implicit in the data. In the grand simulations contemplated here, we must allow for the possibility of any and all of these data formats (and others!) being freely mixed in the I/O stream.

Some basic design considerations then are: (1) How do we accommodate a broad range of data formats in a flexible way? (2) Should we dis-
tistinguish between complete data streams used to reconstruct entire calculations and much simpler “snapshot” files used to checkpoint and restart simulations? (3) How much data should be saved in a file, and how much should be recomputed when the file is read? (4) How do we represent the data in space and in time? The choice leads to the interpretation of position and time as either array indices or particle attributes. We must be able to support both descriptions. (5) How do we represent particle attributes at each sampling point, however defined: as an array of physical quantities (homogeneous particle data), or as a collection of tagged properties (inhomogeneous data)? (6) Finally, since we expect to be dealing with very large amounts of data, how can we accomplish these goals efficiently? We discuss some examples of data formats (FITS, NEMO, Starlab/story, Starlab/tdyn,...), each of which addresses one or more some of the above points.

10.2. The tdyn Data Representation

A simple “snapshot” data format is adequate for many types of simulation. Systems with small dynamic range are easily synchronized (or are synchronous by construction), making it both straightforward and efficient to save data at regular predefined intervals. Not all calculations lend themselves to this approach, however. N-body simulations, for example, naturally produce data in a quite different format. Their large dynamic range means that individual particle time steps are the norm, meaning that particle trajectories are updated non-uniformly, each at the “right” rate, as defined by its own local time scales. A complete description of the dynamical evolution requires that we find a convenient way of saving and reproducing this level of detail in a data file.

Of course, one can easily produce snapshots of an N-body system for check-pointing and restart purposes. In practice, time steps are chosen to be powers of two, greatly improving scheduling efficiency by organizing particles into blocks which can be updated simultaneously. This also means that the system is necessarily synchronized at regular intervals, allowing snapshots to be produced. (Even without block time steps, synchronization can be forced at any desired output time, but block steps are widely used and facilitate the process.) However, for many purposes, such as analyzing particle motion or visualizing the evolving data set, it is desirable to reproduce trajectories in detail. Snapshots are poorly suited to this task, as they are severely limited by the dynamic range of the data, which generally makes interpolation impossible unless the interval between snapshots is made impractically small.

We have developed a data structure that allows us to save and manipulate enough detail to reproduce the native N-body structure without significant loss of resolution. In essence, rather than trying to sample the positions and velocities of stars at fixed, finely spaced intervals, we provide these data asynchronously, at key points along each orbit, then use this information to interpolate each trajectory to any specified time. As an analogy, if we wished to provide a means of specifying the locations of all trains within the New York subway system at any given time, we probably would not opt to publish a long list of all train positions on a second-by-second basis. Rather, it would be much more efficient to provide a timetable stating when trains arrive and depart from each station, together with some simple rule for computing a train’s movement en route from one station to the next. The description below is couched in the terminology of the Starlab environment within which this approach was developed. However, the basic ideas are common to all N-body codes.

In the simplest approach, we might save particle positions and velocities at the end of every N-body step. Then, to determine the particle’s position at any intermediate time, we use a fourth-order interpolation scheme (in fact, the same scheme used in the N-body code) to fit the position and velocity at each end of the stored interval spanning the desired time. In this way we can reconstruct a continuous, differentiable trajectory that reproduces the original N-body track. The choice of time step ensures that the sampling interval is adequate. As a practical matter, we find that sampling every step is unnecessary. Sub-sampling the trajectory—saving data only every 20–30 time steps—allows adequate re-

13See http://www.manybody.org/modest.html
production for most purposes, ensuring energy conservation to better than 0.1%. The resulting volume of saved data is large (about 100 bytes per particle per \(N\)-body time unit), but manageable given the proper tools, provided with Starlab.

