# **CirBinDis**Circumbinary disk analyser

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https://bitbucket.org/paulmag/circumbinarydisk

## 1 Introduction

A small piece of software for receiving an artificial light curve from a simulated density map of a gas disk around a binary star.

This explains certain procedures with bash-commands, which exists in Linux/UNIX and Apple OS. CirBinDis should work in Windows as well, but some bash-commads may be different.

# 2 Installing

#### 2.1 Downloading

The source code is available at this Bitbucket repository:

https://bitbucket.org/paulmag/circumbinarydisk

Provided that Git is installed on you computer you can easily get all the files by running the following command inside the folder where you want the repository: git clone https://paulmag@bitbucket.org/paulmag/circumbinarydisk.git This link will be updated if the location of the repository or the installation process in any other way changes (note that at this moment the repository is private). Should it not work then contact the authors via email.

#### 2.2 Alias

We recommended to make the alias "cirbindis" for the command python ~/path\_to\_repository\_folder/circumbinarydisk/src/main.py. F. ex. place this in your .bashrc or .bash\_aliases: alias cirbindis="python /GitHub/circumbinarydisk/src/main.py" This alias will be assumed for the rest of this manual.

# 3 Preparing your data

The format of the input data must be an ASCII/CSV-file with three columns where each line represents a datapoint in space. The two first columns of each line represent the position of a datapoint. (x, y) if using cartesian coordinates and  $(r, \theta)$  if using polar coordinates. The last column represents the density in this position.

Any units can be used for the input data. How to specify units are covered in the section **Configuring and running CirBinDis**.

## 4 Processing algorithm

CirBinDis produces artificial lightcurves by analysing the provided dataset according to given configurations. In this section the process for extracting the lightcurve from the dataset is explained.

#### 4.1 Loading data

**TODO** 

#### 4.2 Cropping

The space covered by the dataset may represent a larger area than the disk you want to analyse. The dataset is cropped to an inner and outer radius such that the shape of the remaining datapoints resembles a donut. The outer radius represents the size of the disk and makes sure that the disk is circular. The inner radius is necessary to avoid treating the stars themselves as dust, and the density of the dust is very low close to the stars anyway.

## 4.3 Rotating

The coordinates of all datapoints are rotated stepwise with the rotation matrix  $R_z$  for  $\theta = [0, 360)$ °.

$$R_z = \begin{bmatrix} \cos(\theta) & -\sin(\theta) & 0\\ \sin(\theta) & \cos(\theta) & 0\\ 0 & 0 & 1 \end{bmatrix}$$

This rotation simulates the physical orbital rotation of the dircumbinary disk. The reason we get a variation in the lightcurve is because when the disk rotates we see the stars through different areas of the disk with different densities.

A rotation also happens around the y-axis due to the inclination angle  $\phi$ .  $R_y$  is the rotation matrix which would perform this rotation. However,  $R_y$  is not used, and the y-rotation is never performed directly. It is implicitly done in a very different way in a following section Mean density of bins, weighted and integrated.

$$R_y = \begin{bmatrix} \cos(\phi) & 0 & \sin(\phi) \\ 0 & 1 & 0 \\ \sin(\phi) & 0 & \cos(\phi) \end{bmatrix}$$

# 4.4 Sylinder

A section of the datapoints are cropped out, which represents only the sylinder of gas that is between an observer on Earth and the star. These are the datapoints that fall within the sylinder whose base area is defined by the stellar surface and which extends from the stellar surface and infinitely along the x-axis in positive direction (the de facto limit is the outer radius of the disk). In other words, the observer's position is assumed to be  $(\infty, 0, 0)$ . A sylinder like this is made once for each azimuthal rotation of all the points. Thus, each sylinder will be a little different from the previous one (if  $d\theta$  is small). If there are two (or even more) stars a sylinder will be created for the line of sight of each star, so there can be two (or even more) sylinders at the same time.

#### 4.5 Binning

#### 4.5.1 Algorithm

Each sylinder is sliced up into  $n_{steps}$  bins along the line of sight, where  $n_{steps}$  is given by the field radiussteps in input.xml.  $N_{sylinder}$  is the number of datapoints contained within a sylinder. For each bin the mean density is computed. The binning algorithm works like this:

- 1. Sort all datapoints in sylinder according to x-component.
- 2. Find  $N_{bin} = N_{sylinder}/n_{steps}$ .
- 3. First  $N_{bin}$  (sorted) datapoints goes in the first bin, next  $N_{bin}$  datapoints go in the second bin, etc.
- 4. Create corresponding dr array, where the dr corresponding the each bin is the difference between the x-component of the first and last datapoint in that bin.

#### 4.5.2 Reasoning

An alternative way this could be done is have a static  $\Delta r$  and check which points fall within [r, r + dr] for r in  $[0, 1, 2, 3...] \cdot \Delta r$ , but this requires a boolean test on the entire sylinder for each radius. It is much faster to sort the datapoints in the sylinder once and then just slice it with indices. There could be even smarter ways to do it, but this has worked well for now. A side effect of this method is that  $\Delta r$  is smaller in areas where there are more datapoints. If the grid of datapoints is spaced denser in central areas where the most interesting features are this is a bonus compared to a static  $\Delta r$ .

