Active Learning for Computationally Efficient Distribution of Binary Evolution Simulations

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Received 2022 March 29; revised 2022 August 2; accepted 2022 August 17; published 2022 October 13

Abstract

Binary stars undergo a variety of interactions and evolutionary phases, critical for predicting and explaining observations. Binary population synthesis with full simulation of stellar structure and evolution is computationally expensive, requiring a large number of mass-transfer sequences. The recently developed binary population synthesis code POSYDON incorporates grids of MESA binary star simulations that are interpolated to model large-scale populations of massive binaries. The traditional method of computing a high-density rectilinear grid of simulations is not scalable for higher-dimension grids, accounting for a range of metallicities, rotation, and eccentricity. We present a new active learning algorithm, psy-cris, which uses machine learning in the data-gathering process to adaptively and iteratively target simulations to run, resulting in a custom, high-performance training set. We test psy-cris on a toy problem and find the resulting training sets require fewer simulations for accurate classification and regression than either regular or randomly sampled grids. We further apply psy-cris to the target problem of building a dynamic grid of MESA simulations, and we demonstrate that, even without fine tuning, a simulation set of only ~1/4 the size of a rectilinear grid is sufficient to achieve the same classification accuracy. We anticipate further gains when algorithmic parameters are optimized for the targeted application. We find that optimizing for classification only may lead to performance losses in regression, and vice versa. Lowering the computational cost of producing grids will enable new population synthesis codes such as POSYDON to cover more input parameters while preserving interpolation accuracies.

Unified Astronomy Thesaurus concepts: Astronomical simulations (1857); Classification (1907); Regression (1914); Multiple star evolution (2153)

1. Introduction

Theoretical studies in astronomy often include simulations of physical processes that are not well constrained or understood with the purpose of making predictions and comparisons against observations. To account for model or physical uncertainties, simulations are carried out with various combinations of unconstrained parameters to explore the space of possible outcomes. However, in many cases, the computational cost per simulation is large and limits the exploration of the parameter space. For example, three-dimensional (3D) hydrodynamic supernova (SNe) simulations can take up to tens of millions of CPU-hours per simulation on current facilities (e.g., Müller et al. 2017; Vartanyan et al. 2019; Bollig et al. 2021). Large-scale cosmological simulations such as those from the Illustris project performed with the AREPO code (Springel 2010), demand similarly high computational costs (Nelson et al. 2018; Wang et al. 2021). Another area where large computational resources are expended is in modeling single and binary stars with one-dimensional (1D) stellar structure and evolution codes such as Modules for Experiments in Stellar Astrophysics (MESA; Paxton et al. 2011, 2013, 2015, 2018, 2019) especially when considering galactic populations. Large-scale computing is not only a significant investment of time and financial resources, but also comes with a carbon footprint; hence, optimization also minimizes our environmental impact (Portegies Zwart 2020). In this work, we aim to reduce the cost of producing data sets of binary evolution simulations for large populations through active learning (AL).

Binary population synthesis (hereafter population synthesis) codes simulate large numbers of binary star systems and their interactions to compare their statistics against observations (e.g., Han et al. 2020). While population synthesis has broad applicability, there are large computational hurdles to overcome when modeling binary populations. Not only does one have to model complex physical processes like mass transfer, SNe, and common envelope evolution, but one also needs to model large numbers (1010–1012) of systems since many astrophysically interesting phenomena are also rare (e.g., Kruckow et al. 2018; Breivik et al. 2020). Therefore, the computational cost to model each binary must be low to evolve a reasonably sized population with tens of millions of binary systems. Given a population synthesis code, the number of binaries that need to be simulated to understand an astrophysical population may be reduced by...
using targeted sampling methods (e.g., Andrews et al. 2018; Broekgaarden et al. 2019); however, there can still be considerable computational cost associated with evolving a binary accurately. To address this computational challenge, fitting formulae were developed to reproduce single star evolution (SSE) across a range of masses and metallicities (Hurley et al. 2000). The SSE equations were then combined with recipes to approximate the evolution of binary stars in the population synthesis code Binary Stellar Evolution (BSE; Hurley et al. 2002). This approach was computationally efficient enough for population studies, but came at the cost of physical approximations (e.g., simple parameterizations for the stability of mass transfer, lack of self-consistency in the treatment of stars out of thermal equilibrium and of stellar wind mass loss).

Since the development of BSE, similar rapid population synthesis codes have been developed using the same underlying methodology of relying on the SSE fitting formulae for single star evolution and different recipes to approximate binary evolution. These include SeBa (Portegies Zwart & Verbunt 1996; Toonen et al. 2012), Scenario Machine (Lipunov et al. 1996, 2009), StarTrack (Belczynski et al. 2002, 2008), binary c (Izzard et al. 2004, 2006, 2009), COMPAS (Stevenson et al. 2017; Barrett et al. 2018; Riley et al. 2022), MOBSE (Giacobbo et al. 2018), and COSMIC (Breivik et al. 2020). There are other approaches to population synthesis diverging from using SSE fitting formulae, aiming to increase the accuracy of physics being implemented. The ComBinE (Kruckow et al. 2018) and SEVN (Spera et al. 2015, 2019) codes use dense grids of single star evolutionary sequences and performs linear interpolation between models. BPASS provides more accurate modeling of stellar populations as it uses detailed binary models including their interactions although without simultaneously evolving both stars (Eldridge et al. 2017). Although stellar evolution codes such as MESA can evolve binary star systems self consistently, few population synthesis codes integrate detailed binary evolution models into their framework.

Population SYnthesis with Detailed binary evolution simulatIOns (POSYDON) is a new population synthesis code that uses full detailed binary evolution simulations evolved with MESA to perform population synthesis (Fragos et al. 2022). The 1D stellar evolution code MESA solves the equations of stellar structure and composition as a function of time, allowing MESA to self-consistently evolve two stars and their orbit, which is not done with the aforementioned population synthesis codes. A few binary effects that are important to model self consistently include the structural response to mass loss in each star, as well as angular momentum transport between the orbit and stellar rotations. Modeling binary systems with MESA provides greater physical accuracy than binary population synthesis modeling, but at a comparatively high computational cost.

Although a single 1D stellar evolution simulation may have a modest cost ranging from tens to hundreds of CPU-hours, it is common to compute thousands to tens of thousands of models due to the intrinsically large number of parameters that describe stellar and binary physics. (e.g., Fragos et al. 2015; Farmer et al. 2015; Choi et al. 2016; Misra et al. 2020; García et al. 2021; Marchant et al. 2021; Román-Garza et al. 2021; Gallegos-García et al. 2021). Although the aforementioned studies show that MESA simulations can be scaled to run large numbers of systems, they are still too computationally expensive to directly model galactic populations.