As the simulation proceeds, particle data are saved in a more or less unstructured way, as follows. At the end of each chosen step, the system simply writes a self-contained record specifying particle ID, mass, position, velocity, and other properties (e.g. stellar evolutionary state) to the output stream. The task of reconstituting this “stream of consciousness” into a usable data structure is left entirely to the program reading and interpreting the data. As a practical matter, it is convenient to print complete snapshots of the system at regularly spaced synchronization times, typically separated by a few dynamical time scales. The complete external representation then consists of segments of data starting and ending with full snapshots defining the hierarchical tree structure of the entire \(N\)-body system, connected by asynchronous sequences of particle records spanning each trajectory. These data segments, typically a few tens megabytes in size for 10k particles, form the basic unit of external data.

Changes in tree structure resulting (for example) from binary formation and destruction are handled in a manner analogous to the segments forming the full data set. When two particles combine to form a binary, the event is signaled by terminating records marking the end of the individual particle trajectories, followed by a snapshot of the new binary tree (center of mass plus components) marking the start of a new trajectory family. The reverse occurs when a binary splits back into components. Thus all structural changes in the tree, large and small, are clearly delineated by “bookends” defining the old and new tree structures.

Internally, as the stored data are read in, they are assembled into a four-dimensional tree structure, mimicking the standard Starlab linked-list describing spatial structure at any given time, but with the added dimension of forward and backward pointers in time allowing navigation along a given trajectory. (The \(\text{dyn} \) in \(\text{tdyn}\) refers to the basic Starlab data structure; the \(t\) refers to time.) Determining a particle’s position at any given time then amounts to identifying its trajectory and the interval along it spanning the desired time, then interpolating from the saved position and velocity data to obtain the desired information. By construction, the tree structure at any instant is completely defined by the tree structure on the most recent snapshot along each particle’s trajectory.

We note that most non-dynamical data vary slowly, or may even be constant, across a given interval or segment, so simple linear interpolation is usually adequate. Finally, since step-by-step treatment of internal binary motion generally requires too much storage in this treatment, most binaries are treated as evolving “kepler” structures describing their slowly varying orbital elements. A binary is reconstituted by first interpolating its orbital structure, then locating its components in the interpolated orbit.

10.3. Mixed Particle Type Snapshots

Here we would like to argue that for data exchange (initial conditions, archiving, Virtual Observatory, etc.), one should use a simple, lowest common denominator description of how the data will be stored “offline”. This means leaving out the complexities of varying time in the dataset, and adopt the notion of a snapshot.

Thinking of a snapshot as a matrix, where the columns are particle attributes and the rows individual particles, a programmer still has the choice to write the matrix column- or row major wise. In addition, modern \(N\)-body codes often deploy a small number of different types of particles (e.g. pure gravity, SPH, stellar evolution etc.) which can also evolve one type into another and create new types as the system evolves. So we arrive at a description of a snapshot as a set of differently sized matrices, organized in the following hierarchy:

\[
\begin{align*}
\text{Attribute} & = \text{a named Quantity (can also be vector)} \\
\text{Particle} & = \text{set of Attributes} \\
\text{Family} & = \text{set of Particles (same set of Attributes)} \\
\text{Snapshot} & = \text{set of Families} \\
\text{Album} & = \text{set of Snapshots}
\end{align*}
\]

where at each level a number of “header” variables are needed to describe the items and the
Figure 3. A “PAT” cube, with for Na attributes in Np particles in Nt SnapShots. Shown here are 4 SnapShots, with each 4 different type of Families (dyn, hydro, no-name and star). Hierarchically: Attribute ∈ Particle ∈ Family ∈ SnapShot ∈ Album

lower level items.

At this stage we do not want to suggest a particular implementation for that data-format yet, as there are several possibilities, including possibly a new one. Ideally we probably would want a self-descriptive format, such as XML, NEMO’s dataformat is also of this nature, and one can equally well think of implementations in FITS (BINTABLE) and HDF, both of which have had previous proposals floating around. (Teuben 1995)

10.3.1. Naming Conventions

Apart from deciding the basic layout of the data, all header variables and columns need to get names on which everybody can agree and give the same meaning to. Instead of the usual Fortran unformatted I/O

we would envision some structured I/O routines, which could look as follows

CALL NBAWRITE (UNIT,’Position’,POS,NBODY,3)