#### 4.6 Mean density of bins, weighted and integrated

#### 4.6.1 Mass integral

For each bin a mean density is produced from all the datapoints in that bin. This is done by dividing the total mass of the sylinder with its volume. The mass of a

sylinder can be calculated from the following integral.

$$M_{bin,j} = \int \int \int_{V_{bin,j}} \rho(x,y,z) \, dx \, dy \, dz$$

All our density datapoints are in the xy-plane, so the x and y part of the integral can be evaluated by summing the density of each datapoint  $\rho_{0,i}$  multiplied with its respective discrete  $dx_i dy_i$ . We will assume that the datapoints in the grid is spaced evenly. This is not necessarily true, but it should be approximately true for most cases, especially if the size of the bin is much smaller than the whole dataset. In this case  $dx_i$  and  $dy_i$  is the same for every datapoint.

Whe have no data for density variation in the z-dicetion. Instead we assume a gaussian decrese of density with increasing distance from the midplane.  $\rho(z)$  is the assumed density at a point with altitude z above a point i in the midplane with density  $\rho_0$ .

$$\rho(x_i, y_i, z) = \rho_0(x_i, y_i) \cdot \exp\left(-\frac{z^2}{2H^2}\right)$$

Thus the mass integral has a discrete part and an analytical part. The limits of the discrete part is the area of the midplane  $S_{bin}$  contained in the sylinder and is such that  $N_{bin} dx_i dy_i = S_{bin}$ . The limits for z are explained in the following subsection.

$$M_{bin} = \int \int_{S_{bin}} \rho_0(x_i, y_i) \, dx_i \, dy_i \cdot \int_{z_{i,a}}^{z_{i,b}} \exp\left(-\frac{z^2}{2H^2}\right) \, dz$$
$$= \sum_i \left(\rho_{0,i} \, dx_i \, dy_i \cdot \frac{\sqrt{\pi}}{2} \sqrt{2H^2} \left[ \operatorname{erf}\left(\frac{z_{i,b}}{\sqrt{2H^2}}\right) - \operatorname{erf}\left(\frac{z_{i,a}}{\sqrt{2H^2}}\right) \right] \right)$$

#### 4.6.2 The *z*-limits

Each density point is given a weight according to where it is in the sylinder. Points closer to the middle of the sylinder gains larger weight because they represent its full height and thus a larger area than points near the edges. (**TODO:** There needs to be a figure of this to illustrate it more cleary.)

$$W_i(y) = \frac{\sqrt{r_{star}^2 - (y_i - y_{star})^2}}{\cos(\phi)}$$

The factor  $1/\cos(\phi)$  adjusts the height of the sylinder if it is inclined so that it is always shaped like a circular sylinder.

Figure 1: How 
$$z_{i,a}$$

To get the density inside the entire area of the slice of the sylinder and the variations in density from different altitudes we integrate the density for each point, projected from the bottom  $(z_a)$  to the top  $(z_b)$  of the sylinder. The distance to integrate is  $2W_i$  for each point, centered around  $z_i$ .

$$z_i = (x_i - x_{star}) \cdot \tan(\phi)$$
$$z_{i,a} = z_i - W_i$$
$$z_{i,b} = z_i + W_i$$

#### 4.7 Integrating intensity

For each bin j in each sylinder, from the inside to he outside of the disk, the ratio of intensity transferred from one bin to the next is given by the following expression.

$$\tau_j = \kappa \cdot \rho_{bin,j} \cdot \Delta r_j$$
$$I_{j+1} = I_j \exp(-\tau_j)$$

The resulting intensity passed on ny the outermost bin  $I_{end}$  is the intensity of the star's radiation that escapes the disk and is observed by the observer on the current line of sight. If there are several stars and thus several sylindres, the total perceived intensity is simply the sum of the  $I_{end,k}$  for each sylinder k.

$$I_{total} = \sum_{k} I_{k,end}$$

#### 4.8 Full algorithm summary

This is how one analysis is performed, and the product is one lightcurve. If providing different values for certain parameters, like different inclination angles or different outer radii then this analysis will be performed once for each different value of each parameter (different inclinations are actually analysed in quasi-parallell for efficiency).

```
for each \theta in [0,\ldots,2\pi] do rotate density datapoints angle \theta with R_z rotate stars angle \theta with R_z for each star k do extract sylinder bin sylinder for each bin j in sylinder do \rho_{bin,j} = \frac{\int \int \int_{V_{bin,j}} \rho(x,y,z) \; \mathrm{d}x \; \mathrm{d}y \; \mathrm{d}z}{\pi r_{star,k}^2 \Delta r_j} \tau_j = \kappa \cdot \rho_{bin,j} \cdot \Delta r_j I_{k,j+1} = I_{k,j} \exp(-\tau_j) end for end for I_{\theta,total} = \sum_k I_{k,end} end for
```

## 5 Configuring and running CirBinDis

How to make necessary configurations and then run CirBinDis to perform an analysis.

This is the most practical and maybe the most important section, as it explains how to actually use the software.