The POSYDON framework uses large data sets of precomputed binary evolution simulations to evolve a population through various phases of evolution. To approximate the evolution of a binary system for which no precomputed model exists, interpolation is employed. To maximize the interpolation accuracy, and the subsequent population synthesis, the detailed binary simulations must cover an adequate range of parameter space to resolve complex behavior. Since binary stellar evolution has many intrinsic input parameters (e.g., component masses \(M_1, M_2\), orbital period \(P_{\text{orb}}\), eccentricity \(e\), and metallicity \(Z\)), POSYDON requires orders of magnitude more models compared to the single star approach of previous codes.

A common procedure for building these data sets is to choose the most important parameters in the problem and create a dense regular (rectilinear) grid of simulations. Then, the density is generally chosen based upon a combination of physical intuition about the problem and manual inspection (e.g., Wellstein & Langer 1999; Nelson & Eggleton 2001; de Mink et al. 2007; Farmer et al. 2015; Misra et al. 2020; Marchant et al. 2021). However, POSYDON data sets of binary simulations will have prohibitively large computational cost to produce when we consider expanding the dimensionality of our data sets. For example, to construct a grid of detailed binary simulations with 10 points in each of five dimensions, assuming a typical computation time of 12 hr per simulation, it would cost 1.2 million CPU-hours to complete. Here we study how to overcome these computational bottlenecks to use detailed simulations in every major phase of binary evolution. We propose AL as a solution for building grids of binary simulations, at a fraction of the cost of current methods of sampling parameter space.

AL is the process of intelligently sampling points of interest to build a custom training data set for machine-learning (ML) algorithms (Settles 2009). AL is useful in situations where labeling instances (e.g., running a simulation, and human annotation of data) is expensive or the number of samples to label is large. Depending on the application, AL algorithms optimize for either classification or regression with applications including speech recognition, image classification, and information extraction (Kumar & Gupta 2020). In astronomy it has been proposed as a way to optimize spectroscopic follow-up for labeling SNe light curves (Ishida et al. 2019; Kennamer et al. 2020), as well as mitigating sample selection bias for photometric classification of stars (Richards et al. 2012). It has also been applied to multiple computational-astrophysics problems to build training sets for interpolation or increased performance over standard sampling methods (e.g., Solorio et al. 2005; Doctor et al. 2017; Daningburg & O’Shaughnessy 2022; Ristic et al. 2022). Here we apply AL to population synthesis.

In this paper we present an AL framework for dynamically constructing data sets of binary simulations to be used in population synthesis codes such as POSYDON. In Section 2 we introduce our new AL algorithm, psy-cris (PoSydon Classification and Regression Informed Sampling), and describe our implementation. Then we present two tests of psy-cris: first on a synthetic data set in Section 3 and then in a real-world scenario using MESA to run binary evolution simulations in Section 4. In Section 5 we discuss the results of
our tests. We find that psy-cris performs well when optimized on synthetic data, and we see promising performance when applying our method to construct a data set of MESA simulations. The psy-cris code is open source, and will be available as a module of POSYDON. The psy-cris algorithm is broadly applicable to other AL problems with both classification and regression data.

2. The psy-cris Algorithm

2.1. Defining the Problem

To describe the psy-cris algorithm, we first provide a description of the data set and the problem we are trying to solve in the context of binary population synthesis with POSYDON.

An individual simulation corresponds to a vector $\mathbf{S}(t) = [v_1(t), \ldots, v_n(t)]$ that evolves in time $t$ based on some physical model $\mathbf{p}$. We can write the evolution of a simulation from an initial state $\mathbf{S}_i$ to some final state $\mathbf{S}_f$ with our model $\mathbf{p}$ as $\mathbf{p}(\mathbf{S}_i) = \mathbf{S}_f$. Our task is to predict the outcomes of all simulations in a given simulation domain $D$, without manually running simulations at all points in the space due to the prohibitive computational cost.

In the case of POSYDON, $D$ would consist of parameters describing binary stellar evolution (e.g., component masses $M_1$ and $M_2$, orbital period $P_{\text{orb}}$, metallicity $Z$, stellar rotation $\omega_1$ and $\omega_2$, and eccentricity $e$). Our physical model $\mathbf{p}$ is MESA, evolving a binary system from some initial configuration to a final state. Then, $\mathbf{S}(t)$ is anything reported by, or post-processed from the MESA simulations, that changes over the evolution of a binary. For example, it could be a real number, like the mass of the star at time $t$, or it could be a classification, such as whether a star is a black hole (BH) or a neutron star. Finally, $\mathbf{S}(t)$ is the set of all of these output parameters for a single binary. In POSYDON, we use ML techniques to predict the outcomes of these MESA simulations to evolve a binary population through various phases of evolution.

It is helpful to split our domain into two categories: the set of labeled data $\mathcal{L}$, completed simulations with inputs and outputs, and unlabeled data $\mathcal{U}$, the space of possible simulations that have not been computed. To evolve simulations $\mathcal{S}_i \in \mathcal{U}$, we can use an ML model $\Gamma$ trained on a labeled data set $\mathcal{L}$, denoted as $\Gamma_L$. Then the predicted evolution of a simulation is given by:

$$\Gamma_L(\mathbf{S}_i) = \mathbf{S}_f^\mathcal{L},$$

(1)

where the star in $\mathbf{S}_f^\mathcal{L}$ denotes predicted final properties.

Many computational-astrophysics problems, the computational cost of evaluating $\Gamma_L(\mathbf{S}_i)$ is much less than $\mathbf{p}(\mathbf{S}_i)$. For example, getting a prediction from an ML algorithm trained on MESA data is much less expensive than running MESA. To be applicable in any scientific study, we must also ensure that our ML model has high accuracy, i.e., $\mathbf{S}_f^\mathcal{L} \approx \mathbf{S}_f$. However, we must also consider the cost of computing our training set $\mathcal{L}$ and then optimizing $\Gamma$, each of which is not always negligible (e.g., for training Gaussian processes; Rasmussen & Williams 2006).

