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1 Introduction

1.1 What is MilkyWay@home?

At best, "MilkyWay@home" is a nebulous term that can refer to several different projects, softwares, and/or groups of people. At its core, MilkyWay@home is a crowd-sourced supercomputer that takes volunteer computing time to perform complex and time-consuming calculations that are designed to improve our understanding of the Milky Way Galaxy. At the moment, this includes both the MilkyWay@home Separation application and the MilkyWay@home N-body application. The MilkyWay@home project was developed at Rensselaer Polytechnic Institute under the direction of Dr. Heidi Jo Newberg, and has utilized countless hours of work from many graduate students and researchers. For more information regarding Milky-Way@home, I direct the user to the MilkyWay forums at https://milkyway.cs.rpi.edu/milkyway/. All of the MilkyWay@home code is publicly available at https://github.com/Milkyway-at-home.

While the Separation application is an important part of MilkyWay@home, the Separation application is difficult to apply to other projects. Additionally, the tools that I have developed for the Separation application are quite situation specific and not designed with user friendliness/adaptability in mind. For this reason, *mwahpy* does not include functionality for the Separation application. For Separation application tools and documentation, see the Separation application github page at https://github.com/Milkyway-at-home/milkywayathome_client/tree/master/separation.

What I refer to as "MilkyWay@home" in this document is actually the MilkyWay@home N-body software. This software was originally designed to generate model dwarf galaxies in the Milky Way gravitational potential and integrate them forwards in time. This evolved dwarf galaxy would then be compared to observations of stellar streams in the sky in order to determine a quality of fit. By optimizing over the parameters that were used to generate the dwarf galaxy, one could in theory determine the parameters of the progenitor dwarf galaxy. This could even be done for streams with no obvious progenitor (e.g. the Orphan Stream).

It was realized shortly thereafter that the N-body application could be used to integrate the orbits of any bodies in the Milky Way, not just those that were generated as part of a dwarf galaxy. The ability to insert list of bodies to integrate forwards in time was provided. Additionally, the ability to control the size of the timestep of the simulation, the orbit of a dwarf galaxy progenitor, the timescale of the simulation, the dwarf galaxy parameters, the underlying gravitational potential, and other MilkyWay@home functionality made it a versatile and robust tool for N-body integration.

While the MilkyWay@home N-body application is currently used by many graduate and undergraduate students at Rensselaer Polytechnic Institute, the MilkyWay@home team is making an effort to encourage more widespread usage of the MilkyWay@home N-body software. Our goal is to provide a user-friendly, comprehensive experience for N-body integration. This will decrease the time needed to get new students and researchers up to speed with N-body software, which has become commonplace in the dynamical astrophysics community.

We expect that the MilkyWay@home N-body application will become more widely used, and that proper documentation and accompanying tools will quickly become necessary. Additionally, new applications of the N-body software have expanded simulations for upwards of one million bodies. Analysis of these large simulations requires quick, robust software. It is for these reasons that mwahpy was developed.

1.2 What is mwahpy?

The MilkyWay@home N-body software has been used by the Galactic dynamics research group at Rensselaer Polytechnic Institute for years. As each new student comes into contact with the software, each one has had to develop their own tools for analyzing the outputs of the software. Each individual software has its own set of bugs, sets the student back a few weeks while developing their own tools, and by not using a standardized set of analysis software we open up the group to problems with sharing data and code. Additionally, MilkyWay@home is written in C, a language which can be daunting to the typical undergraduate student (and is often unecessarily unwieldly if you are just trying to cut data or make a plot). My goal to alleviate these issues was a compilation of the tools that I had built in python and tested over my tenure as an undergraduate (and improved during my time as a graduate student) in the MilkyWay@home dynamics group. This became the mwahpy package.

Before we talk about what mwahpy is, it is good to clarify what mwahpy is not. This package is not able to do N-body integration. Things like generating dwarf galaxies, running simulations, and performing routines on the MilkyWay@home supercomputer are to be delegated to the proper MilkyWay@home software. Additionally, I would like to express that because the Separation application is interacted with by many fewer people than the N-body application, mwahpy will not support Separation application functionality.

There is plenty that mwahpy is meant to be used for, though. At its very base, mwahpy is a python package that is designed to easily and quickly read in N-body data output from the MilkyWay@home N-body software. Once read in, it is fairly simple to cut the data, plot the data, and save the data in a variety of formats. This package is also able to output data in a format that is readable by the MilkyWay@home N-body software. The software will automatically calculate complicated values such as proper motion and energies as needed, instead of computing everything up front. The major benefit of these routines is that new users can be confident that the values that are produced by mwahpy have been tested over several years and are known to be accurate.

There are several associated auxiliary packages in *mwahpy* as well, such as the coordinate transformation subpackage and the orbit fitting subpackages. These are provided in the package because while working with N-body simulations, I often ran into situations where I used the functionality provided in the auxiliary subpackages. The auxiliary subpackages are less streamlined and complete compared to the main subpackages, but are still fairly well tested and can be trusted.

The code for *mwahpy* is publically available and can be found at https://github.com/thomasdonlon/mwahpy. Not only is collaboration on the code allowed, it is encouraged! If you would like functionality added to *mwahpy* or if you find any bugs in the code, you can either leave a comment on the github page or you can write the code yourself and make a pull request. This code is still being actively maintained, and I plan on eventually getting around to any bug fixes or desired functionality that are brought to my attention.

1.3 What can mwahpy do?

The following functionality is provided in *mwahpy*:

- ullet Easily & quickly read in data from a MilkyWay@home N-body .out file
- MilkyWay@home N-body output only provides 3D Cartesian positions and velocities, as well as mass (in MilkyWay@home structural units). The *mwahpy* package converts this data into many other useful forms, such as R.A. & Dec., Galactic longitude and latitude, angular momenta about the Galactic center, line-of-sight velocity, proper motions, and others.
- Make cuts based on any of *mwahpy*'s supported values, cut the data based on individual components of the data, or subsample the data randomly or symmetrically.
- Quickly plot N-body data in any of the supported values
- A variety of coordinate transformations for typical coordinate systems used in Galactic astronomy
- Two separate orbit fitting routines that have been used in publications

The proper syntax, usage, and details of each of the above capabilities will be outlined in the following sections.

1.4 Citing mwahpy

If mwahpy or any of its related code is used in a publication or project, I simply ask that you state that mwahpy was used in your work and provide the link for the mwahpy github repository located at https://github.com/thomasdonlon/mwahpy. There is currently no publication that should be cited for this package. In the future, the requirements for proper citation may change, so please check again before final publication. Thank you for using mwahpy!