#### 5.1 Input parameters

The input parameters for each run of CirBinDis is configured in an XML file. Inside the repository you will find /xml/input.xml. Copy this file to the path where you will run CirBinDis from (normally the folder where the input datafile is) and change the value of the fields as required (it must still be called input.xml). Specifically, this is where you provide the filename of the dataset to analyse (if input.xml is in another folder than the input datafile you need to provide the relative path).

**TODO**: Here I plan to more or less copy the explanations that are in input.xml already.

unit-mass Dorem
unit-distance ipsum
unit-intensity dolum

## 5.2 Executing the code

When you have prepared input.xml and you are located in it's folder, type the command cirbindis

(or python ~/path\_to\_repository\_folder/circumbinarydisk/src/main.py). Then the software will run until it has completed the analysis of your datafile with all the parameters you specified.

## 5.3 Output

The output after a CirBinDis analysis is a comma-separated-value (csv) file. The output file will be placed in the path you specified in input.xml. The filename contains the value of the parameters H,  $r_{in}$ ,  $r_{out}$ , and inclination ( $\phi$ ), separated by double underscores "\_\_". A filename can be f.ex. "H=0.1\_r\_in=0.75\_r\_out=3\_inc=5.csv". If you perform several analysis of the same dataset at once, f.ex. by providing several values for  $r_{out}$ , then one outfile will be produced for each different value of  $r_{out}$ .

The first line of the output is a header containing all the physical and numerical parameters for the simulation. There are two columns. The first column lists rotation angles  $\theta$  in units of degrees. The second column lists the observed intensities

given the respective angles, normalised so the mean intensity is 1. A output file with azimuthsteps=8 can look like this:

```
#H=0.1,_kappa=10,_r_star=0.21-0.19,_r_in=0.75,_r_out=3,_dr=0.75,
dtheta=45deg,_inc=5deg
0.000000,0.000000
45.000000,0.000000
90.000000,0.000000
135.000000,0.000000
180.000000,7.644186
225.000000,0.000000
270.000000,0.000000
```

# 6 Bibliography

# 7 Acknowledgments

#### 8 Source code

#### 8.1 input.xml

The file for the user to provide input parameters to CirBinDis . You can copy and modify it. It is not a part of the source code itself.

```
<?xml version="1.0"?>
<!--This is a comment.-->
<!--With this file you can set all the input parameters for analysing a
   circumbinary disk. Make a copy of this file and place it in the directory
   you run the program from. Then you can just run 'python somepathname/main.py'. You can edit all the leaf nodes in this file,
   that is any field between two tags. F.ex. you can replace '0 10 30' with
    '5 15 25' in '<inclination' 10 30</inclination'. There are
   descriptions of the meanings of all the elements, and some elements have
   examples under them, where the different examples are separated with
   vertical | bars | like | this.
<input>
   <!---==Input units======--->
   <!--The units provided here are the units of the input values provided in this file, except for kappa. You can choose between the suggested
   units under each field (and other standard units). You can also use a
   quantity of a unit, f.ex. you can measure distance in units of '10.5
   solRad' and mass in units of '1e27 kg'. Output units are always in
   CGS-units and degrees.
   -->
   <unit>
       <mass>solMass</mass>
          <!--Examples: solMass | kg | g -->
       <distance>10.921 solRad</distance>
```

```
<!--Examples: AU | solRad | km | m | cm -->
   <intensity>erg s^-1 cm^-2</intensity>
      <!--Examples: erg s^-1 cm^-2 | J / (m2 s) -->
   <angle>deg</angle>
      <!--Examples: rad | deg | arcmin | arcsec -->
</11nit>
<!--The pathname to the dataset to be analysed. The pathname must be
   relative to where you run the program from, or it can be an absolute
   path. It can be an ACII file with (x,y,density) columns or a pickle
   file. If it is ASCII you must also specify if the coordinates are
   cartesian (x,y) or polar (r,theta).
<datafile>data/data_cropped.p</datafile>
<system>cartesian</system>
   <!--Examples: cartesian | polar -->
<!---==0utput data======-->
<!--The pathname to the folder to contain the output files. The pathname
   must be relative to where you run the program from, or it can be an
   absolute path. If the folder does not already exist it will be
   created automatically.
<outfolder>results</outfolder>
<!--Define the inner and out radius for where the disk exists. The
   dataset will be cropped to these limits, so it becomes a "donut".
   You can provide several inner and outer radiuses by separating them
   with spaces it you want to analyse different sized disks.
   If radius_in is not provided it will default to just outside the
   position \ of \ the \ stars. \ If \ radius\_out \ is \ not \ provided \ it \ will \ \frac{default}{}
   to the radius that defines the largest circle that can fit within
   the dataset.
<radius_in>1</radius_in>
   <!--Examples: 1 | 1.5 | 1 1.25 1.5 -->
<radius_out>3</radius_out>
   <!--Examples: 3 | 5 | 4 6 | 3 5 7.5 10 -->
<!--Which inclinations to analyse the system in. Several inclinations
   can be separated with spaces. If no inclinations is provided a
   default of 0 will be chosen.
<inclination>0 10 30</inclination>
   <!--Examples: 10 | 0 2.5 5 | 0 5 10 20 30 -->
<!--=====0ther physical disk parameters===============-->
<!--Total mass of disk, excluding the stars.
Opacity, in units of [cm^2/g].
   Thickness of disk.
<diskmass>0.01</diskmass>
<kappa>10.0</kappa> <!--[cm^2/g].-->
<H0>0.1</H0>
<! --There can be one or more stars with different positions, radiuses
   and intensities. They will be integrated separately and have their
   resulting fluxes added together.
   * If you want to add another star:
     Copy-paste a star element and change its parameters.
   * If you want to remove a star:
     Delete a star element.
   If no stars are provided a default star will be made at origo with
   radius=radius_in and solar luminosity.
   Provide EITHER cartesian (x,y) OR polar (r,theta) coordinates for
   each star.
```

```
-->
   <star>
       <position>
           <x>-0.5</x>
           <y>0</y>
           <r></r>
           <theta></theta>
       </position>
       <radius>0.21</radius>
       <intensity>1.1</intensity>
   </star>
   <star>
       <position>
           < x > 0.5 < /x >
           <y>0</y>
           <r></r>
          <theta></theta>
       </position>
       <radius>0.19</radius>
       <intensity>0.9</intensity>
   </star>
   <!--Example:
       <star>
           <position>
              <x></x>
               <y></y>
               <r>0.5</r>
               <theta>5</theta>
           </position>
           <radius>0.2</radius>
           <intensity>1.0</intensity>
       </star>
   <!--How many azimuthal rotation steps to use. This is the resolution of
       the resulting lightcurve.
       dtheta = 360deg / azimuthsteps
   <azimuthsteps>18</azimuthsteps>
   <!--How many radial steps to use in each line-of-sight integration. This
       defines dr and the accuracy of each fluxpoint in the resulting
       lightcurve.
       dr = (radius_out - radius_in) / radiussteps
   <radiussteps>10</radiussteps>
</input>
```