10.3.2. Attribute (Column) Names

As an initial suggestion here are some potential names one could agree on that would give meaning to the associated data in a snapshot:

// SnapShot and Family header variables
int Npoint // 922372036854775808
int Ndim // 3 (number of dimensions)
int Nattributes // 3 (number of attributes)
string FamilyName // ‘disk’, ‘halo’, ‘gas’, ‘bulge’
string CoordSys // ‘cartesian-xyz’, ‘polar-rtp’
string CodeName // ‘arik’
string CodeAuthor // ‘Roald Teuben’
string CodeVersion // ‘3.0’
string CodeDate // ‘23-jun-2014’
string Hardware // ‘grape12’

// Particle Attributes (~vector means Ndim applies)
real-vector Pos[] // simple dynamics
real-vector Vel[]
real-vector Pos[]
real Mass[]
real-vector Acc[]
real Potential[]
real Density[]
real SPHEntropy[] // gas properties (SPH)
real SPHTemperature[] //
real SPHSmothingLength[] // needs SPH kernel type
real SPHDensity[] //
real SPHAcc[] // pressure gradient
int SPHnneib[] // number of neighbors
real Age[] // stellar evolution
real Temperature[]
real Metallicity[]
real Radius[]
real SemiMajorAxis[] // orbital elements
real Eccentricity[]
real Inclination[]
real LongAscendNode[]
real LongPeriapse[]
real TrueLong[]

11. Conclusions

Dynamical simulations of dense star clusters have reached the point where detailed treatments

\[\text{WRITE (UNIT) ((POS(K,J),K=1,3),J=1,NBODY)}\]

\[\text{\textsuperscript{14}See e.g. http://xml.gsfc.nasa.gov/XDF/XDF_home.html}\]

\[\text{\textsuperscript{15}see also http://zeus.ncsa.uiuc.edu:8080/data_format/data_format.html for a proposal by Bryan & Summers}\]
of many aspects of stellar physics must be included. A significant fraction of stars in globular clusters and galactic nuclei are expected to experience close encounters or actual physical collisions with other stars at some time during the evolution of their parent system. At the same time, collisions and the effects of stellar and binary evolution can strongly influence cluster dynamics, and may lead to the formation of objects whose properties provide key insights into a cluster’s past. Population synthesis studies have reached a similar conclusion from the opposite direction: dynamical interactions can be vitally important in determining the observed properties of dense stellar systems.

The dynamics of dense stellar systems is also essential for understanding star cluster formation. While protostars in a dense cluster environment build up, they are likely to interact strongly or even merge, and in general they will compete with each other for gas accretion. This has important consequences for the stellar mass spectrum and for the subsequent dynamical evolution of the cluster.

In the workshop MODEST-1 (for MOdeling DEnse STellar systems) the participants discussed many possible avenues for combining stellar physics with stellar dynamics. Options considered ranged from simple rules and heuristic recipes, to extensive look-up tables using precomputed data, to full-blown “live” simulations of stellar and binary evolution and stellar hydrodynamics embedded in a dynamical code. The following is a consensus view of the current state of the art and an assessment of feasible future developments in the various subfields represented at the meeting.

Dynamics. Traditionally, treatments of stellar and binary evolution and simple recipes for collisions have been realized as modules attached to existing dynamical integrators. In part, this is historical—dynamicists have had the most pressing reasons to incorporate these effects into their simulations. However, it is also a fairly natural way to proceed, as the dynamical portion of a large N-body calculation is usually also the part principally concerned with large-scale structure, scheduling, and the orchestration of “local” events, such as binary formation and destruction, stellar interactions, stellar evolution, and so on. One might imagine constructing a more democratic system in which the dynamics, stellar evolution, and hydrodynamics are handled on an equal footing. However it seems likely that, for the foreseeable future, the dynamical integrator will continue to provide the framework within which other physical effects are incorporated.

Evolution of isolated stars. For “canonical” stars that start their lives on the main sequence with more or less normal compositions and never experience close encounters with other stars, there seems to be no strong reason to perform on-the-fly computations of stellar evolution. Such calculations will almost certainly be of lower precision and contain less physics than existing published calculations. Rather, the most practical approach involves the use of look-up tables and fitting formulae based on precomputed tracks, essentially as already implemented in current N-body codes.