2 Installing mwahpy

or

Installing mwahpy is as simple as installing any other python package. For linux machines, go to your terminal and type

```
Code Block 1: Installing mwahpy
>>> pip3 install mwahpy
```

Alternatively, you can also install mwahpy using

```
Code Block 2: Installing mwahpy

>>> python3 -m pip install mwahpy
```

It may be necessary to install *mwahpy* only in your user directory due to user access restrictions. It is strongly recommended you do not use sudo to install python packages. Using sudo makes it difficult to track who has the permissions to use what package on your machine, and on machines with multiple users, it can mean you have to install it for each user individually anyways. Misuse of sudo can also result in accidentally using a different version of *mwahpy* than intended, particularly after updates. To install *mwahpy* for only the active user, you can instead type

```
Code Block 3: Installing mwahpy
>>> pip3 install mwahpy --user
```

```
Code Block 4: Installing mwahpy
>>> python3 -m pip install mwahpy --user
```

Any of these lines of code will install mwahpy on your machine, as well as install all of the prerequisite packages needed for mwahpy to work.

If you use Anaconda as a package manager (as is becoming more and more common these days), it is recommended that you still use pip to install mwahpy as is shown above. Hoewever, this can have a couple disadvantages: For starters, the Conda package manager will not manage mwahpy. However, mwahpy will still be managed by the Anaconda environment. Additionally, you can run into some problems if your default installation of python is different than your Anaconda installation of python (a more common problem than it should be). If your default python installation is different than your Anaconda installation, I strongly suggest that you go through and fix your python configuration, as you will almost certainly run into other problems eventually. However, if you don't want to do that (or have your python installation purposefully configured that way for some reason), you can use

```
Code Block 5: Installing mwahpy

>>> conda skeleton pypi mwahpy

>>> conda build mwahpy

>>> conda install --use-local mwahpy
```

This will install *mwahpy* and avoid issues with a misconfigured Anaconda/python installation. Additionally, if *mwahpy* is installed this way, the Conda package manager will be responsible for managing *mwahpy* instead of pip. Be aware that this method may not automatically update *mwahpy* when a new vrsion of the package is released.

In order to use *mwahpy* after it is installed, simply import the subpackages as you would any other other package in your python script. For example, the following lines are all syntactically acceptable imports:

```
Code Block 6: Importing mwahpy

import mwahpy.output_handler
from mwahpy.data import Data
from mwahpy.coords import *
```

It should be noted that *mwahpy* is only built for and maintained for python v3.2.3 and above. You should get an error when installing if your python installation is not recent enough for *mwahpy*. Depending on your installation, you may instead have to type >>> python -m pip install mwahpy instead of python3, or use pip3 instead of pip. If you are experiencing errors, asking questions on the *mwahpy* github page or trying different permutations of these installation methods are recommended.

If you wish to install <code>mwahpy</code> for development, then you should clone the most recent <code>mwahpy</code> github repository (found at https://github.com/thomasdonlon/mwahpy). Feel free to make your own fork of the master branch if that's what you would prefer. Then, you should download the source code wherever you wish on your machine. If you want to import mwahpy into a python script after doing this, you will have to add a couple extra lines to the import portion of your script:

```
Code Block 7: Using mwahpy as a developer

import sys
sys.path.insert(1, '/home/path/where/init.py/is/located/')
import mwahpy_subpackage
```

In this case, you would replace the path with the path to your **init.py** file is located (it usually ends with .../mwahpy/mwahpy/), and you would replace import mwahpy_subpackage with whatever subpackage you wish to use (for example, import output_handler as oh).

If you then wish to add, remove, or change code in your distribution of *mwahpy*, that can be done easily in the folder where you cloned *mwahpy*. Any changes that you feel are beneficial to the package should be sent as a pull request to the master branch.

3 Core Functionality

WARNING:

MilkyWay@home uses a right-handed Galactic Cartesian coordinate system. This is defined as positive X being in the direction of the Sun towards the Galactic center, positive Y being in the direction of the disk spin at the location of the Sun, and positive Z being in the direction of the right-handed cross product $X \times Y$ (often called the "Galactic north"). In our coordinate system, the Sun is located at the position (X,Y,Z) = (-8,0,0).

This is in contrast to many other Galactic scientists, who prefer a left-handed coordinate system where the X-axis is flipped and the Sun is located at (X,Y,Z) = (8,0,0). In a left-handed coordinate system the physical interpretation of certain quantities are not always clear (such as the right-handed angular momentum cross product). Many coordinate transformations in mwahpy allow for left-handed coordinates. However, be aware that by default, mwahpy (and MilkyWay@home) output is right-handed.

3.1 The *Timestep* Class

The Timestep class is the heart of *mwahpy*, and it's where the majority of the important and useful calculations in the package are performed. An instance of the Timestep class represents a single timestep of an *N*-body simulation. In other words, a Timestep instance is the data from a single MilkyWay@home .out file. The code for the Timestep class can be found in *mwahpy*'s timestep.py file.

3.1.1 Initializing a *Timestep*

The most basic usage of Timestep object is a blank instance, to which you can then manually add data:

```
import numpy as np
from mwahpy.timestep import Timestep

t = Timestep()

t.id = np.array([0, 1, 2])
t.x = np.array([10, 50, 100])
t.y = np.array([12, 15, 17])
t.z = np.array([1, 2, 3])
t.vx = np.array([13, 183, 102])
t.vy = np.array([0, 50, 180])
t.vy = np.array([23, 69, 12])
t.vz = np.array([21, 1, 1])
```

These 8 values are all that you need to specify for the Timestep class to do its job. These 8 values (ID, x, y, z, vx, vy, vz, and mass) will be referred to as "provided values". This is in contrast to the "supported values", which are any values that mwahpy can calculate for you (a full list of the supported values is provided in Section 6.7). In the Timestep implementation, all of the particle IDs, x positions, etc. are stored in order as a numpy array of those values. As such, the mwahpy package heavily utilizes features of the numpy package, and some knowledge of numpy can be helpful for those working with mwahpy.

If you wish to print the information of a single particle, you can do so with printParticle by specifying the ID of the particle you are interested in:

Code Block 9: Printing data for a single particle >>> t.printParticle(1) Printing data for Particle 1: (id:1, x:50, y:15, z:2, vx:183, vy:50, vz:1, mass:1,)

WARNING:

The provided values should **always** be numpy arrays of identical length. Attempting to calculate values, plot data, or write out data when the provided values have mismatched length or are not numpy arrays will typically result in an error. It is strongly recommended that the user uses the built-in Timestep methods for cutting or adding data instead of doing it manually.