#### 8.2 main.py

The script that is called when running CirBinDis .

```
import xmltodict
from DensityMap import DensityMap
import Functions as func

if __name__ == "__main__":
    infile = open("input.xml", "r")
    input_ = xmltodict.parse(infile)["input"]
    infile.close()

for radius_in in func.to_list(input_["radius_in"], float):
```

```
for radius_out in func.to_list(input_["radius_out"], float):
    dataset = DensityMap(
       filename=input_["datafile"],
        coordsystem=input_["system"],
        unit=input_["unit"],
        inclinations=func.to_list(input_["inclination"], float),
        radius_in=radius_in,
       radius_out=radius_out
        diskmass=float(input_["diskmass"]),
        H=float(input_["H0"]),
        kappa=float(input_["kappa"]),
    )
    for star in func.to_list(input_["star"]):
        dataset.add_star(star)
    dataset.make_lightcurve(
       n_angle=int(input_["azimuthsteps"]),
        n_radius=int(input_["radiussteps"]),
        unit=input_["unit"]["angle"],
        show=True,
        save=True
        outfolder=input_["outfolder"],
   )
```

#### 8.3 DensityMap.py

Contains the class DensityMap for making an instance of a dataset representing a circumbinary disk. It contains most methods that can be performed on the data. Also contains a subclass Sylinder. Sylinders a sub-sets of a full dataset.

```
import sys
   # For doing meta things like receiving command-line arguments and exiting
   # the program.
import numpy as np
   # Numerical Python. Contains mathematical functions for performing
    # computations on arrays/matrices fast.
import cPickle as pickle
   # Can be used to save (dump) and load Python objects to files. Much
   # faster than reading and writing ASCII tables.
import time
   # Used to time parts of code to look for bottlenecks.
import matplotlib.pyplot as plt
   # For plotting results.
from scipy import integrate
from scipy import special
import astropy.units as u
from Star import Star
import Functions as func
class DensityMap:
   def __init__(self,
        data=None,
        filename=None,
        coordsystem="cartesian",
        outfolder=None,
        unit=None,
        inclinations = None,
        radius_in=0,
        radius_out=np.inf,
        diskmass=.01,
        H = 1.
```

```
kappa=10.
):
    self.data_rotated = None
    # If the inclination is a single number, put it in a list:
        iter(inclinations)
        self.inclinations = inclinations
    except TypeError:
        self.inclinations = [inclinations]
    self.outfolder = outfolder
    self.unit = unit
    self.stars = []
    self.radius_in = radius_in
    self.radius_out = radius_out
    self.diskmass = diskmass
    self.H = H
    self.kappa = kappa # [cm^2 / g]
        # Between 5 and 100 according to Boubier et al. 1999.
    if data is not None:
        self.data = data
    elif filename is not None:
        self.load(filename)
    # Convert density to physical units related to the total mass and
    # size of the disk:
    self.data[:, ~0] /= self.data[:, ~0].mean()
self.data[:, ~0] *= (
        self.diskmass / (
            np.pi * u.Quantity(50, "AU").to(
                u.Unit(self.unit["distance"])
            ).value**2 * 2*self.H
    if coordsystem == "cartesian":
    elif coordsystem == "polar":
        x, y = func.pol2cart(self.data[:, 0], self.data[:, 1])
        self.data[:, 0], self.data[:, 1] = x, y
        raise KeyError("Coordinate system must be 'cartesian' or 'polar'.")
def add_star(self, d=None, position=None, radius=None, intensity=None):
    """Make a Star instance and store it in a list of stars for the disk.
    d: (dictionairy) Must contain position, radius and intensity and can
        be provided instead of giving these other arguments individually.
    position: (float, array-like) Coordinates of the star.
    radius: (float) Radius of the star.
    intensity: (float) Intensity of the star.
    self.stars.append(Star(
        \verb|d=d|, position=position|, radius=radius|, intensity=intensity|
def load(self,
    filename.
    method=None,
separator=" ",
    """Load a dataset to analyse from a file.
    Assumed to be on the form 'x,y,density' or 'x,y,z,density' which
    represents a point in cartesian space and the density at that point.
    If the z-coordinate is not given it will be assumed to be 0 for \,
```

```
every point. Resulting data is an array of shape (N, 4), where N is
    the number of data points.
    filename: (string) Full pathname to file containing dataset.
    method: (string) What kind of loading algorithm to use. Can be
        'ascii' or 'pickle', If none is given, will try to automatically find out by looking at file ending.
    separator: (string) If method='ascii' this is the separator