Our study focuses on identifying an optimal $\mathcal{L}$, without knowing a priori what our simulations across $D$ may look like, at a minimized computational cost. Our AL algorithm psy-cris takes an initially sparse training data set $\mathcal{L}$, and identifies new points $\mathcal{S}_i \in \mathcal{U}$, such that when they are labeled ($\mathbf{p}(\mathcal{S}_i) = \mathbf{S}_f^\mathcal{L}$) and added to the training set for $\Gamma$ ($\mathcal{L} = \mathcal{L} \cup \mathcal{S}_i$), we achieve high accuracy ($\Gamma_L(\mathcal{S}_i) \approx \mathbf{p}(\mathcal{S}_i)$) with a low computational cost compared to standard methods of constructing the training set.

2.2. Algorithm and Workflow

AL algorithms take as input an initial training data set and output query points to be labeled by an oracle (Settles 2009). The oracle may be a human manually labeling data or the results of simulations that we treat as the truth (as in POSYDON). AL algorithms work in a loop with the oracle to iteratively build a training data set designed to achieve high accuracy in interpolation with an economical number of simulations. It is common for AL algorithms to be applied in either classification or regression problems (Kumar & Gupta 2020), but in POSYDON, our data naturally span both regimes. For example, one can classify binary simulations using their mass-transfer histories and perform regression on their continuous output quantities like final orbital separation, age, and component masses. Therefore, we designed psy-cris to consider both classification and regression simultaneously.

We achieve this by first combining AL heuristics, which are estimates for uncertainty in classification or regression (Section 2.3.2). Next, we sample from this new, combined AL heuristic and generate an uncertainty distribution (Section 2.3.3). Finally, we draw query points from the uncertainty distribution in serial or batches. This completes a single AL loop of psy-cris, which will repeat until a termination condition is met. This condition can simply be a maximum number of sampled points, or something more complex. The details of the psy-cris algorithm are described in Section 2.3, our exact implementation is described in Section 2.4, and our results from applying psy-cris are presented in Sections 3 and 4.

Figure 1 shows a schematic diagram of psy-cris integrated with POSYDON to run binary stellar evolution simulations with MESA. The green rectangle shows the scope of the psy-cris algorithm and how it integrates with other portions of POSYDON infrastructure. POSYDON runs binary simulations with MESA queried by psy-cris and then post-processes the output into our pre-defined classes and regression quantities. The blue square represents the oracle in an AL loop and can be replaced with other simulation software.

2.3. Components and Concepts

2.3.1. Classification and Regression AL Heuristics

Classification deals with categorical data while regression is used in cases where data are continuously valued. For example, we may classify a binary evolution simulation based on its mass-transfer history while a regression quantity may be the final mass of each component. Conceptually, a classification or regression AL heuristic is designed to locate unlabeled samples of interest based on a given ML method applied to $L$. If the ML algorithm has a measure of uncertainty in its predictions, or if one can be constructed, then this can be used as an AL heuristic. We denote a general classification or regression AL heuristic as $C$ and $R$, respectively.

2.3.2. Combining Classification and Regression AL Heuristics

We designed the psy-cris algorithm to optimize for both classification and regression simultaneously. To enable this coupled optimization, we propose that a linear combination of individual heuristics can also be used as a heuristic. We denote
our new heuristic as $F_{CR}$ defined as:

$$F_{CR}(x) = (\tau F_C(x) + (1 - \tau) F_R(x))^\beta$$  

(2)

where $x$ is a point in parameter space, $\tau \in [0, 1]$ sets the fractional contribution of the classification and regression terms, and $\beta$ controls the sharpness of the distribution (similarly to an inverse temperature; Mehrjou et al. 2018). The terms $F_C$ and $F_R$ are normalized heuristics. Since we combine $C$ and $R$ terms, their relative scaling matters.\(^8\) Equation (2) is designed to be high valued in regions with the largest estimated uncertainty in classification and regression. This is just one way to combine $C$ and $R$, but many other choices could be made.

### 2.3.3. Sampling Query Points

Once the heuristic has been defined, we must select query points for the oracle to label. AL algorithms can return either a single query point (serial) or a set of query points (batch) to label. Batch proposal schemes introduce more complexity, but allow multiple queries to be labeled in parallel. This may be less efficient in terms of the total number of simulations needed than ever picking the best point, but provides an advantage in terms of computational wall time (Schohn & Cohn 2000; Sener & Savarese 2017; Cai et al. 2017; Kennamer et al. 2020). With batch proposals, we may also consider more than just the maximum of the AL heuristic, but also the diversity of samples in the batch to avoid redundancy (King et al. 2004; Hoi et al. 2006; Demir et al. 2011; Wang et al. 2017).

In psy-cris, we adopt a flexible approach, which allows either serial or batch proposals, by creating an uncertainty distribution from which any number of query points can be drawn. We use our combined heuristic in Equation (2) as the target distribution for a parallel tempered Markov Chain Monte Carlo (PTMCMC) sampling algorithm (Swendsen & Wang 1986). PTMCMC (Earl & Deem 2005) combines multiple MCMC sampling chains at different temperatures (where the coldest temperature corresponds to the target distribution, and an infinite temperature to a uniform distribution), to improve the exploration of parameter space, which can be challenging for standard MCMCs if the target distribution is sharply peaked and multimodal. We assume that psy-cris may query any point in the input space, and is not constrained to a fixed or discrete pool of unlabeled samples. Drawing query points is the final step in one psy-cris iteration.

### 2.4. Implementation

#### 2.4.1. Classifiers & Regressors

We perform one-against-all binary classification. For data with $n$ classes (where $n > 2$), we train $n$ binary classifiers and define the predicted class $\hat{y}$ and classification probability $P(\hat{y}|x)$, respectively,

$$\hat{y} = \arg\max_{i=1,...,n}[\theta_i(x)],$$

(3)

$$P(\hat{y}|x) = \max_{i=1,...,n}[\theta_i(x)],$$

(4)

where $x$ is a query point in parameter space, and $\theta_i$ denotes a binary classifier for the $i$th class. The classifier $\theta_i(x)$, returns the probability that a query point $x$ corresponds to class $i$.

To construct our classification heuristic, we use a least confident measure for classification (Lewis & Gale 1994; Settles 2009; Kumar & Gupta 2020):

$$F_C(x) = \frac{n}{n - 1} \left(1 - P(\hat{y}|x)\right),$$

(5)

where $x$ is a query point, $n$ is the total number of classes in the data set, and $P(\hat{y}|x)$ is the classification probability in Equation (4). The prefactor is a pseudo-normalization term in the event all $n$ classes intersect at some point in the domain, which need not occur.