From this point forward, the user can ask for any mwahpy supported value, and it will be calculated for them. For example, if you wanted the line-of-sight velocities of the particles, you could type:

```
Code Block 10: Calculating supported values

>>> t.vlos
array([ 7.894394 , 235.8965911 , 128.90974589])
```

Note that if you call t.printParticle again (this time using the optional dec argument to shorten the output), it now shows more data for the particle!

```
Code Block 11: Printing data for a single particle

>>> t.printParticle(1, dec=2)
Printing data for Particle 1:
(id:1, x:50, y:15, z:2, vx:183, vy:50, vz:69, mass:1, msol:222288.47,
1:0.29, b:0.04, ra:266.54, dec:-28.67, dist:59.94, lx:935, ly:3084,
lz:-245, lperp:3222.62, ltot:3231.92, r:52.24, R:52.2, vlos:235.9,
vgsr:247.14, rad:192.15, rot:-4.69, distFromCOM:52.24, )
```

This was all calculated in the background when you asked the package to calculate vlos for this Timestep. Due to the overhead on some calculations, the more computationally complex calculations are avoided until the user requests those values. This is explained in more detail in Section ??.

The Timestep class has one very unique property: attributes of a Timestep instance can be accessed via the usual method, or as the key to a dictionary. In fact, comparing the two methods shows that these two actions produce equivalent results.

Code Block 12: Accessing Timestep attributes >>> t.x #accessing data as an attribute array([10, 50, 100]) >>> t['x'] #accessing data as a dict key array([10, 50, 100]) >>> np.all(t.x == t['x']) True >>> t.x[0] = 1 #changing the value for Particle 0's x position >>> t.x[0] #the same value is accessed by both methods 1 >>> t['x'][0] 1 >>> t.x[0] == t['x'][0] True

At first this property may seem confusing and not particularly useful. What is the point of being able to access the same data in two different ways? It turns out that adding this functionality to Timestep allows for some rather powerful behavior. Of the greatest importance to the typical user is the implementation of the iterator for Timestep, which iterates over the *keys* of the class. In this case, that has been implemented as the names of all of the supported values (that have been calculated so far!).

```
Code Block 13: Iterating over a Timestep

>>> outstr = ''
>>> for key in t:
...    outstr += (key + ',_')
>>> print(outstr)
id, x, y, z, vx, vy, vz, mass, msol, l, b, ra, dec, dist, lx, ly,
lz, lperp, ltot, r, R, vlos, vgsr, rad, rot, distFromCOM,
>>> outstr = ''
>>> for key in t:
...    outstr += (str(round(t[key][0],2)) + ',_')
>>> print(outstr)
0, 10, 12, 1, 13, 0, 23, 1, 222288.47, 0.88, 0.06, 266.86, -28.15, 21.66,
276, 217, -156, 351.09, 384.19, 15.65, 15.62, 7.89, 21.43, 9.77, -9.99,
15.65,
```

This may not seem like a big deal at first. However, if one tries to reproduce this behavior for a class that doesn't have the property c.x == c['x'], then you will quickly run into several problems. I expect that the typical reader would try to iterate over the class' built-in attribute dictionary, which is what I initially tried. What happens if you want to add attributes to the class that cannot be iterated over at the same time as your arrays, such as a single identifying string c.name? I suggest that the reader play around with this idea on their own if they are so inclined.

3.1.2 Reading In & Writing Out Data

So far, we know how to initialize a Timestep, how to add data to it manually, and how to access this data. This is all fine and good, but most of the time a user will be interested in using data that has already been generated by the MilkyWay@hoe N-body application. Conveniently, there is a mwahpy function built specifically for this in the mwahpy.output_handler subpackage:

The oh.readOutput function takes in the path to a MilkyWay@home .out file and outputs a Timestep instance of the data from that file. In the above code block, we read in the test ,out file, which is provided in .../mwahpy/test/. The oh.readOutput function provides you with a progress bar, which is useful for large files. In this case, the progress bar finished almost instantly since the file was small. Typically, the progress bar will reach 100%. The oh.readOutput function also provides you with the number of particles that were read in from the file, and a brief update on when it is converting the data (again, useful for very large files). From this point forward, you can perform any normal Timestep operations on the new Timestep object.

The oh.readOutput function has been heavily optimized, and operates extremely quickly. Even for files with millions of values, reading in the data only takes a few seconds on most machines. If you are working with N-body output, it is strongly recommended that you use this function to initialize your Timestep instances.

The *mwahpy* package also offers a few different options for printing out Timestep data to a file. Notably, *mwahpy* can print out the data to a .csv file,

```
Code Block 15: Writing out a Timestep

>>> oh.makeCSV(t, '/path/to/my/file.csv')
Writing header...
Printing data...
Timestep output to /path/to/my/file.csv
```

Saving the data to a .csv saves all of the supported values that has been calculated so far. Alternatively, you can write out data from a Timestep to a MilkyWay@home .in file,

```
Code Block 16: Writing out a Timestep

>>> oh.makeNbodyInput(t, '/path/to/my/file.in')
Writing Timestep as N-body input to /path/to/my/file.in...
done
```

Note that a MilkyWay@home .in file will only include the data about the 8 provided values, and not any of the other supported values. This is due to the format that MilkyWay@home requires readable files to be in. Any file that you generate with makeNbodyInput is immediately ready to be used as the manual body input for a MilkyWay@home N-body simulation.

If a user wishes to suspend a Timestep object for later use, I recommend the pickle package (https://docs.python.org/3/library/pickle.html). The pickle package allows for object serialization and saving out/reading in arbitrary objects as binary code.

3.1.3 Manipulating Data

After N-body data has been read in, it is useful to be able to only select the data that you are interested in. There are a few different routines for this, namely subsetRect() and subsetCirc(). These methods are n-dimensional cutting routines. For example, say that you took the test data and only wanted particles with a positive X value.

Note that subsetRect() has cut along all axes, not only the axis that we instructed it to cut along. As such, it is easy to see that tcopy.y has the same length as tcopy.x. It is also important to note that we made a copy of t before using subsetRect(), because the subset methods treat the Timestep object as directly mutable.

While cutting the data, we provided 1000 as an upper value for X. This was because subsetRect() requires both a lower and upper limit for each axis when cutting the data. To get around this, we just chose some arbitrarily large value that would not cut any data off at the high end of X values.

You can also cut on multiple axes with subsetRect(). Say that you wanted to only include particles with positive X and negative Y values:

```
Code Block 18: Cutting data in a Timestep

>>> tcopy = t.copy()
>>> tcopy.subsetRect(['x','y'], [(0,1000),(-1000,0)])
>>> tcopy.x
array([4.53813066, 0.38487162, 0.52268562])
>>> tcopy.y
array([-0.05701174, -0.43899208, -0.34261058])
```

As all methods in Timestep, these routines can be performed on any supported values **after they have been calculated**. Attempting to cut on line-of-sight velocity before it has been calculated, for example, would result in a KeyError.