        between the values each line. Usually a space or comma. Ignored if
        method='pickle'.
    t_start = time.time()
        # Just to time the loading, in case of large dataset.
    infile = open(filename, "r")
    if method is None:
        if filename.endswith(".p") or filename.endswith(".pickle"):
            method = "pickle"
        else:
            method = "ascii"
    if method == "pickle":
        data = pickle.load(infile)
        mask = (
            (np.linalg.norm(data[:, 0:2], axis=1) >= self.radius_in) *
            (np.linalg.norm(data[:, 0:2], axis=1) <= self.radius_out)</pre>
        self.data = data[np.where(mask)]
    elif method == "ascii":
        data = []
        for line in infile:
            line = line.rstrip().split(separator)
            if len(line) >= 3:
                line = [float(value) for value in line]
                if (self.radius_in <=</pre>
                    np.linalg.norm(line[0:2]) <=</pre>
                    self.radius_out
                    data.append(line)
        self.data = np.array(data)
    infile.close()
    t_end = time.time() # End of timer.
    print "Loading took %f seconds." % (t_end - t_start)
def writeto(self, filename, method="pickle", separator=" "):
    """Write self.data to a file for later use.
    Assumed to be on the form 'x,y,z,density' which represents a point in
    cartesian space and the density at that point.
    filename: (string) Full pathname to outfile for writing data.
    method: (string) What kind of writing algorithm to use. Recommended to
        use 'pickle' if it will be loaded by this program later (faster) and
        'ascii' for any other purpose.
    separator: (string) If method='ascii' this is the separator between the
    values each line. Usually a space or comma. Ignored if method='pickle'.
    t_start = time.time()
        # Just to time the writing, in case of large dataset.
    if method == "pickle":
        outfile = open(filename, "wb")
        pickle.dump(self.data, outfile)
    elif method == "ascii":
```

```
outfile = open(filename, "w")
        for line in self.data:
            outfile.write("%f%s%f%s%f\n" % (
                 line[0], separator,
                 line[1], separator,
                line[2],
            ))
    outfile.close()
    t_end = time.time() # End of timer.
    print "Writing took %f seconds." % (t_end - t_start)
def set_r0(self, r0=1.49597871e13):
    self.r0=r0 # [centimeters]
def set_H(self, H0=0.03, H_power=1/4.):
    self.H0 = H0
    self.H_power = H_power
def get_H(self, r):
    return r * self.H0 * (r / self.r0)**self.H_power # [centimeters]
def set_sigma(self, sigma0=1700., sigma_power=-3/2.):
    self.sigma0 = sigma0
    self.sigma_power = sigma_power
def get_sigma(self, r):
    return self.sigma0 * (r / self.r0)**self.sigma_power # [g / cm^2]
def rotate(self, angle_z=0, unit="deg"):
    """Rotate entire dataset by an angle around any axis.
    The original data is not changed. Rotated version of data stored in
    self.data_rotated. self.stars are also rotated.
    angle_z: (float) Angle to rotate around z-axis. This is the rotational
        axis for the disk. It can be gradually increased to simulate the
        orbital rotation of the system.
    angle_y: (float) Angle to rotate around y-axis. The inclination between
        the disk and the field of view. This angle should always be the same
        for one analysis if the disk is not wobbling.
    angle_x: The x-axis is the line of sight. Rotations around this axis
        would have no effect on the received flux, therefore this angle is
        ignored.
    unit: (string) What unit angles are given in.
       'rad', 'deg', 'arcmin' or 'arcsec'.
    # Transform angles into radians:
if unit == "rad":
       factor = 1.
    elif unit == "deg":
        factor = np.pi / 180.
    elif unit == "arcmin":
        factor = np.pi / 180. * 60
    elif unit == "arcsec":
       factor = np.pi / 180. * 3600
    angle_z *= factor
    # Make rotation matrix:
    rotation_matrix = np.matrix([
        [ np.cos(angle_z), -np.sin(angle_z)],
[ np.sin(angle_z), np.cos(angle_z)],
    # Rotate the disk:
    coords_in = self.data[:, :~0]
    coords_out = \
```

```
np.asarray(rotation_matrix * coords_in.transpose()).transpose()
    self.data_rotated = np.hstack((coords_out, self.data[:, ~0, None]))
    # Rotate the stars:
    for star in self.stars:
        star.position_rotated = np.asarray(
            rotation_matrix * star.position[:, None]
        ).transpose()[0]
def distance(self, p1, p2=None):
    """Returns the distances from the (rotated) datapoints to a line.
    p1: (float, array) A point in space which defines a line with p2.