We train regression algorithms on data separated by class, taking into account both the different numbers of outputs and unique outputs per class. For $n$ classes that all have $m$ outputs, there are $n \times m$ interpolation algorithms trained.
For AL regression problems, approaches for estimating uncertainty include minimizing estimates of model variance or training multiple models to identify areas of disagreement. In psy-cris, we use a simple and computationally inexpensive measure: the average difference in the output between three nearest neighbors in the training set within the same class. We use the standard Euclidean metric to find nearest neighbors in parameter space. This is essentially a probe of the local function change in a given class. We calculate the average distance between $N$ nearest neighbors in the $i$th class as follows:

$$g^i_j(x) = \frac{1}{N} \sum_{k=1}^{N} |f^i_j(x) - f^i_j(x_k)|,$$

where $x \in L$ is a point in the labeled training set, $f^i_j(x)$ is the regression output in the $j$th output variable in that class, and $k$ iterates through the $N$ nearest neighbors in the class. Then we calculate Equation (6) for all points in the training set $L$.

Since Equation (6) can only be calculated at points in the training set, we interpolate between values of $g^i_m$ to give a continuous distribution for any unlabeled query point $x \in U$. This will be used during sampling where we must calculate Equation (2) at any point in parameter space.

In order to combine Equation (6) with a classification term, we pseudo-normalize it so that both terms are of the same order:

$$F_R(x) = A_1 \log_{10} \left[ A_0 \max_{j=1,...,m} [g^i_j(x)] + 1 \right].$$

The constant $A_0$ sets the scale of important absolute differences in the data set, while $A_1$ sets the scale of the function itself. We set $A_0 = 0.5$ and use $A_1$ as a normalization term, calculated by inverting the maximum value possible for Equation (7) across all classes, which is specific to the training data set. We take the maximum absolute difference across multiple regression outputs in the event that a class contains more than one. Numerically, Equation (7) is monotonically increasing, similar in form to the softplus activation function.

Our choices for $F_C$ and $F_R$ in our implementation of psy-cris are motivated by simplicity and are not to be taken as standard or optimal. For a more complete review of AL heuristics, see Settles (2009).

### 2.4.2. Sampling Method

After defining $F_C$ and $F_R$, we combine the two terms using Equation (2) to use as the target distribution for a PTMCMC sampling algorithm. Our implementation of PTMCMC uses one walker for each temperature, where each walker’s temperature is given by the relation $T_{i+1} = T_i^{1/c}$, where the spacing constant $c = 1.2$. The number of chains is then defined from the maximum temperature $T_{\text{max}}$ down to $T = 1$. We use the Metropolis-Hastings jump proposal scheme and take three steps before chain swap proposals (Metropolis et al. 1953; Hastings 1970). The stochastic sampling of points may help with exploring the parameter space, and discovering new regions with class boundaries (Joshi et al. 2009; Yang & Loog 2018).

The psy-cris algorithm generalizes for both serial and batch proposal schemes since any number of points can be drawn directly from the uncertainty distribution. Although this approach does not guarantee optimally spaced points within a batch, more complicated algorithms can be easily adopted into our formalism.

### 2.4.3. Handling Small Classes

It is possible for the psy-cris algorithm to discover classes by random chance, which were not in the original training set. Initially, new classes may contain only one sample. Our one-against-all binary classification scheme can handle this, but our regression algorithms are separately trained by class. Therefore, fitting most regression algorithms for a small class (one data point) is not well defined and fails. We could set the regression term $F_R$ to zero in Equation (2) if small classes are unimportant, but in our application, many astronomically interesting systems are also rare and may be subject to this edge case.

Therefore, to reflect the importance of small classes (and to avoid the numerical limitation of fitting small classes), we implement a separate proposal technique that supersedes the regular psy-cris algorithm in the event a small class is present in the training set. We draw points from a multivariate normal centered on the small class (one point) with a length scale set by the 10 nearest neighbors in input space (regardless of class) and scaled arbitrarily by 1/30, which was found to work well in tests. Then if the first proposal iteration does not populate the small class, subsequent iterations will tend to decrease the length scale. In practice, we would take care to construct our initial training set to contain an adequate sample of all relevant classes. The procedure described here is only necessary to test psy-cris in nonideal conditions.

### 2.5. Performance Metrics

To evaluate the performance of our algorithm, we compare training sets built with psy-cris to baseline configurations including regularly spaced grids and random uniformly distributed data. We use a validation set with known labels to determine how well classification and regression algorithms extrapolate after being trained on the different training sets.

For classification, we calculate the overall classification accuracy (CA), which we define as

$$CA = \frac{N_{\text{pred},C}}{N_{\text{total}}},$$

where $N_{\text{pred},C}$ is the number of correct predictions, and $N_{\text{total}}$ is the total number of queries.

We also use the per-class CA (PCA), which we define as

$$PCA = \frac{N_{\text{pred},C}^i}{N_{\text{total}}^i},$$

where $N_{\text{pred},C}^i$ is the total number of correct predictions for the $i$th class, and $N_{\text{total}}^i$ is the total number of points that truly belong to that class (true positives and false negatives). Our definition for PCA is consistent with what is often referred to as recall of a classifier. Although similar to CA, PCA can trivially approach 1 by overpredicting the classification region, which often happens for small training sets: PCA is sensitive to false negatives but insensitive to false positives.

We also calculate the F1-score, the harmonic mean of recall and precision, to compare with Equation (9), and find both PCA and the F1-score have qualitatively similar behavior for the tests we perform.
Figure 2. The number of training points needed to achieve a given classification (left panel) and regression (right panel) performance in our synthetic data set test. We define overall classification accuracy (CA) using Equation (8) and define regression accuracy (RA) as the 90th percentile of the distribution of absolute differences. We compare our AL algorithm psy-cris (solid lines) to regularly spaced grids (Grid), and random sampling (RS) while varying \( \tau \) and fixing \( \beta = 2 \). In both panels, high-performance data sets exist in the bottom-right corner, reaching high accuracy or low error with a small training set. Error bars in the combined regression error for Grid and RS are comparable to the size of their respective markers. The psy-cris configuration with \( \tau = 0.5 \) outperforms the regular grid and RS in both classification and regression simultaneously.

For regression accuracy (RA), we use the 90th percentile of the distribution of absolute differences between interpolation predictions and true values from the validation data set. In psy-cris, we train regression algorithms on data organized by class, so differences calculated at any point must first be classified to select the appropriate interpolator. Therefore, errors in classification can affect RA by requiring predictions in regression for points well outside the true classification region. We could consider calculating absolute differences only for true positive classifications, but in practice, we will not know the true outputs at every point. We choose to calculate absolute errors for points predicted to be in a given class, and only remove data if a class has no true regression data to compare against the prediction. We calculate a combined RA, where all absolute differences are combined into a single distribution regardless of class, and a per-class RA, where each class has its own absolute difference distribution.