Similar results can be obtained from subsetCirc. If we wanted all particles with X values within 1 kpc of the Galactic center:

Or if we wanted all particles with X and Y values within 1 kpc of the Galactic center:

This routine is a bit of a misnomer, as it can be extended to any n-dimensional spheroidal volume of your phase space. For example, say that we wanted particles located within a spheroid defined with an semiaxis of length 1 on the X-axis, a semiaxis of length 2 on the Y-axis, and a semiaxis of length 5 on the Z-axis, centered at (X, Y, Z) = (0, 1, 3). This can be done with

```
Code Block 21: Cutting data in a Timestep

>>> tcopy = t.copy()
>>> tcopy.subsetCirc(['x', 'y', 'z'],[1,2,5],[0,1,3])
>>> tcopy.x
array([-0.59395581, -0.00613003, 0.11393187, 0.35803147, 0.3565866])
>>> tcopy.y
array([0.87634781, 0.43348783, 0.39540971, 0.14577695, 0.0082146])
>>> tcopy.z
array([ 0.55739028, -0.18463063, 0.32116868, 0.17315727, -0.09600029])
```

Unlike subsetRect(), this routine becomes somewhat unclear when you begin mixing values with different units. In that case, it is suggested that you either make multiple circular cuts where the units of each value match, or make rectangular cuts instead.

Other data manipulation routines include sampling and splitting Timesteps. There are three types of sampling that *mwahpy* supports: manual, incremental, and random sampling, shown below.

Code Block 22: Sampling a Timestep >>> #manual sampling >>> tcopy = t.copy() >>> tcopy.take([0,2,3,6,9]) #take particles at indices 0, 2, 3, 6, and 9 array([4.53813066, -1.41000385, -0.00613003, 0.35803147, 3.57179938]) >>> #incremental sampling >>> tcopy = t.copy() >>> tcopy.subsample(2) #take every 2nd particle, starting at 0 >>> tcopy.x array([4.53813066, -1.41000385, 0.11393187, 0.35803147, 0.52268562]) >>> #random sampling >>> tcopy = t.copy() >>> tcopy.randSample(5) #randomly take 5 particles from the data array([4.53813066, -0.59395581, 0.38487162, -0.00613003, 0.35803147]) >>> #run again to demonstrate randomness >>> tcopy = t.copy() >>> tcopy.randSample(5) >>> tcopv.x array([0.52268562, -0.59395581, 0.3565866, -0.00613003, 0.11393187])

It is easy to split a Timestep by providing the index at which you want to split:

```
Code Block 23: Splitting a Timestep

>>> tcopy, tcopy2 = t.split(4) #split at index 4
>>> tcopy.x
array([ 4.53813066, -0.59395581, -1.41000385, -0.00613003])
>>> tcopy2.x
array([ 0.11393187,  0.38487162,  0.35803147,  0.3565866,  0.52268562,  3.57179938])
```

Finally, if you would like to add two Timestep instances together, use the appendTimestep method:

Note that actions like just replacing a single x value in a Timestep does not update the rest of the calculated values in the Timestep. That Timestep will need to be updated manually using Timestep.update(). This method will update all values associated with the Timestep in order to keep them in sync with the provided values.

WARNING:

Note that if the autoUpdate flag is turned on (it is on by default), *mwahpy* functions will automatically keep the provided and calculated values in sync. However, if you manually update the provided values after the calculated values have been computed, you will need to run Timestep.update().

```
Code Block 25: Updating a Timestep
>>> t = oh.readOutput('../test/test.out')
Reading in data from ../test/test.out...
[-----
                    ] 73%
10 objects read in
Converting data...done
>>> t.x
array([ 4.53813066, -0.59395581, -1.41000385, -0.00613003, 0.11393187,
        0.38487162, 0.35803147, 0.3565866, 0.52268562, 3.57179938])
>>> t.distFromCOM
Calculating basic values...
array([4.53771154, 1.20665236, 2.01267674, 0.48791978, 0.51046082,
       0.99322694, 0.39540509, 0.34974823, 0.85795412, 3.73818636])
>>> t.x[0] = 0
>>> t.distFromCOM
array([4.53771154, 1.20665236, 2.01267674, 0.48791978, 0.51046082,
       0.99322694, 0.39540509, 0.34974823, 0.85795412, 3.73818636])
>>> t.update()
>>> t.distFromCOM
array([0.80324921, 1.29094239, 2.05506687, 0.36725208, 0.51176956,
       0.99371407, 0.34057982, 0.29000981, 0.81102414, 3.37390626])
```

You should now have a fundamental undertsanding of how to use the Timestep class. The rest of the methods that were not covered in this section can be found in Section 6.7. Continue reading for information about handling entire simulations with multiple timesteps and plotting timesteps and simulations.

3.2 The *Nbody* Class

The Nbody class is one rung above the Timestep class in the *mwahpy* hierarchy; one Nbody instance is composed of many individual Timestep instances. This is useful when you need to compare many timesteps from a single MilkyWay@home simulation. The simplest example of a Nbody instance is the default instance:

```
Code Block 26: Initializing an Nbody

>>> from mwahpy.nbody import Nbody
>>> n = Nbody()
```

From here, you can add a Timestep to the dictionary in the Nbody instance:

Note that the Nbody instance itself does not carry the Timestep data, but just points to it.

```
Code Block 28: An Nbody does not carry simulation data

>>> n.x
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
AttributeError: 'Nbody' object has no attribute 'x'
```

In this case, the Timestep t still exists on its own, but the Nbody instance forms a nice functional container for it. This can be seen by altering t, as the change is also present when accessing the data through the Nbody. Thus, the Nbody structure does not copy the Timestep data, it just references it.

The $\mathsf{Timestep}$ does recognize that it belongs to an Nbody class however, and it remembers its designated time and the Nbody object which it belongs to:

```
Code Block 30: Accessing an Nbody

>>> t.time
1
>>> t.nbody
<mwahpy.nbody.Nbody object at 0x[...]>
```

WARNING:

Currently, Timestep instances only weakly retain information about parent Nbody instances. If a Timestep is added to a second Nbody, it will replace its self.time and self.nbody from the first Nbody instance with the new information from the second Nbody. This is made more confusing by the fact that multiple Nbody instances can point to the same Timestep, or a single Nbody can point to the same Timestep more than once. In these cases, the data can be properly accessed through the Nbody objects, but is not guaranteed to refer back to the desired information when accessed from the Timestep object.