    p2: (float, array) A point in space which defines a line with p1. If
        p2 is not provided it is assumed that the line is parallell to
        the x-axis.
    return: (float, array) The shortest euclidian distances between
    points and the line (p1, p2).
    if p2 is None:
        p2 = p1.copy()
        p2[0] += 1.
    return np.abs(
        np.cross(
           p1 - self.data_rotated[:, :~0],
p2 - self.data_rotated[:, :~0],
        np.linalg.norm(p2 - p1),
def get_sylinder(self, starno=None, star=None):
     ""Slice out a sylinder shape from the data based on a star.
    The sylinder is always oriented along the x-axis. Its size and
    position is determined by a star. It spans from the position of the
    surface of the star until x=\inf.
    starno: (int) Index to get star from self.stars. Ignored if star is
       given.
    star: (Star instance) A star to base the sylinder on.
    return: (float, array) The slice of the dataset contained in the
    sylinder.
    if star is None:
        star = self.stars[starno]
    mask = (
        (self.data_rotated[:, 0] > 0) *
        (self.distance(star.position) <= star.radius)</pre>
    data_sylinder = self.data_rotated[np.where(mask)]
    return data_sylinder
def get_density_profile(self, sigma=1, skip=2000, show=True):
    """Returns and displays the density profile of self.data.
    TODO: Finish this docstring.
    from scipy.ndimage.filters import gaussian_filter1d
    radiuses = np.linalg.norm(self.data[:, 0:2], axis=1)
    indices_sorted = np.argsort(radiuses)
```

```
radiuses = radiuses[indices_sorted][::skip]
    densities = gaussian_filter1d(
        self.data[:, 3][indices_sorted],
        sigma=sigma,
       mode="nearest",
   )[::skip]
    if show:
        plt.plot(radiuses, densities, "b+")
        plt.xlabel("radius")
       plt.ylabel("density")
       plt.show()
    return radiuses, densities
def make_lightcurve(self,
    inclinations = None,
   H=None,
    n_angle=None,
    dtheta=None,
    theta=None,
    unit="deg",
   n_radius=None,
    dr=None,
    save=False,
    show=False,
    outfolder=None,
    """Makes a lightcurve by calling the other methods for each orientation
    of the dataset. Sort of a main method.
    TODO: Complete docstring.
    # Default setting for theta is a full revolution:
    if theta is None:
       if unit == "rad":
            theta = 2*np.pi
        elif unit == "deg":
            theta = 360.
        elif unit == "arcmin":
           theta = 360. * 60
        elif unit == "arcsec":
            theta = 360. * 3600
    if inclinations is None:
       inclinations = self.inclinations
    \mbox{\tt\#} If the inclination is a single number, put it in a list:
    inclinations = func.to_list(inclinations)
    if H is None:
       H = self.H
    if n_angle is None:
       n_angle = int(round(float(theta) / dtheta))
    elif dtheta is None:
        dtheta = float(theta) / n_angle
    angles = np.linspace(0, theta-dtheta, n_angle)
    lightcurve = np.zeros((len(inclinations), n_angle))
    for i, angle in enumerate(angles):
        print "%f / %f" % (angle, theta)
        self.rotate(
            angle_z=angle,
            unit=unit,
        for k, star in enumerate(self.stars):
            sylinder = Sylinder(
```

```
star=star,
             data=self.get_sylinder(star=star),
             unit=self.unit,
             radius_in=self.radius_in,
             radius_out=self.radius_out,
             kappa=self.kappa
        for j, inclination in enumerate(inclinations):
             sylinder.space_sylinder(
                 inclination=inclination,
                 unit=unit,
                 H=H,
                 n_steps=n_radius,
                 dr=dr,
             lightcurve[j, i] += sylinder.integrate()
print "%f / %f" % (theta, theta)
lightcurve /= lightcurve.mean(axis=1)[:, None]
for j, inclination in enumerate(inclinations):
    starradius = ""
    for star in self.stars:
        starradius += "%g-" % star.radius
    starradius = starradius.rstrip("-")
    header = (
        "H=%g, kappa=%g, "
        "r_star=%s, r_in=%g, r_out=%g, dr=%g, "
        "dtheta=%g%s, inc=%g%s"
        % (Н,
             self.kappa,
             starradius,
             self.radius_in,
             self.radius_out,
             (self.radius_out - self.radius_in) / n_radius,
             float(theta) / n_angle,
             unit,
             inclination,
             unit,
        )
    if save:
        outname = (
             "H=%g__"
"r_in=%g__r_out=%g__"
             "inc=%g"
             % (Н,
                 self.radius_in,
                 self.radius_out,
                 inclination,
             )
        if outfolder is None:
             outfolder = self.outfolder
        func.make_folder(outfolder)
        outfile = open("%s/%s.csv" % (outfolder, outname), "w")
        outfile.write("#" + header + "\n")
for angle, flux in zip(angles, lightcurve[j]):
    outfile.write("%f,%f\n" % (angle, flux))
        outfile.close()
    if show:
        plt.plot(
             angles,
             lightcurve[j],
             label="inc=%g" % inclinations[j],
if show:
```

```
plt.title(header)
            plt.xlabel("rotational angle [degree]")
            plt.ylabel("relative intensity")
            plt.legend()
            plt.show()
class Sylinder(DensityMap):
   def __init__(self,
        star,
        data.
        unit=None,
        inclinations=None,
        radius_in=0,
        radius_out=np.inf,
        kappa=10.
   ):
        # If the inclination is a single number, put it in a list:
           iter(inclinations)
            self.inclinations = inclinations
        except TypeError:
           self.inclinations = [inclinations]
        self.star = star
        self.unit = unit
       self.radius_in = radius_in
self.radius_out = radius_out
        self.kappa = kappa # [cm^2 / g]
        self.data = data
   def space_sylinder(self,
        inclination=0,
        unit="deg",
       H=1.,
        n_steps=None,
        dr=None,
        """Bin this sylinder's datapoints into a set of mean densities.
        The sylinder is first sorted along the x-axis and is then cut along
        the x-axis like a loaf of bread. Each point is integrated
        analytically through its projected density from the bottom to the
        top of the sylinder. The mean density is then computed from each
        slice of the sylinder/bread.
        This is stored temporarily as self.densities and self.drs: (float,
        array), (float, array) A list of mean densities and the
        corresponding list of delta radiuses for each bin. Both are arrays
        of length n\_step. The arrays are order FROM inside of disk TO
        oustide of disk.
        inclination: (float) The angle to incline the line of sight on the
            svlinder.
        deg: (string) Unit of the angle.
        H: (float) Thickness of the disk. Necessary for integral.
        n_steps: (int) How many slices to divide the sylinder in. Affects
           accuracy of integral.
        dr: (float) The width of each sylinder section. Ignored if n_steps
            is provided.
        if unit == "rad":
            factor = 1.
```

```
elif unit == "deg":
       factor = np.pi / 180.
    elif unit == "arcmin":
        factor = np.pi / 180. * 60
    elif unit == "arcsec":
        factor = np.pi / 180. * 3600
    inclination *= factor
    if n_steps is None:
       n_steps = int(round((self.radius_out - self.radius_in) / dr))
    dpoints = int(round(self.data.shape[0] / float(n_steps)))
        # How many datapoints to include in each bin.
    densities = np.zeros(n_steps)
    drs = np.zeros(n_steps)
    data = self.data[np.argsort(self.data[:, 0])]
    g = np.sqrt(2) * H # Constant used several times in calculations.
    for i in xrange(n_steps):
        start = i*dpoints
        if i == n_steps-1:
            # If it is the last step, make sure the last few points are
            # included (in case there are some rounding problems).
            end = data.shape[0]
            drs[i] = data[end-1, 0] - data[start, 0]
            s = data[start:end].shape[0]
            drs[i] *= (s + 1.) / s
        else:
            end = (i+1)*dpoints
            drs[i] = data[end, 0] - data[start, 0]
        W = np.sqrt(
            self.star.radius**2 -
            (data[start:end, 1] - self.star.position[1])**2
        ) / np.cos(inclination)
        z = (
            (data[start:end, 0] - self.star.position[0]) *
            np.tan(inclination)
        z1 = z - W
        z2 = z + W
        densities[i] = (
            np.sum(
                # \int_z1^z2 \rho_0 * e^{- z^2 / (2*H^2)} dz
g * data[start:end, ~0] * 0.5 * np.sqrt(np.pi) *
                (special.erf(z2 / g) - special.erf(z1 / g))
            ) / (2. * np.sum(W))
    self.densities = densities
    self.drs = drs
def integrate(self):
    """Integrates the intensity through the layers of dust.
    Assumes that space_sylinder has just been called and used its results.
    return: (float) Perceived intensity outside the disk
    kappa = self.kappa * u.Unit("cm2/gram").to(
        u.Unit(self.unit["distance"])**2 / u.Unit(self.unit["mass"])
    intensity = self.star.intensity
    for density, dr in zip(self.densities, self.drs):
        tau = kappa * density * dr
        intensity *= np.exp(-tau)
```

```
return intensity * (u.Unit(self.unit["intensity"])).to("erg / (cm2 s)")
```