For our regression algorithm, we use radial basis function (RBF) interpolation, which is parameterized by a length scale. By default, the length scale is set to the average distance between training data, which is assumed to be a good starting point. For the synthetic data test in Section 3, we set the length scale manually to the wavelength from Equation (A2), as it was found to achieve higher accuracies in small tests. For the MESA test in Section 4, we allow the default behavior, instead of explicitly attempting to fit for an optimal length scale.

3. Synthetic Data Test

3.1. Test Setup

To determine how psy-cris performs compared to standard methods of constructing training data sets, we first perform a test applying psy-cris on synthetic data. The use of synthetic data with known underlying distributions allows us to test the performance of the algorithm without resolution concerns. We construct a 3D synthetic data set including both classification and regression data drawn from analytic functions. The data contain six unique classes and one regression output, which is continuous across class boundaries. A detailed description of the synthetic data set, including the analytic functions and a visualization of the classification space, can be found in the Appendix.

Using the aforementioned synthetic data set, we start every AL run from a sparse, regularly spaced \( 5 \times 5 \times 5 \) grid. We let psy-cris iteratively query new points, which are labeled by the oracle in the standard AL loop, until reaching a combined total of 10,000 points in the final labeled data set. In addition to testing a default configuration of psy-cris, we run a suite of different models varying the contribution factor \( \tau \) and the sharpness parameter \( \beta \), as in Equation (2), to demonstrate their effect on our results. When evaluating performance, we compare our resulting training set from psy-cris \( L_{\text{AL}} \) to randomly distributed training data \( L_{\text{RS}} \) as a benchmark and regularly spaced training data \( L_{\text{Grid}} \). The test set for each configuration of training data is an overdense regular grid with \( 10^6 \) points.

3.2. Synthetic Data Test Summary

Figure 2 shows aggregate performance metrics comparing AL to random sampling (RS) and regularly spaced grids at various densities. Since the psy-cris algorithm contains inherent variability from PTMCMC and query point proposals, we show the average of 15 psy-cris runs where the solid lines are the mean, and shaded areas show 68th percentile contours. To simulate unique psy-cris runs starting from the same starting grid density, we vary the center of the starting grid.
randomly by ±1/2 the bin width in each dimension. This procedure is also used to determine the variance for regular grid configurations at different densities.

For overall CA (left panel of Figure 2), we find that psy-cris performs significantly better than RS and regularly spaced grids (Grid) when $\tau = 1$. As $\tau$ approaches 1, we find that each subsequent psy-cris configuration achieves higher CAs than the last (on the order of ~0.5%) or reaches a target accuracy with significantly smaller number of training points (e.g., for a CA = 96%, a factor of two reduction in $N_{\text{train}}$ between Grid and the $\tau = 0.75$ model).

For combined regression error (right panel), we see the opposite trend: $\tau$ approaching 0 gives optimal regression performance. However, the best-performing psy-cris configurations do not significantly outperform Grid or RS in regression, and most models converge after $\tau = 0.75$. This large-scale behavior when changing $\tau$ reflects our design principles when constructing the target distribution in Equation (2).

When considering classification and regression simultaneously with $\tau = 0.5$, psy-cris outperforms Grid and RS in classification, and performs similarly to Grid and RS in regression. Conversely, considering only classification ($\tau = 1$) or regression ($\tau = 0$) alone during AL may lead to significant losses in performance for the neglected category.

We may also consider performance on a class-by-class basis, allowing a more granular perspective on AL performance across the domain. In Figure 3 we show the PCA defined in Equation (9). We find the trends between models at different $\tau$ remain similar to those in overall classification, but some classes achieve larger (Class 5, $\tau = 0.5$) or smaller (Class 1, $\tau = 0.5$) gains from AL, which is not apparent from Figure 2. Class 5 presents a challenging classification problem because it contains an extended narrow region for $\tau$ in parameter space that is hard to see in the overall regression performance. For example, Class 5 with $\tau = 0.85$. In summary, psy-cris performs similarly to regularly spaced grids and RS even in the worst cases (e.g., Class 3 and Class 6) and can significantly outperform in others (e.g., Class 5).

We interpret the fact that Grid and RS perform similarly in regression performance to mean that our data can be fit easily with uniformly distributed training data. This is sensible given the regression function (Equation (A2)) is highly symmetric and smooth, which is not typical for real data.

We also ran models varying the sharpening parameter $\beta$ while fixing $\tau = 0.5$, and present aggregate performance results in Figure 5. We find that $\beta$ has a stronger effect on the classification performance than regression performance, and that a larger $\beta$ leads to higher CA. Although it is difficult to see from Figure 5, there appears to be a difference of ~1000 points in $N_{\text{train}}$ for a constant regression performance from $\beta = 2$ to $\beta = 3$. This suggests that $\beta = 2$ is close to the optimal value for this data set, and is likely to be a good starting place for problems with similar complexity.

4. Building a MESA Grid

4.1. MESA Test Setup

The primary goal for our AL algorithm is to construct an optimal set of detailed binary evolution simulations for use in population synthesis codes, such as the MESA simulations being used in POSYDON. An optimal training set provides a target CA and RA at the lowest computational cost (lowest number of simulations) possible. In this test, we demonstrate psy-cris working with POSYDON infrastructure to propose and label MESA simulations (as in Figure 1). We use a Python implementation of message passing interface (MPI), combining the software used to run MESA in POSYDON with psy-cris. Implementing distributed computing is critical, as it acts as the oracle in the AL loop, automatically running new MESA simulations and post-processing their output.

We model binary systems with a helium (He) main-sequence star and BH companion using MESA’s binary module. We use POSYDON default MESA controls and input parameters, which are described in detail in the POSYDON instrument paper (Fragos et al. 2022) with an additional change. We set the maximum radiative opacity to 0.5 cm$^2$ g$^{-1}$ for all MESA simulations to alleviate numerical convergence issues as described in Fragos et al. (2022). Our parameter space is the same as the grid of He-rich stars with compact object (CO) companions from Fragos et al. (2022), which covers the initial masses $M_1$ of $[0.5M_\odot, 80M_\odot]$, $M_2$ of $[1M_\odot, 35.88M_\odot]$, and the initial orbital period $P_{\text{orb}}$ of $[0.02$ day, $1117.2$ day].