3.2.1 Reading In & Writing Out Data

As with the Timestep class, the Nbody class is typically not created manually, but is initialized by reading in data. This is also done through the mwahpy.output_handler subpackage by providing a folder with the oh.readFolder() function.

```
Code Block 31: Reading data into an Nbody
>>> n = oh.readFolder('<...path/to/mwahpy>/mwahpy/test/nbody_test/')
Reading in data from directory ../test/nbody_test/...
Reading in data from ../test/nbody_test/1...
[-----
                    ] 73%
10 objects read in
Converting data...done
Reading in data from ../test/nbody_test/2...
[---->
                    ] 73%
10 objects read in
Converting data...done
Reading in data from ../test/nbody_test/3...
[---->
                    ] 73%
10 objects read in
Converting data...done
>>> n[1].x
array([ 4.53813066, -0.59395581, -1.41000385, -0.00613003, 0.11393187,
       0.38487162, 0.35803147, 0.3565866, 0.52268562, 3.57179938])
>>> n[2].x
array([21.51019042, 17.97537383, 15.99136357, 12.30622142, 10.80896895,
      10.56226125, 12.94243578, 11.16187291, 11.17781449, 8.04042707])
```

This data has been read in from the .../mwahpy/test/nbody_test/ folder that is included in mwahpy.

WARNING:

When reading in a Nbody from simulation data, the **only** files in the provided folder must be MilkyWay@home .out files, and **must** be named only their time **without the** .out **extension**. This is the default output scheme of MilkyWay@home N-body.

Currently, there is not support for writing out Nbody data. This can be done fairly simply by iterating over the Nbody (see the next subsection for details) and writing out each Timestep. It is also not recommended that a user writes out Nbody data using the pickle package.

3.2.2 Functionality of *Nbody* Objects

Nbody objects have a few nice capabilities that allow for robust treatment of simulation data. To begin, iterating over an Nbody object returns its component Timestep objects sorted by time.

```
Code Block 32: Iterating over an Nbody

>>> for ts in n:
... print(ts.time)

1
2
3
```

Being able to read in and iterate over large groups of Timestep instances from the same simulation is much easier than going through each one individually, not to mention it is often faster.

The times of each timestep can be scaled in order to reflect the actual physical simulation time of each timestep using the scaleTimes() method. For example, if one timestep is equal to 0.5 Myr in physical time, then we can scale the corresponding timesteps. We see that the original information is retained, although renamed:

```
Code Block 33: Scaling Nbody times

>>> n.scaleTimes(0.5)
>>> for ts in n:
... print(ts.time)
0.5
1.0
1.5
>>> n[0.5].x
array([ 4.53813066, -0.59395581, -1.41000385, -0.00613003, 0.11393187, 0.38487162, 0.35803147, 0.3565866 , 0.52268562, 3.57179938])
```

3.3 Plotting

Text

4 Auxiliary Subpackages

The *mwahpy* package comes with a few auxiliary subpackages with additional content that the typical user might find useful. While the mwahpy.orbit_fitting and the mwahpy.orbit_fitting_gc subpackages have been used in academic contexts, the packages have not yet been refactored for general use. After the packages have been made consistent with the rest of the *mwahpy* code standards, then documentation for those subpackages will be released.

4.1 Coordinate Transformations

The coordinate transformation package mwahpy.coords is made up of many first and second-order transformations for position, velocity, and other values. A complete list of the routines and functions provided in this package is given in Section 6.1.

This subpackage contains most of the coordinate transformations that a typical person will need to do Galactic astronomy, excluding some common transformations (such as R.A., Dec. to l, b) that are implemented in the astropy python package.

4.2 Orbit Fitting

The subpackages mwahpy.orbit_fitting and mwahpy.orbit_fitting_gc are both still under development. Once the code has been refactored and brought to an acceptable level of quality, the documentation for these subpackages will be provided.

5 Flags & Settings

5.1 flags

The flags and settings for *mwahpy* can be found in flags.py. Currently, there are only a few options, but more may be added in the future.

• mwahpy.flags.autoUpdate (default value = 1)

This flag determines whether or not to automatically update an entire Timestep whenever the provided values for that Timestep change due to a *mwahpy* method or function. Having this flag turned on will keep things like the center of mass and momentum accurate even after changing the constituent data, and will keep all of the calculated values in sync with the provided values throughout *mwahpy* functions.

If you are frequently updating certain values in a Timestep, this flag can occasionally cause performance issues (if this flag is causing performance issues, that's probably a sign that there's a better way to do what you are trying to do). It is strongly recommended that this flag is only turned off if the user understands what updating means, how to update the data manually, and what data needs to be updated.

• mwahpy.flags.progressBars (default value = 1)

If this flag is turned on, *mwahpy* will use progress bars when performing certain actions (such as reading in files) in order to avoid long wait times without updating the terminal. If turned off, *mwahpy* functionality will not be changed; however, be aware that you may occasionally experience long periods of time where your terminal is silent.

• mwahpy.flags.verbose (default value = 1)

This flag dictates how much is printed out into the terminal when *mwahpy* code is running. If turned on, functions are much noisier, and will update you on their current activity. If this is turned off, the overall functionality of *mwahpy* will remain unchanged, but the terminal will be less informative.

$5.2 \quad mwahpy_glob$

Constants and functions that are universally accessed in mwahpy are stored in mwahpy_glob.py.

mwahpy.mwahpy_glob.fileLen(f)

Given a file, this function will determine the number of lines in that file. Useful for implementing mwahpy_glob.progressBar() while reading in files.

Parameters:

- f (str): The filename (should be a path).

Returns:

- len (int): The number of lines in the file.
- mwahpy.mwahpy_glob.G

Newton's gravitational constant is provided in astropy units of m³/kg s².

mwahpy.mwahpy_glob.kmsToKpcgyr

The conversion factor from km/s to kpc/Gyr. Roughly equal to unity (1.023).

mwahpy.mwahpy_glob.kpcgyrToKms

The conversion factor from kpc/Gyr to km/s. Roughly equal to unity (0.978).

• mwahpy.mwahpy_glob.progressBar(value, endvalue, bar_length=20)

Adapted from https://stackoverflow.com/questions/6169217/replace-console-output-in-python. This function can be placed inside a loop and will output a nice progress bar in the terminal, provided you know the end value that the loop should terminate on.

Parameters:

- value (int): The value that is being iterated.
- endvalue (int): The value at which the loop terminates.
- bar_length (int): How many characters long the progress bar is in the terminal output.

Returns:

• mwahpy.mwahpy_glob.structToSol

This is the conversion factor between MilkyWay@home structural masses and solar masses. Equal to 222,288.47 solar masses/structure unit.

5.3 pot

Constants and functions related to the Milky Way's gravitational potential are found in mwahpy.pot.py. The default potential that is used in mwahpy is the Orphan Stream Fit #5 Potential from Newberg et al. (2010). This potential is very similar to Law et al. (2005).

• mwahpy.pot.energy_offset

Due to the logarithmic halo potential term in our gravitational potential for the Milky Way, the potential does not disappear as one travels infinitely far from the center of the potential. A side effect of this type of halo potential is that it is not clear what structures are bound or unbound to the potential, as the zero-point of the energy is not representative of the escape velocity. This adjusts the calculated potential energy so that it is consistent with Donlon et al. (2019), and so that clearly bound structures will not have positive energy values. The default value is -60000 km²/s².