Evolution of isolated binaries. Binary evolution is too complex for live binary evolution programs, and is expected to remain so for the foreseeable future. No such programs currently exist, and even simplified versions would likely be too fragile for standalone use. The physics can be very sensitive to small perturbations and in many cases is not sufficiently well defined for encapsulation in a program to be possible; the number of binary configurations in which the detailed physics is simply unknown is depressingly large. For the same reasons, no definitive precomputed binary evolutionary tracks exist. The parameter space is probably too large for look-ups analogous to those used in stellar evolution to be practical in any case. We thus expect continued use of recipes and heuristic rules of increasing sophistication, again more or less as implemented in existing N-body codes. We note that this approach has the added benefit of allowing an investigator to identify and parametrize key binary properties, and to vary and study their effects in a controlled way.

Hydrodynamics. Some integrated treatment of stellar collisions is clearly required. Many collisions involving main-sequence stars can be ad-
equately handled by rules and recipes currently under development, but it seems inevitable that others will have to be performed on the fly, probably using SPH as the description of fluid dynamics best suited to incorporation into a dynamical integrator. Existing codes do not include such modules; most resort to (over)simplified “sticky” criteria for stellar mergers. Basic self-contained SPH (or shortcuts such as entropy-sorting) treatments of two-body collisions could in principle be added to existing codes in a relatively straightforward way. Integration of arbitrary stellar encounters within a full N-body environment is probably a feasible, but much longer-term, goal.

Collision Products. Collisions—either direct, between unbound stars, or indirect, resulting from binary evolution or temporary capture of stars in binaries—will give rise to “non-canonical” stars quite unlike those normally studied by stellar evolution codes or reported in the literature. They will be out of thermal equilibrium, will probably be rapidly rotating, and will have unusual composition and entropy profiles. We will not be able to precompute and interpolate all the possibilities. Here we really do need live stellar evolution codes to study the appearance and evolution of the collision products. However, such studies pose a severe challenge to existing techniques, and lie beyond the capabilities of current stellar evolution codes. The creation of a robust, standalone module to handle the evolution of collision products is a high priority.

How to make the pieces communicate? It is unrealistic to expect researchers to completely rewrite their codes (no matter how attractive such a prospect might be...) in order to merge them with other programs. Rather, it is better to create modular programs by encapsulating parts or all of existing computer codes and define robust interfaces specifying clearly the functionality of each module and the data that must be provided and returned for each to work. Such an approach is vital, as it will facilitate controlled comparison of competing techniques. Behind the interface, the structure of each module will be entirely up to the programmer, so long as it conforms rigorously to the agreed-upon interface specifications. We have begun a study of the interfaces, data structures, and communication protocols needed to realize this goal. The external representation of simulation data is also an important and unresolved issue—we need to share data between programs in an efficient, extendible, and non-destructive way.

The first MODEST workshop was successful in bringing together the three astrophysics communities of researchers working the fields of stellar evolution, stellar hydrodynamics, and stellar dynamics. We will continue to hold these workshops twice yearly, thereby providing a meeting point for those who are actively involved in simulating dense stellar systems. For further details, see the MODEST web site.\[16\]

Acknowledgments. We acknowledge the input of the participants of the MODEST-1 workshop. Here is the complete list of those who attended part or all of the workshop:


In addition, we also acknowledge comments on the manuscript by Douglas Heggie and Pavel Kroupa. R.S.K. acknowledges financial support by the Emmy Noether Program of the Deutsche Forschungsgemeinschaft (DFG, grant KL1358/1).

\[16\]http://www.manybody.org/modest.html
J.C.L. acknowledges support from NSF grant AST 00-71165. His work was also partly supported by the National Computational Science Alliance under grant AST 98-0014N and utilized the NCSA SGI/Cray Origin2000 parallel supercomputer. S.M. acknowledges support from NASA ATP grant NAG5-10775. R.F.W.'s participation is supported in part by NASA grant NAG 5-11016 to the University of Illinois.

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