Part of the POSYDON infrastructure includes parsing MESA outputs to categorize simulations into one of five classes, used to organize data for interpolation (Fragos et al. 2022). The classes are based upon a binary’s characteristic evolution: initial_MT (Roche-lobe overflow at zero-age main-sequence), stable_MT (dynamically stable Roche-lobe overflow mass transfer), unstable_MT (dynamically unstable Roche-lobe overflow mass transfer), no_MT (no Roche-lobe overflow mass transfer), and not_converged (numerical error or exceeded maximum computation time of 2 days). Some systems labeled as unstable_MT can start their mass-transfer sequences stably but eventually evolve to become unstable. Any binary sequence labeled as stable_MT remains stable throughout the entire evolution. Numerical convergence
issues arise in MESA due to limitations in modeling and often occur in specific regions in parameter space. In the AL phase, we use all five interpolation classes (including the not_converged systems) and only consider regression for the final orbital period for the stable_MT and unstable_MT classes. The not_converged class was included to keep psy-cris from continuously proposing simulations in these problematic regions since any failed run would not add information to the training set, wasting computing resources. We also log-normalize the inputs and outputs from $[-1, 1]$ before entering psy-cris and transform the proposal points back into linear space before evolving the system with MESA.

Our parameter space for this test is 3D but can be extended into a higher-dimensional problem, including covering multiple metallicities to give one example. The design of psy-cris is generalized to higher dimensions and can readily be applied in such cases. We compute a high-density regular grid with dimensions $41 \times 21 \times 49$ in $\log_{10}(M_1/M_\odot)$, $\log_{10}(M_2/M_\odot)$, $\log_{10}(P_{\text{orb}}/\text{day})$, respectively, totaling 42,113 successful MESA simulations (76 were unable to be parsed due to corrupt output files). We choose these dimensions for the regular grid by inspection based on previous tests in the same parameter space. We start psy-cris from a subset of the full-density grid by taking every other point in $M_1$ and $M_2$ and every fourth point in $P_{\text{orb}}$, retaining end points such that the range in each axis matches the full grid. We chose a different sampling for the orbital period because this dimension has a higher resolution because of the large classification changes that occur with small changes in $P_{\text{orb}}$. This procedure creates the starting, low-density grid with 3003 MESA simulations. To create the medium-density grid (which we use to compare performance in Section 4.2), we again use a subset of the full-density regular grid, taking every other point in $P_{\text{orb}}$, creating a grid with 21,525 MESA simulations.

During the beginning of AL, the parent MPI process runs one psy-cris iteration, which is used to propose the initial parameters for all child processes waiting to run MESA. Once any child process finishes running MESA, the parent process runs psy-cris again with the updated data set and proposes a new point in parameter space to run with MESA (Figure 1). This process is effectively a serial proposal scheme after the initial startup phase, although it depends on the rate at which MESA simulations finish. In our test, psy-cris terminates when a maximum number of proposed points is reached. We stop this test when we could see trends in our performance metrics since this demonstration is computationally expensive. In practice, psy-cris is meant to be used for grid sampling until a threshold accuracy in classification, regression, or a
increases in classification performance when using AL compared to RS and regular grids while varying \( \tau \) and fixing \( \beta = 2 \). We find that for some classes, our best-performing configurations of psy-cris slightly outperform Grid and RS. In most cases, the best-performing configuration of psy-cris is not \( \tau = 0 \) (which we might expect after seeing the trends in Figure 3), but an intermediate value of \( \tau \), which itself is not consistent across classes. Class 5 shows significant gains in regression performance compared to Grid and RS, but with the best-performing psy-cris configurations focusing more on classification (\( \tau > 0.5 \)). However, this behavior is reasonable in cases where classification is challenging, and we see Class 5 has the greatest increases in classification performance when using AL compared to RS and regular grids (Figure 3). The errors for grid points are comparable to the size of the markers for all classes. The errors for RS points are similarly small except for Class 2 and Class 4, which exhibit errors on the order of the \( \tau = 0 \) line.

4.2. MESA Test Summary

In our MESA test, we demonstrate how a fiducial configuration of psy-cris with parameters taken from the synthetic data tests with \( \tau = 0.5 \) and \( \beta = 2 \).

Unlike with the synthetic data set, the exact outcomes of all MESA simulations in our simulation domain are not known exactly. In order to calculate our standard performance metrics, we create a test set by running 10,000 random uniformly distributed MESA simulations in \( \log_{10}(M/M_\odot) \), \( \log_{10}(M_2/M_\odot) \), and \( \log_{10}(P_{\text{orb}}/\text{day}) \), which are kept separate from the regular grid and AL data sets until performance evaluation.

In Figure 4 we show the total number of training points needed to achieve a given regression performance (defined as the 90th percentile of the distribution of absolute differences) in our synthetic data set, separated by class. We compare our AL algorithm psy-cris (solid lines) to regularly spaced grids (Grid), and RS while varying \( \tau \) and fixing \( \beta = 2 \).

In Figure 6 we show the overall classification and regression performance between psy-cris and a regular grid for our BH–helium-main-sequence binary simulations. With psy-cris, we achieve overall CA comparable to the densest regular grid with \( \sim 4 \) times fewer MESA simulations. We also show the combined regression error for the final orbital period \( P_{\text{orb}} \) only, because this is the only quantity we use when considering regression during AL. We see that psy-cris reaches errors of \( \sim 5 \) day compared to \( \sim 2.4 \) day for the medium-density regular grid. While we see that the RA is decreasing, we do not have enough data to identify a strong trend that can be extrapolated to high \( N_{\text{train}} \). We assess that the variability seen in combined (and per-class) regression error is caused by our RBF interpolation, which sets the length scale as the average distance between training data. We do not optimize for nor fix the length scale since this is only a demonstration of psy-cris. Overall, the performance is consistent with expectations, but to gain a more complete picture, we consider per-class performance for both classification and regression.

In Figure 7 we show the PCA for our MESA test. The stable- and unstable-mass-transfer classes have the lowest accuracies while the no-mass-transfer class starts at nearly 98% accuracy. This is due to the relative size of classes in the parameter space, where small classes generally have lower performance. The unstable-mass-transfer class has the lowest accuracy and sees the largest gains from AL, similar to the challenging case of Class 5 in the synthetic data set. We find that all classes achieve better or comparable PCA with psy-cris at a lower computational cost than the medium-density grid. The reduction in number of points required for psy-cris to achieve a comparable per-class accuracy to the medium-density regular
and terms, and vary the sharpening parameter \( \beta \) as in Equation (2). Increasing \( \beta \) results in a higher CA while not significantly affecting the regression performance. However, at \( \beta = 3 \) we see the regression performance drop slightly. The right choice of \( \beta \) will depend on the data set as well as the classification and regression algorithms implemented.

In Figure 8 we show the absolute error in final orbital period in days for the stable-, unstable-, and no-mass-transfer classes. During AL, we ignore regression for systems undergoing mass transferring at zero-age main sequence (initial MT) because we expect these systems to result in stellar mergers early in their evolution, in which case their final orbital period is undefined. We also ignore regression for systems that do not have Roche-lobe overflow mass transfer (no_MT) as we assume these systems have comparatively simple evolution, which we are less interested in resolving in this demonstration psy-cris.

The unstable-mass-transfer class sees the greatest improvements in RA, achieving comparable performance to the full-density regular grid with 5.8 times fewer \texttt{MESA} simulations. This large increase in performance is driven by the fact that the unstable-mass-transfer class is least represented in the data (see Table 2), and AL necessarily populates underrepresented classes (Ertekin et al. 2007). Just as in the synthetic data test, we see that each class has different characteristic performances in regression on the final orbital period; the stable-mass-transfer class has errors on the order of days compared to tens of days for the unstable-mass-transfer class. The no-mass-transfer class does not achieve any significant improvements in regression, having a nearly constant performance throughout the duration of the AL phase. This is consistent with trends from the synthetic data test where disregarding regression during AL impacts the AL phase. This is consistent with trends from the synthetic data test where disregarding regression during AL impacts the AL phase. This is consistent with trends from the synthetic data test where disregarding regression during AL impacts the AL phase. This is consistent with trends from the synthetic data test where disregarding regression during AL impacts the AL phase. This is consistent with trends from the synthetic data test where disregarding regression during AL impacts the AL phase.

Finally, Figure 9 shows a slice of our parameter space visualizing the predicted classification after training on the full-density regular grid and psy-cris. The left panel shows the full-density regular grid, which is missing about five simulations at \( \log_{10}(P_{orb}/\text{day}) \sim -0.75 \) due to errors parsing the \texttt{MESA} simulations. The starting grid for \texttt{psy-cris} is a subset of the full-density regular grid and contains 3003 points. With AL, we resolve classification boundaries and even discover a new region of unstable-mass-transfer systems at \( \log_{10}(P_{orb}/\text{day}) \sim 2.5 \), all while requiring \( \leq 50\% \) the total number of training points of the regular grid. This classification performance has been achieved without optimizing \texttt{psy-cris} for this data set.

5. Discussion

We performed two tests of our AL algorithm \texttt{psy-cris}\texttt{.} on a synthetic data set as well as a real-world application of creating a grid of \texttt{MESA} simulations.

In the synthetic data test, we find that \texttt{psy-cris} can outperform standard methods of sampling in classification and regression by changing the primary free parameters in our algorithm. We also find that only focusing on classification or regression during AL may significantly hinder performance in the neglected category. We then use a fiducial configuration of \texttt{psy-cris} taken from the synthetic data test to construct a dynamic grid of \texttt{MESA} simulations. In this demonstration, we find \texttt{psy-cris}, while not optimized for this data set, is able to achieve comparable overall CA to the full-density regular grid with a factor of \( \sim 4 \) fewer simulations. We also see significant improvements in per-class regression performance for the unstable-mass-transfer class, achieving comparable performance to the full-density regular grid with a factor of \( \sim 5.8 \) fewer simulations. However, not all classes see large improvements in performance (e.g., stable-mass-transfer class in regression), so proper optimization of \texttt{psy-cris} is still necessary and will be explored in future work.

The \texttt{psy-cris} algorithm shows promising results for use with \texttt{MESA} grids and is currently being optimized for the \texttt{POLYDON} project. For use in other problems, there are several considerations one should take into account. We use linear interpolation and RBF interpolation from \texttt{scipy} (Virtanen et al. 2020) to perform classification and regression, respectively.
also use the same interpolation schemes when evaluating all performance metrics for classification and regression. There is a free parameter in the RBF interpolation, the length scale, which by default is set by the average distance between training data. We do not attempt to optimize the length scale in the MESA test, which is a naive way to perform regression, and we suspect that this causes poor regression performance in general. In all of our tests, we use the same interpolation and classification scheme across all classes, but in practice, a more fine-tuned approach is expected to achieve higher accuracies. A more targeted AL regression heuristic may also contribute better performance than shown here.

Concerning AL performance metrics themselves, there are also caveats one should consider. Although we present common performance metrics in this study, there are multiple ways to characterize performance where each metric may highlight specific features while hiding others. For example, our per-class classification definition trivially approaches unity for overconverged classificers. Imbalanced data sets also present challenges since minority systems may be neglected without affecting cumulative performance metrics significantly (Attenberg & Ertekin 2013). Therefore, it is important that we explore other performance metrics for AL in POSYDON as we further optimize psy-cris.

It is impossible to guarantee that our AL algorithm, like any AL algorithm, will always outperform standard sampling (e.g.,

| Class          | PCA  | F1-score |
|----------------|------|----------|
| initial_MT     | 4.2  | 4.2      |
| no_MT          | 4.9  | 3.7      |
| stable_MT      | 2.1  | 3.3      |
| unstable_MT    | 2.5  | 4.2      |

Note. This is simply $N_{\text{grid}}/N_{\text{AL}}$ at a fixed PCA (or F1-score). We calculate F1-score as a secondary metric to compare with our PCA, with both metrics showing similar reduction factors. All psy-cris curves except for unstable_MT for PCA, and initial_MT for F1-score, achieve comparable accuracy to the highest-density regular grid by the end of AL to within <0.5%.
three dimensions, matching the accuracy with AL to high-density rectilinear grids. Factors of four reduction in the number of simulations to achieve results.

The starting density of this grid impacts the future evolution of AL sampling, as every subsequent query point is based upon the information present in the initial training data set. For example, if a class does not exist in the initial training set because the starting grid is too sparse, it may never be discovered. Characterizing all possible outcomes in the parameter space is a problem faced by any discrete sampling method (e.g., Andrews et al. 2018; Broekgaarden et al. 2019), including regular grids and AL. However, in practice, we can use our domain knowledge of binary stellar evolution to check that our initial starting grid covers a reasonable range of physical outcomes, which is what is already done with regular grids at arbitrary densities.

6. Conclusions

Our aim is to efficiently construct data sets of binary evolution simulations with reliable interpolation accuracy to be used in the population synthesis codes such as POSYDON. Regular grids of binary evolution simulations can require prohibitively large computational resources to construct due to the cost per simulation combined with the parameter space we must cover to adequately characterize binary stellar evolution. We propose AL as a solution for constructing high-performance training sets of binary evolution simulations, at a reduced cost compared to standard methods. We demonstrate our new AL algorithm psy-cris tested on a synthetic data set and a realistic data set of binary evolution simulations with MESA.

In our initial synthetic data set test, we find that psy-cris can be optimized to outperform standard methods of sampling by varying the free parameters in our model. We find that focusing just on classification or regression alone may lead to significant losses in performance in the neglected category. Therefore it is essential to consider both to obtain well-adapted results. When considering per-class performance, we see major gains from AL in both classification and regression for classes, which present challenging classification problems, characterized by extended and narrow shapes in parameter space (Class 5). We also see low performance occurs for a minority classes in imbalanced data sets, as well as a class’s presence (or absence) in the initial training set (Class 2). psy-cris does indirectly address data imbalances by weighting each class equally (Table 2), but a more active approach may achieve better results. Different classes will have different characteristic performances based on their size and shape in parameter space, and may also benefit the most from AL in challenging cases.

We also demonstrate psy-cris working with POSYDON infrastructure to evolve binary systems consisting of a CO and
helium-main-sequence star with MESA. We use an unoptimized configuration of psy-cris, determined from the synthetic data set tests, and find that psy-cris outperforms the highest-density regular grid in overall classification using a quarter the total number of training points. In regression we see psy-cris is reducing the regression error but not yet to the level of the medium-density regular grid.

Creating the oracle in our AL loop was critically important for running simulations with MESA. Not only does the oracle label simulations automatically, but it ensures that we encounter fewer numerical difficulties that commonly arise in MESA. In our MESA test, 14% of our AL data set consists of failed simulations, which do not terminate normally due to numerical convergence issues (Table 2). Minimizing numerical convergence issues in our binary simulations with MESA will only further improve the AL results in POSYDON.

Since each POSYDON data set will exhibit unique classification and regression challenges, psy-cris may benefit from further optimization specific to each binary evolution grid. For instance, one could consider different AL metrics such as the classification entropy (e.g., Gal et al. 2017; Cai et al. 2017; Yang & Loog 2018). Currently, psy-cris has been designed to consider initial–final quantities alone, but our MESA simulations have time series evolution in between these end points. Therefore, in the future we are interested in seeing how psy-cris will respond to the complexity of their time series evolution as a metric for simulation proposals. Furthermore, we could factor in the varying computing cost of different simulations to pick the most informative points in parameter space per unit CPU time (Tomanek & Hahn 2010; Kumar & Gupta 2020).

Although we have designed it to optimize our generation of binary evolution grids as part of the POSYDON code, psy-cris is a general algorithm that can be applied to data sets with classification and regression outputs. To be applied directly in another similar problem, one must construct the oracle (which will label and run new simulations) to completely automate the AL loop. The psy-cris AL approach will work best for optimizing large grids of simulations where the number of simulated points is too large to optimize manually, and brute-force oversampling is too computationally expensive. Examples are numerical relativity simulations for building gravitational-wave approximants (Varma et al. 2019; Healy & Lousto 2022), or large grids of kilonova models (Ristic et al. 2022). The psy-cris code is open source, and exists as a module in POSYDON, which will be available in the next release of POSYDON.

We thank Monica Gallegos-Garcia for useful conversation during the development of psy-cris. K.A.R. is supported by the Gordon and Betty Moore Foundation (PI Kalogera, grant award GBMF8477). K.A.R. also thanks the LSSTC Data Science Fellowship Program, which is funded by LSSTC, NSF Cybertraining grant No. 1829740, the Brinson Foundation, and the Moore Foundation; their participation in the program has benefited this work. J.J.A. acknowledges support from CIERA and Northwestern University through a Postdoctoral Fellowship. P.M. acknowledges support from the FWO junior postdoctoral fellowship No. 12ZY520N. C.P.L.B. and Z.D. acknowledge support from the CIERA Board of Visitors Research Professorship. V.K. was partially supported through a CIFAR Senior Fellowship and a Guggenheim Fellowship. S.B., T.F., K.K., D.M., Z.X., and E.Z. were supported by a Swiss National Science Foundation Professorship grant (PI Fragos, 2022).
project No. PP00P2 176868). K.K. was partially supported by the Federal Commission for Scholarships for Foreign Students for the Swiss Government Excellence Scholarship (ESKAS No. 2021.0277). Z.X. was supported by the Chinese Scholarship Council (CSC). The computations were performed at Northwestern University on the Trident computer cluster (funded by the GBMF8477 award). This research was supported in part through the computational resources and staff contributions provided for the Quest high-performance computing facility at Northwestern University, which is jointly supported by the Office of the Provost, the Office for Research, and Northwestern University Information Technology.

Software: NumPy (van der Walt et al. 2011), SciPy (Virtanen et al. 2020), matplotlib (Hunter 2007), pandas (McKinney 2010), scikit-learn, (Pedregosa et al. 2011), POSYDON (Fragos et al. 2022).

Appendix
Constructing the Synthetic Data Set

In Section 3 we test our AL algorithm psy-cris in a 3D synthetic data set by comparing classification and regression performance of training sets built with psy-cris to regularly spaced grids and random sampling. We designed the classification space to contain multiple challenging classification features including a spatially small class, as well as extended and narrow classes throughout the parameter space. Our choices were also inspired in part by visual inspection of the binary evolution data sets in Fragos et al. (2022). We provide a pseudo-code function for the classification of an arbitrary point in our domain,

```python
def classification(x,y,z):
    if psi_5(x,y,z) < 0.3:
        return "Class 2"
    elif psi_4(x,y,z) > 1.8 and z < 0:
        return "Class 4"
    elif psi_1(x,y,z) > 1 and psi_2(x,y,z) > 2.5:
        return "Class 1"
    elif psi_2(x,y,z) > 2.5:
        return "Class 6"
    elif psi_3(x,y,z) > -0.5:
        return "Class 3"
    else:
        return "Class 5"
```

where methods psi_1 through psi_5 correspond to the functions in Equation (A1a). Finally, we have the regression function $\psi_6$, which is a 3D sinusoid, spherically symmetric about the origin. We chose to use the same function for all classes even though psy-cris treats each class’s regression output separately.

\[
\psi_6(x, y, z) = \sin(8\pi \sqrt{x^2 + y^2 + z^2}/3). \tag{A2}
\]

Figure 10. The true classification space for the synthetic data set, constructed with the analytic functions provided in Equation (A1a). The regression output from Equation (A2) is defined in the same range, and together they make up the synthetic data set.