• mwahpy.pot.m_bulge

The mass of the bulge in astropy solar masses. Equal to 3.4e10 M_{\odot}

• mwahpy.pot.m_disk

The mass of the disk in astropy solar masses. Equal to 1.0e11 M_{\odot}

• mwahpy.pot.v_halo

The Milky Way halo dispersion in astropy km/s. Equal to 74.61 km/s.

• mwahpy.pot.plotPotential(potential, Rrange=[0.01,10.])

This method will generate and show a figure with the rotation curve of the potential in the plane of the Milky Way.

Parameters:

- potential (galpy potential object) : the potential that you wish to graph
- Rrange (list of floats): the Galactocentric cylindrical radius of the points at which you want to evaluate the potential

Returns:

• mwahpy.pot.pot

The total gravitational potential of the Milky Way. Given as a galpy potential object.

- mwahpy.pot.pot_bulge

 The gravitational potential of the Milky Way due to the bulge. Given as a galpy potential object.
- mwahpy.pot.pot_disk

 The gravitational potential of the Milky Way due to the disk. Given as a galpy potential object.
- mwahpy.pot.pot_halo

 The gravitational potential of the Milky Way due to the halo. Given as a galpy potential object.

6 Functions & Methods

Below is a complete list of functions and methods implemented in the files of *mwahpy* that perform actual routines. Note that only the functions and methods that are meant for the typical end user are provided in detail here, and therefore some helper functions/methods that are present in the code are not gone over in this document. For a tutorial of using the main parts of *mwahpy*, one should look at Section 3.

6.1 coords

Text

$6.2 \quad nbody$

Text

$6.3 \quad orbit_fitting$

The subpackage mwahpy.orbit_fitting is still under development. Once the code has been refactored and brought to an acceptable level of quality, the documentation for this subpackage will be provided.

$6.4 \quad orbit_fitting_gc$

The subpackage mwahpy.orbit_fitting_gc is still under development. Once the code has been refactored and brought to an acceptable level of quality, the documentation for this subpackage will be provided.

$6.5 \quad output_handler$

Text

6.6 *plot*

Text

6.7 timestep

These functions and methods are all related to the timestep class, and are located in the mwahpy/timestep.py file. These functions and methods make up the main core of mwahpy functionality, and rely on a lot of the other code in the package.

Additionally, a partial list of the many attributes of the timestep class are given below, and what they store is described in detail. Note that many attributes will return empty lists, empty arrays, None, or 0 initially if accessed without actually reading in or supplying data.

There are several other flags, functions, and attributes that are implemented in the timestep.py file that are not detailed below. These are not meant to be accessed by end users, and are explained by comments in the code.

6.7.1 Initialization

An instance of the Timestep class is initialized by

Note that all of the provided values between id_val and mass must be arrays of the same length, or else there will be errors.

- id_val (list of ints, optional): The id values of each particle (in order).
- x (list of floats, optional): The Galactocentric X positions of each particle in units of kpc (in order).

- y (list of floats, optional): The Galactocentric Y positions of each particle in units of kpc (in order).
- z (list of floats, optional): The Galactocentric Z positions of each particle in units of kpc (in order).
- vx (list of floats, optional): The Galactic standard of rest velocities of each particle in the X direction in units of km/s (in order).
- vy (list of floats, optional): The Galactic standard of rest velocities of each particle in the Y direction in units of km/s (in order).
- vz (list of floats, optional): The Galactic standard of rest velocities of each particle in the Z direction in units of km/s (in order).
- mass (list of floats, optional): The masses of each particle in units of MilkyWay@home structure masses (in order).
- centerOfMass (list of floats, optional): The 3D location of the center of mass of the system (all particles in the Timestep instance) in units of kpc. The first element of the list is the Galactocentric X position of the center of mass, while the second and third elements are the Y and Z positions, respectively.
- mwahpy.timestep.centerOfMomentum (list of floats, optional): The 3D location of the center of momentum of the system (all particles in the Timestep instance) in units of km/s. The first element of the list is the Galactocentric vx position of the center of mass, while the second and third elements are the vy and vz positions, respectively.
- mwahpy.timestep.potential (galpy potential object or list of galpy potential objects, optional): Can be None. If not None, the energy of each particle will be calculated with this potential instead of the default mwahpy potential.
- time (float, optional): The time in Gyr after the beginning of the simulation.
- nbody (Nbody object, optional): The parent Nbody instance for this Timestep.

6.7.2 Attributes

• mwahpy.timestep.b

(numpy array of floats): The Galactic latitude of each particle in degrees, in order. Calculated value.

• mwahpy.timestep.centerOfMass

(list of floats): The 3D location of the center of mass of the system (all particles in the Timestep instance) in units of kpc. The first element of the list is the Galactocentric X position of the center of mass, while the second and third elements are the Y and Z positions, respectively.

• mwahpy.timestep.centerOfMomentum

(list of floats): The 3D location of the center of momentum of the system (all particles in the Timestep instance) in units of km/s. The first element of the list is the Galactocentric vx position of the center of mass, while the second and third elements are the vy and vz positions, respectively.

• mwahpy.timestep.dec

(numpy array of floats): The declination of each particle in degrees, in order. Calculated value. Computed using astropy.

• mwahpy.timestep.dist

(numpy array of floats): The heliocentric spherical radius of each particle in units of kpc, in order. Equal to $\sqrt{(X+8)^2+Y^2+Z^2}$. Calculated value.

• mwahpy.timestep.distFromCOM

(numpy array of floats): The distance of each particle from the center of mass of the Timestep instance, in order. Calculated value.

mwahpy.timestep.energy

(numpy array of floats): The total energy per unit mass of each particle based on the specified potential in units of $\rm km^2/s^2$ (in order). If no potential is specified, this is calculated using the default $\it mwahpy$ potential. Calculated value as part of calcEnergy().

• mwahpy.timestep.id

(numpy array of ints): The id of each particle (in order).

• mwahpy.timestep.indexList

(list of strs): A list of values that are currently calculated for the Timestep. This is automatically updated as new values are calculated. Examples of list members are 'x', 'mass', 'vlos', etc. This is used by *mwahpy* to determine what values need to be calculated and/or iterated over, and therefore it is strongly recommended to not alter this attribute manually.

• mwahpy.timestep.KE

(numpy array of floats): The kinetic energy per unit mass of each particle in units of km²/s² (in order). Calculated value as part of calcEnergy().

• mwahpy.timestep.l

(numpy array of floats): The Galactic longitude of each particle in degrees, in order. Calculated value.