#### 8.4 Star.py

The simple Star class whose intention is to hold the physical parameters of each star.

```
import numpy as np
import Functions as func
class Star:
    def __init__(self, d=None, position=None, radius=None, intensity=None):
    """Make a star instance.
        \ensuremath{\mathtt{d}} \colon (dictionairy) Must contain position, radius and intensity and can
            be provided instead of giving these other arguments individually.
        position: (float, array-like) Coordinates of the star.
        radius: (float) Radius of the star.
        intensity: (float) Intensity of the star.
        if d is not None:
            try:
                 position = np.array([
                     float(d["position"]["x"]),
                     float(d["position"]["y"]),
             except KeyError, ValueError:
                 position = np.array(func.pol2cart(
                     float(d["position"]["r"]),
                     float(d["position"]["theta"]),
                 ))
             radius = float(d["radius"])
            intensity = float(d["intensity"])
        self.position = np.array(position)
        self.radius = radius
        self.intensity = intensity
```

#### 8.5 Functions.py

A file containing some general functions that are used other places in the program.

```
import os
import numpy as np

def to_list(x, dtype=None, separator=" "):
    """Converts any sequence or non-sequence into an array.

The use of a numpy array is primarely because then it is easy to also convert the type. Except for that it couls just as well be a list.

If x is a string, split the string with the separator.
    If x is a sequence, convert it into an array.
```

```
If x is a non-sequence, but it into a size-1 array.
   {\tt X:} (anything) Something to be converted into an array.
   dtype: (type) What type of objects the array contains. F.ex. float. If
       None, numpy will interpret the type itself.
   separator: (string) If x is a list to be splitted, this is the
       separator. Normally a space or comma.
   if isinstance(x, basestring): # If x is a string.
       x = np.array(x.split(separator), dtype=dtype)
       try:
           iter(x) # If x is a sequence.
           x = np.array(x, dtype=dtype)
       except TypeError: # If x is a non-sequence.
           x = np.array([x], dtype=dtype)
   return x
def cart2pol(x, y):
    """Convert cartesian coordinates into polar coordinates."""
   r = np.sqrt(x**2 + y**2)
   theta = np.arctan2(y, x)
   return [r, theta]
def pol2cart(r, theta):
     ""Convert polar coordinates into cartesian coordinates."""
   x = r * np.cos(theta)
   y = r * np.sin(theta)
   return [x, y]
def make_folder(directory, warning=False):
   """Make the directory, but do nothing if it already exists. Optionally
   print a warning if it already exists, if there should be a reason for
   that.
   if not os.path.exists(directory):
       os.makedirs(directory)
   else:
       if warning:
           print "%s already exists." % (directory)
```

#### 8.6 plot.py

A standalone script that can be used to plot the output of CirBinDis . You can just as well use f.ex. TOPCAT.

```
import sys
import numpy as np
import matplotlib.pyplot as plt

def load(filename, inclination, separator=","):
    """Load a dataset to plot from a file.

    Assumed to be on the form 'angle,flux'.
    The first 3 lines are the title, xlabel and ylabel.

filename: (string) or (string, list) Full pathname to file containing
    dataset or a list of pathnames.
separator: (string) This is the separator between the values each line.
    Usually a space or comma.
```

```
if isinstance(filename, basestring):
       filenames = [filename]
    else:
            iter(filename)
            filenames = filename
        except TypeError:
            raise TypeError(
                "'filename' must be a string or sequence of strings."
    if isinstance(inclination, basestring):
       inclinations = inclination.split(" ")
    else:
            iter(inclination)
            inclinations = inclination
        except TypeError:
            raise TypeError(
                "'filename' must be a string or sequence of strings."
   infiles = [open(filename, "r") for filename in filenames]
   titles = [infile.readline() for infile in infiles]
   xlabels = [infile.readline() for infile in infiles]
   ylabels = [infile.readline() for infile in infiles]
   title = titles[0]
   xlabel = xlabels[0]
   ylabel = ylabels[0]
   i = 0
    \begin{tabular}{ll} for infile in infiles: \\ \end{tabular} 
        data = []
        for line in infile:
            line = [float(value) for value in line.split(separator)]
            data.append(line)
        angles, fluxes = np.array(data).T
        infile.close()
        plt.plot(angles, fluxes, label="inc="+inclinations[i])
        i += 1
   plt.title(title)
   plt.xlabel(xlabel)
   plt.ylabel(ylabel)
   plt.legend()
   plt.show()
if __name__ == "__main__":
   load(filename=sys.argv[2:], inclination=sys.argv[1])
```

#### 8.7 make\_testdata.py

A standalone script that can be used to generate artificial datasets that can be analysed by CirBinDis . You can use it for testing and for generating data according to any analytical function that you would like to analyse (then you need to change the function density).

```
"""This script can be used to make an artificial density map with a smooth sinusoidal shape. This can be used to test the analysis program.
```

```
import numpy as np
import cPickle as pickle
Nx = int(6e2 + 1)
Ny = int(6e2 + 1)
strip_x = np.linspace(-4, 4, Nx)
strip_y = np.linspace(-4, 4, Ny)
x = np.zeros(Nx*Ny)
y = np.zeros(Nx*Ny)
z = np.zeros(Nx*Ny)
for i in xrange(Ny):
to i in xrange(Ny).
    x[i*Nx : (i+1)*Nx] = strip_x
for i in xrange(Nx):
    y[i :: Nx] = strip_y
def density(x, y):
    return (1. + np.sin(2*np.arctan2(y, x)))
def add_density(x, y):
    return np.vstack((x, y, z, density(x, y))).transpose()
data = add_density(x, y)
outfile = open("../data/testdata.p", "wb")
pickle.dump(data, outfile)
outfile.close()
```