• mwahpy.timestep.lperp

(numpy array of floats): The magnitude of the projection of the angular momentum of each particle onto the X-Y plane, in order. Has units of kpc·km/s. Equal to $\sqrt{L_x^2 + L_y^2}$. Calculated value.

• mwahpy.timestep.ltot

(numpy array of floats): The magnitude of the total angular momentum of each particle in units of kpc·km/s, in order. Calculated value.

• mwahpy.timestep.lx

(numpy array of floats): The X component of the angular momentum of each particle about the origin in units of kpc·km/s, in order. Calculated value.

• mwahpy.timestep.ly

(numpy array of floats): The Y component of the angular momentum of each particle about the origin in units of kpc·km/s, in order. Calculated value.

• mwahpy.timestep.lz

(numpy array of floats): The Z component of the angular momentum of each particle about the origin in units of kpc·km/s, in order. Calculated value.

mwahpy.timestep.mass

(numpy array of floats): The mass of each particle in units of MilkyWay@home structure masses (in order).

• mwahpy.timestep.msol

(numpy array of floats): The mass of each particle in units of solar masses (in order). Calculated value.

mwahpy.timestep.nbody

(nbody object): If the Timestep instance is part of an nbody instance, i.e. if you read in the Timestep with mwahpy.output_handler.readFolder, then that information is saved here.

• mwahpy.timestep.PE

(numpy array of floats): The potential energy per unit mass of each particle based on the specified potential in units of $\rm km^2/s^2$ (in order). If no potential is specified, this is calculated using the default $\it mwahpy$ potential. Calculated value as part of calcEnergy().

• mwahpy.timestep.pmdec

(numpy array of floats): The proper motion of each particle in the direction of declination in units of mas/yr (in order). Calculated value as part of calcrvpm().

• mwahpy.timestep.pmra

(numpy array of floats): The proper motion of each particle in the direction of right ascension in units of mas/yr (in order). Multiplied by $\cos \delta$. Calculated value as part of calcrvpm().

• mwahpy.timestep.pmtot

(numpy array of floats): The magnitude of the total proper motion of each particle in units of mas/yr (in order). Calculated value as part of calcrvpm().

• mwahpy.timestep.potential

(galpy potential object or list of galpy potential objects): If not None, the energy of each particle will be calculated with this potential. If None, the *mwahpy* default potential will be used to calculate the particle energies.

• mwahpy.timestep.r

(numpy array of floats): The Galactocentric spherical radius of each particle in units of kpc, in order. Equal to $\sqrt{X^2 + Y^2 + Z^2}$. Calculated value.

• mwahpy.timestep.R

(numpy array of floats): The Galactocentric cylindrical radius of each particle in units of kpc, in order. Equal to $\sqrt{X^2 + Y^2}$. Calculated value.

• mwahpy.timestep.ra

(numpy array of floats): The right ascension of each particle in degrees, in order. Calculated value. Computed using astropy.

• mwahpy.timestep.rad

(numpy array of floats): The magnitude of the Galactic standard of rest line-of-sight velocity of each particle as seen from the origin. Has units of km/s. Calculated value.

• mwahpy.timestep.rot

(numpy array of floats): The magnitude of the Galactic standard of rest rotational velocity (projected onto the X-Y plane) of each particle as seen from the origin. Has units of km/s. Calculated value.

• mwahpy.timestep.time

(float): The time in Gyr for this Timestep instance. This will only be initialized if the Timestep was read in as part of mwahpy.output_handler.readFolder() or if manually specified. Otherwise, will return None by default.

• mwahpy.timestep.vgsr

(numpy array of floats): The line-of-sight velocity of each particle in the Galactic standard of rest in units of km/s, in order.

• mwahpy.timestep.vlos

(numpy array of floats): The heliocentric line-of-sight velocity of each particle in units of km/s, in order.

• mwahpy.timestep.vtan

(numpy array of floats): The heliocentric, Galactic standard of rest tangential velocity of each particle in units of km/s (in order). Calculated value as part of calcrvpm().

• mwahpy.timestep.vx

(numpy array of floats): The Galactic standard of rest velocity in the X direction of each particle in units of km/s (in order).

• mwahpy.timestep.vy

(numpy array of floats): The Galactic standard of rest velocity in the Y direction of each particle in units of km/s (in order).

• mwahpy.timestep.vz

(numpy array of floats): The Galactic standard of rest velocity in the Z direction of each particle in units of km/s (in order).

• mwahpy.timestep.x

(numpy array of floats): The Galactocentric X position of each particle in units of kpc (in order).

mwahpy.timestep.y

(numpy array of floats): The Galactocentric Y position of each particle in units of kpc (in order).

• mwahpy.timestep.z

(numpy array of floats): The Galactocentric Z position of each particle in units of kpc (in order).

6.7.3 Functions & Methods

• mwahpy.timestep.appendTimestep(t)

Combines two Timestep objects by appending the particles from the provided Timestep onto this Timestep.

Parameters:

t (Timestep object): The Timestep object that you wish to combine with this one.

Returns:

• mwahpy.timestep.appendPoint(t, n=0, id=None)

Adds a specific particle from the provided Timestep to this Timestep. What particle you wish to append can be specified by the id of the particle, or what position it is located at in the specified Timestep object.

WARNING: This function is extremely time intensive if used repeatedly due to the overhead associated with numpy arrays. One should always try to use combinations of cuts and mwahpy.timestep.appendTimestep() to append multiple particles whenever possible.

Parameters:

- t (Timestep object) : The Timestep object containing the particle that you wish to add to this Timestep.
- n (int, optional): The location of the particle in t that you wish to add to this Timestep. Ignored if id is not None.
- id (int, optional): If not None, this function locates the particle with id equal to this value and adds it to this Timestep.

Returns:

• mwahpy.timestep.copy()

Creates a deep copy of the Timestep instance. No arrays or lists in the copy are aliased to the original respective objects. Will not assign the corresponding "pointer" in a parent Nbody instance to the new copy.

Parameters:

Returns:

- out (Timestep): A deep copy of the Timestep instance.

• mwahpy.timestep.cutFirstN(n)

Removes the first n particles from the Timestep instance, based on the particle order.

Parameters:

- n (int): The number of particles to remove from the front of the Timestep.

Returns:

• mwahpy.timestep.cutLastN(n)

Removes the last n particles from the Timestep instance, based on the particle order.

Parameters:

- n (int): The number of particles to remove from the back of the Timestep.

Returns:

• mwahpy.timestep.hist2d(x, y, **kwargs)

Creates and optionally shows a 2D histogram plot of the particle data in the Timestep. See mwahpy.plot.hist2d() for a more in-depth explanation of the plotting routine.

Parameters:

- x (str): The values that you want to plot on the horizontal axis. For example, if you want to plot vlos on the horizontal axis, you would input 'vlos'.
- y (str): The values that you want to plot on the vertical axis (see x above).
- **kwargs (optional): Keyword arguments that you want to pass to mwahpy.plot.hist2d() or matplotlib.pyplot.hist2d. See the documentation for those functions for more information about what keyword arguments are supported.

Returns:

• mwahpy.timestep.len()

Returns the length of the timestep's values arrays. This is only calculated from the timestep.id array, so if that is out of sync with the other arrays for some reason this will not return an accurate value.

Parameters:

Returns:

- len (int): The length of the timestep's values arrays.
- mwahpy.timestep.printParticle(n, dec=8)

Prints information about all values corresponding to a single particle in the console.

Parameters:

- n (int): The location of the particle in this Timestep that you want to print the information for.
- dec (int, optional): The number of digits after the decimal point to display for each value.

Returns:

• mwahpy.timestep.randSample(n)

Randomly samples the $\mathsf{Timestep}$ down to exactly n particles selected at random from the $\mathsf{Timestep}$. Uses the reservoir theorem to sample the data.

Parameters:

- n (int): The number of particles to include in the random sample.

Returns:

• mwahpy.timestep.recenter()

Shifts the positions and velocities of each particle by the same amount in order to make the center of mass and center of momentum both be located at the origin. This function does not change any physics, it just changes the frame in which the Timestep resides.

Parameters:

Returns:

• mwahpy.timestep.resetIds()

Relabels the id of each particle, in order, starting from 0 and incrementing by 1 each particle. Retains the current order of the particles.

Parameters:

Returns:

• mwahpy.timestep.rotateAroundZAxis(theta, rad=False)

Rotates the particles in the Timestep counter-clockwise around the Z axis by some angle theta.

Parameters:

- theta (float): The angle to rotate the particles around the Z axis. Default unit is in degrees (see rad).
- rad (bool, optional): Whether or not theta is in radians.

Returns:

mwahpy.timestep.scatter(x, y, **kwargs)

Creates and optionally shows a scatter plot of the particle data in the Timestep. See mwahpy.plot.scatter() for a more in-depth explanation of the plotting routine.

Parameters:

- x (str): The values that you want to plot on the horizontal axis. For example, if you want to plot vlos on the horizontal axis, you would input 'vlos'.
- y (str): The values that you want to plot on the vertical axis (see x above).
- **kwargs (optional): Keyword arguments that you want to pass to mwahpy.plot.scatter() or matplotlib.pyplot.scatter. See the documentation for those functions for more information about what keyword arguments are supported.

Returns:

• mwahpy.timestep.split(n)

Splits the Timestep into two new Timestep instances at the nth particle. The first output Timestep will contain the first n objects of the original Timestep, and the second output Timestep will contain the rest of the particles. Does not alter the original Timestep.

Parameters:

- n (int): The location at which to split the Timestep.

Returns:

- Timestep1 (Timestep object): A Timestep instance containing the first n particles from the original Timestep.
- Timestep2 (Timestep object): A Timestep instance containing the rest of the particles from the original Timestep that are not included in Timestep1.

mwahpy.timestep.splitAtIdWrap()

Splits the Timestep at each location where the id wraps back to 0. This is useful for simulations with multiple components. See mwahpy.timestep.split() for more information about splitting Timestep objects. The original Timestep object is not altered.

WARNING: Under some conditions, MilkyWay@home can begin indexing particles in a MilkyWay@home generated dwarf (or dwarf component[s]) starting with 1, instead of 0. In this case, the particles will have to be manually split. This is a known bug in MilkyWay@home.

Parameters:

Returns:

- outlist (list of Timestep objects): A list containing a new Timestep instance made up of each component in the original Timestep object (in this case a component is defined by the particles in between two ids that are 0 in a Timestep). Each particle from the original Timestep belongs to exactly 1 component.

• mwahpy.timestep.subsample(n, offset=0)

Uniformly samples the Timestep by taking every nth particle starting from the offset.

Parameters:

- n (int): The number of particles to skip between each sample particle.
- offset (int, optional): The number of particles to initially skip.

Returns:

mwahpy.timestep.subsetRect(axes, bounds)

Takes a multidimensional rectangular cut on the Timestep. The number of cuts made and their extents are defined by axes and bounds. For each axis specified in axes, all particles with values outside the respective bounds are removed from the Timestep. Axes may be reused (if, for example, one wanted to remove all particles within some rectangular boundary instead of all particles outside it).

If you wish to only declare a minimum or maximum value for your data, then set the other bound as None (see bounds).

Parameters:

- axes (list of strs): Each axis that will be cut on. For example, if you want to make a cut in proper motion, a cut in line-of-sight velocity, and a cut in distance, you would write ['pmra', 'vlos', 'dist']. Order does not matter, but the order of the boundaries in bounds and the axes in axes must be the same.
- bounds (list of tuples of floats): The boundaries for each cut. For the example above in axes, if you wanted to include particles only with proper motions between -2 mas/yr and 2 mas/yr, line-of-sight velocity above 0 km/s, and distance less than 10 kpc, you would write [(-2, 2), (0, None), (None, 10)]. The order of the boundaries must be the same as the order of the axes specified in axes.

Returns:

mwahpy.timestep.subsetCirc(axes, rads, centers)

Takes a multidimensional "circular" cut on the Timestep. The number of cuts made and their extents are defined by axes, rads, and centers. For each axis specified in axes, all particles with values outside the specified region are removed from the Timestep. Axes may be reused.

For example, if one wanted all particles with position within 2 kpc of the point (X, Y) = (10, 15) kpc, one could make axes equal to ['x', 'y'], make rads equal to [2, 2], and make centers equal to [10, 15].

Parameters:

- axes (list of strs): Each axis that will be cut on. Order does not matter, but the order of the boundaries in rads and centers and the order of the axes given in axes must be the same.
- rads (list of floats): The radii of each cut. The order of the radii must be the same as the order of the axes specified in axes.
- centers (list of floats): The centers of each cut. The order of the centers must be the same as the order of the axes specified in axes.

Returns:

• mwahpy.timestep.take(indices)

Cuts the Timestep to only include the particles with the specified values of id.

Parameters:

indices (list of ints): The list of ids. Only particles with id equal to a value in this list will be in the cut Timestep.

Returns:

• mwahpy.timestep.update(force=False)

Updates the centerOfMass and centerOfMomentum attributes of the Timestep instance based on the current positions and velocities of all particles in the Timestep. Also updates all calculated values. Can be time intensive depending on how much needs to be updated, and whether the positions/velocities of the particles were actually changed (this is tracked in the Timestep implementation, and items are only updated if necessary).

If force is set to True, then a complete update of all attributes of the Timestep will happen even if no positions or velocities have changed since the last update or initialization of the Timestep.

Parameters:

 force (bool, optional): If true, force a complete update of the Timestep regardless of what has changed since the last update.

Returns: