CO31: Structure of White Dwarf Stars

Guo-Zheng Theodore Yoong University College, Oxford univ4499

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Abstract

This computing report investigates the equations of state of white dwarf stars. The equations of state are mainly governed by two coupled first-order ordinary differential equations on dm/dr and $d\rho/dr$. The equations of state of white dwarf stars for both non-relativistic and relativistic cases are numerically solved via the fourth-order Runge-Kutta method in Python. The results are also compared and discussed. The critical mass of white dwarf stars, i.e. the Chandrasekhar mass M_C , is also computed.

1 Introduction

In astrophysics, white dwarf stars are one possible evolutionary state at the end of a stellar evolution cycle. These celestial objects generally comprise heavy nuclei, mainly ⁵⁶Fe, although ¹²C may dominate if the nucleosynthesis stops prematurely [1].

The compression forces on the star due to its own gravitational field balance the forces due to the electron degeneracy pressure. The equations of state thereby lead to two coupled first-order differential equations for both the non-relativistic and relativistic treatments of the star. We therefore perform the fourth-order Runge-Kutta method to solve our differential equations and obtain the physical parameters of the star, such as the star's mass and radius.

2 Physical Background

2.1 Stellar Evolution

The stellar birth of low-mass stars begins in protostellar clouds [3]. At this stage, the gravitational force of the dust cloud pulls the dust particles inwards while releasing gravitational potential energy in the form of heat. The temperature of the dust core increases until a critical temperature is reached, in which nuclear fusion of hydrogen atoms begins, which drives stellar evolution.

The dust cloud takes the form of a sphere and a protostar is formed. Further accretion of dust to the core increases the mass and size of the protostar and it becomes a main sequence star. Protostars with insufficient mass will form brown dwarf stars instead of main sequence stars. Different main sequence stars will experience nuclear burning at different rates, and hence different lifetimes. Once the hydrogen fusion at the core of stars is complete, the thermal pressure resisting compression will subside, leading to contraction of the core.

For a low-mass star $(M \lesssim 0.5 M_{\odot})$, the temperature at its outer hydrogen shell cannot sustain fusion, and it thus gradually collapses into a white dwarf as it expels planetary nebulae. For a middle-mass star $(0.5 M_{\odot} \lesssim M \lesssim 8 M_{\odot})$, hydrogen at the outer core will fuse, generating thermal pressure to expand the star into a red giant with a helium core. Eventually, its core will collapse until the electron degeneracy pressure is sufficient to resist further compression. It will then radiate its remnant heat as a white dwarf.

For a high-mass star $(M \gtrsim 8M_{\odot})$, the hydrogen outer shell fuses quickly. Once exhausted, the core must be supported by electron degeneracy pressure alone. Eventually, the contraction stops as the density becomes too large, causing a supernovae explosion. The remnant core is a neutron star, which can collapse further into a black hole if the mass of the neutron star $M \gtrsim 3M_{\odot}$.

In this practical, we focus on white dwarf stars, i.e. the final stage of stars with $M \lesssim 8M_{\odot}$.

2.2 Equations of Equilibrium

In mechanical equilibrium, assuming spherical symmetry, the gravitational force on an element of the star is balanced by its interior pressure [1].

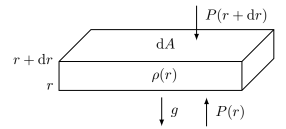


Figure 1: The balance of forces on an element of the star

If the area of element is dA and its density $\rho(r)$ is assumed to be uniform, then its mass m(r) is ρ dA and its weight is $Gm(r)\rho(r)$ dA dr/r^2 . The difference in pressure between r and r + dr is equal to the weight per unit area of the element. We can write this as

$$\frac{\mathrm{d}P(r)}{\mathrm{d}r} = \frac{P(r+\mathrm{d}r) - P(r)}{\mathrm{d}r} = -\frac{G\rho(r)m(r)}{r^2},\tag{1}$$

where G is the gravitational constant, m(r) is the mass of star interior to r, and that the negative sign indicates the pressure increases as r decreases. Applying the chain rule, we rewrite equation (1) as

$$\frac{\mathrm{d}\rho}{\mathrm{d}r} = -\left(\frac{\mathrm{d}P}{\mathrm{d}\rho}\right)^{-1} \frac{G\rho(r)m(r)}{r^2}.$$
 (2)

We can also express the derivative of m(r) with respect to r as

$$\frac{\mathrm{d}m}{\mathrm{d}r} = 4\pi r^2 \rho(r). \tag{3}$$

The goal of this practical is to solve the two coupled first-order ordinary differential equations (2) and (3). However, $dP/d\rho$ will be different for relativistic and non-relativistic electrons. We explore the derivations and physical consequences of both regimes.

2.3 Equations of State

Assume that the star is made up of large ⁵⁶Fe nuclei, which contribute almost all the mass but no pressure, and their electrons, which contribute almost all the pressure but no mass. We therefore model the star as a sphere with a density much larger than normal matter, and assume that the electrons are unbound and

act like a free Fermi gas. Using this model for the electron gas at 0 K [1], the number density of electrons can be expressed as

$$n = \int_0^{p_F} \frac{8\pi}{h^3} p^2 \, \mathrm{d}p = \frac{8\pi}{3h^3} p_F^3,$$

where h Planck's constant. The pressure is then given by

$$P = \frac{8\pi}{3h^3} \int_0^{p_F} p^3 v_p \, \mathrm{d}p \,, \tag{4}$$

where v_p is velocity of electron at a given momentum.

2.3.1 Non-relativistic Treatment

For $v_p \ll c$, we use $v_p = p/m_e$. The electron degeneracy pressure is then

$$P = \frac{8\pi}{15h^3 m_e} p_F^5 = \frac{1}{20} \left(\frac{3}{\pi}\right)^{\frac{2}{3}} \frac{h^2}{m_e} n^{\frac{5}{3}}.$$
 (5)

Except for hydrogen (which is not present in white dwarves), each electron is accompanied by a proton and very nearly one neutron in a electrically neutral atom. The density in the electrically neutral atom is therefore $\rho = n_e(m_e + m_p + m_n)$. Assuming that $m_e \ll m_p, m_n$, and $m_p \approx m_n$, we have $n_e \sim \rho/2m_p$. Substituting our approximation into equation (5) for n_e , we obtain

$$P = \frac{1}{20} \left(\frac{3}{\pi} \right)^{\frac{2}{3}} \frac{h^2}{m_e} \left(\frac{\rho}{2m_p} \right)^{\frac{5}{3}},$$

and hence

$$\frac{\mathrm{d}P}{\mathrm{d}\rho} = \frac{1}{48} \frac{h^2}{m_e} \frac{2^{\frac{1}{3}}}{m_e^{\frac{5}{3}}} \left(\frac{3\rho}{\pi}\right)^{\frac{2}{3}}.$$
 (6)

Substituting equation (6) into equation (2), we get

$$\frac{\mathrm{d}\rho}{\mathrm{d}r} = -k\frac{\rho^{\frac{1}{3}}m}{r^2}, \quad \text{where } k = \frac{48\pi^{\frac{2}{3}}}{2^{\frac{1}{3}}3^{\frac{2}{3}}} \frac{Gm_e m_p^{\frac{5}{3}}}{h^2}.$$
 (7)

2.3.2 Relativistic Treatment

In this regime, we use $v_p = pc^2/\sqrt{p^2c^2 + m_e^2c^4}$ instead. Substituting this expression into equation (4) yields

$$P = \frac{8\pi c^2}{3h^3} \int_0^{p_F} \frac{p^4}{\sqrt{p^2 c^2 + m_e^2 c^4}} \, \mathrm{d}p.$$
 (8)

Using the substitution $p = m_e c \sinh \theta$, we have $dp = m_e c \cosh \theta d\theta$. Equation (8) becomes

$$P = \frac{8\pi m_e^4 c^5}{3h^3} \int_0^{\theta_F} \sinh^4 \theta \, d\theta \,, \quad \text{where } \theta_F = \sinh^{-1} \frac{p_F}{m_e c}. \tag{9}$$

Substituting the relation $\sinh 4\theta = \frac{1}{8}\cosh 4\theta - \frac{1}{2}\cosh 2\theta + \frac{3}{8}$ into equation (9), we can easily perform the integral to obtain

$$P = \frac{8\pi m_e^4 c^5}{3h^3} \left(\frac{1}{32} \sinh 4\theta - \frac{1}{4} \sinh 2\theta + \frac{3}{8} \theta_F \right). \tag{10}$$

The chain rule for $dP/d\rho$ gives

$$\frac{\mathrm{d}P}{\mathrm{d}\rho} = \frac{\mathrm{d}P}{\mathrm{d}\theta_F} \frac{\mathrm{d}\theta_F}{\mathrm{d}p_F} \frac{\mathrm{d}p_F}{\mathrm{d}\rho}.$$

Noting that $p_F = (3h^3n_e/8\pi)^{\frac{1}{3}}$ and using the same approximation of $n_e \sim \rho/2m_p$, we have

$$\begin{split} \frac{\mathrm{d}P}{\mathrm{d}\theta_F} &= \frac{8\pi c}{3h^3} p_F^4, \\ \frac{\mathrm{d}\theta_F}{\mathrm{d}p_F} &= \frac{1}{\sqrt{p_F^2 + m_e^2 c^2}}, \\ \frac{\mathrm{d}p_F}{\mathrm{d}\rho} &= \frac{1}{2^{\frac{4}{3}} 3^{\frac{2}{3}} \pi^{\frac{1}{3}}} \frac{h \rho^{\frac{2}{3}}}{m_n^{\frac{1}{3}}}. \end{split}$$

Putting these all together and substituting into equation (2), we obtain

$$\frac{\mathrm{d}\rho}{\mathrm{d}r} = -\frac{k\sqrt{a+b\rho^{\frac{2}{3}}}}{r^2}\rho^{\frac{1}{3}}m,\tag{11}$$

where

$$k = 2^{\frac{7}{3}} 3^{\frac{1}{3}} \pi^{\frac{1}{3}} \frac{G m_p^{\frac{4}{3}}}{h^2 c},$$

$$a = 2^{\frac{8}{3}} \pi^{\frac{2}{3}} m_e^2 m_p^{\frac{2}{3}} c^2,$$

$$b = 3^{\frac{2}{3}} h^2$$

We have now obtained the forms of the coupled differential equations for both the non-relativistic and relativistic regimes. The Runge-Kutta method will be used to solve both cases, as outlined in the subsequent sections.

3 Runge-Kutta Method

3.1 Single Differential Equation

Consider the differential equation dy/dt = f(y,t). Taking the integral with respect to t from y_n to y_{n+1} , we obtain

$$\int_{y_n}^{y_{n+1}} \frac{\mathrm{d}y}{\mathrm{d}t} \, \mathrm{d}t = \int_{y_n}^{y_{n+1}} f(y, t) \, \mathrm{d}t \,. \tag{12}$$

We can convert (12) to a second-order Runge-Kutta method via Simpson's method with step size h, as outlined in [2], which gives

$$\int_{y_n}^{y_{n+1}} f(y,t) dt = \frac{h}{6} [f(y_n, t_n) + 4f(y_{n+\frac{1}{2}}, t_{n+\frac{1}{2}}) + f(y_{n+1}, t_{n+1}) + \dots] + \mathcal{O}(h^5).$$
 (13)

To improve this to fourth-order, we require four calculations of f(y,t) per iteration, which are

$$k_1 = f(y_n, t_n),$$

$$k_2 = f(y_n + \frac{h}{2}k_1, t_n + \frac{h}{2}),$$

$$k_3 = f(y_n + \frac{h}{2}k_2, t_n + \frac{h}{2}),$$

$$k_4 = f(y_n + hk_3, t_n + h),$$

and therefore

$$y_{n+1} = y_n + \frac{h}{6}(k_1 + 2k_2 + 2k_3 + k_4) + \mathcal{O}(h^5).$$
(14)

We then let $t_n \leftarrow t_n + h$ and $y_n \leftarrow y_{n+1}$, and continue iterating.

3.2 Coupled Differential Equations

We can generalise the fourth-order Runge-Kutta method to two coupled first-order ordinary differential equations. Let dx/dt = f(x, y, t) and dy/dt = g(x, y, t). Similar to what we did in the previous section, we have

$$\begin{aligned} k_1 &= f(x_n, y_n, t_n), \\ l_1 &= g(x_n, y_n, t_n), \\ k_2 &= f(x_n + \frac{h}{2}k_1, y_n + \frac{h}{2}l_1, t_n + \frac{h}{2}), \\ l_2 &= g(x_n + \frac{h}{2}k_1, y_n + \frac{h}{2}l_1, t_n + \frac{h}{2}), \\ k_2 &= f(x_n + \frac{h}{2}k_2, y_n + \frac{h}{2}l_2, t_n + \frac{h}{2}), \\ l_3 &= g(x_n + \frac{h}{2}k_2, y_n + \frac{h}{2}l_2, t_n + \frac{h}{2}), \\ k_4 &= f(x_n + hk_3, y_n + hl_3, t_n + h), \\ l_4 &= f(x_n + hk_3, y_n + hl_3, t_n + h). \end{aligned}$$

Similar to what we had in equation (14),

$$x_{n+1} = x_n + \frac{h}{6}(k_1 + 2k_2 + 2k_3 + k_4) + \mathcal{O}(h^5), \tag{15}$$

$$y_{n+1} = y_n + \frac{h}{6}(l_1 + 2l_2 + 2l_3 + l_4) + \mathcal{O}(h^5).$$
(16)

Once again, we let $t_n \leftarrow t_n + h$, $x_n \leftarrow x_{n+1}$, and $y_n \leftarrow y_{n+1}$ and continue iterating.

The Runge-Kutta can be applied to any number of coupled first-order ordinary differential equations. For example, if we have three coupled first-order differential equations, we let dx/dt = f(x, y, z, t), dy/dt = g(x, y, z, t), dz/dt = h(x, y, z, t) and solve them in a similar fashion by letting $k_1 = f(x_n, y_n, z_n, t_n)$, $l_1 = g(x_n, y_n, z_n, t_n)$ and $m_1 = h(x_n, y_n, z_n, t_n)$, iterate towards k_4 , l_4 and m_4 , update x_n , y_n and z_n , and repeat.

3.3 Error Considerations

We have used Simpson's Rule throughout the derivation, resulting in a local truncation error of order $\mathcal{O}(h^5)$. This results in a global accumulated error of $\mathcal{O}(h^4)$. We will observe later in our computational analysis that $\rho(r)$ and m(r) vary slowly with r, and hence the fourth derivative is very small. The Runge-Kutta method therefore provides a good approximation to the coupled first-order ordinary differential equations.

It is also worth noting that the time complexity of the Runge-Kutta method is $\mathcal{O}(n)$. For astrophysical problems which require many iterations, the linear time can be computationally taxing.

4 Computational Analysis

For this practical, we use Python as our programming language of choice. We create three functions outlined by [1]. The first is a generic fourth-order Runge-Kutta method solver, which has been customised slightly for the case of our coupled differential equations. The remaining two define the relevant differential equations for both the non-relativistic and relativistic cases. The step size we choose is $\delta r = 5000$ m.

4.1 Non-relativistic case

For the non-relativistic case, we choose an input central density $\rho_c = 10^{14} \text{ kg m}^{-3}$, and solve for $\rho(r)$ and m(r). To avoid any singularities, we let ρ_c be measured at 10 m $\ll 5 \times 10^7$ m. The plots of $\rho(r)$ and m(r) are shown in Figure 2.

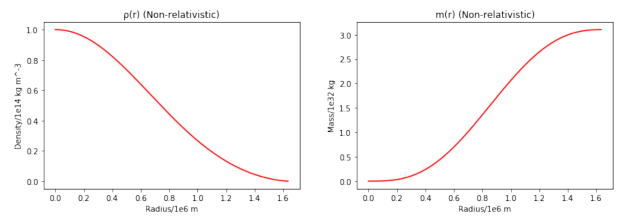


Figure 2: Plots of $\rho(r)$ and m(r) for the non-relativistic case

We notice that using numpy is to our advantage, as an input r of order 5×10^7 m for the non-relativistic case will result in a RuntimeWarning error being raised due to an underflow with double_scalars. This is because the density goes to zero at a point long before 5×10^7 m. The values beyond that are NaN, and are not present in the plots. Overall, we can conclude that for a star with $\rho_c = 10^{14}$ kg m⁻³, it has overall radius $R \sim 1.6 \times 10^6$ m and total mass $M \sim 3.2 \times 10^{32}$ kg.

We are also interested in the total mass M and overall radius R as a function of ρ_c for $10^6 < \rho_c < 10^{14}$ kg m⁻³. We sample 50 points evenly spaced along the log ρ_c axis and obtain their corresponding masses and radii. This time, to filter out the NaN values, we perform the list comprehension [val for val in nonrel_mass if val == val] as NaN has the property NaN != NaN. Maintaining our log-scale, the corresponding plots are shown in Figure 3.

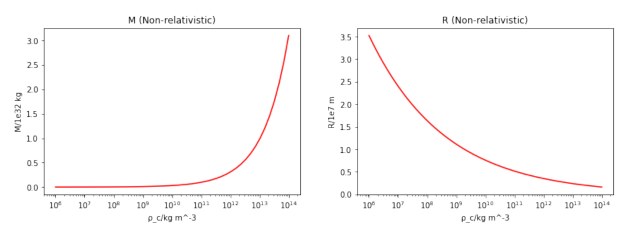


Figure 3: Plots of M and R for the non-relativistic case

As we can see in the plots, with increasing central density, the total mass and radius of the white dwarf increases and decreases respectively. However, there is no upper limit to M, which is not consistent with astronomical data and the Chandrasekhar limit, which we discuss in the next section. This is where the relativistic considerations come into play.

4.2 Relativistic case

Our other function adjusts for our relativistic $d\rho/dr$. Our input ρ_c is the same as the non-relativistic case. The new plots of $\rho(r)$ and m(r) are shown in Figure 4.

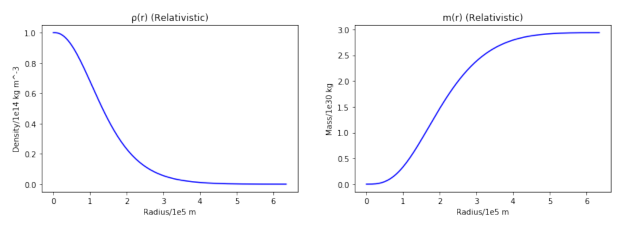


Figure 4: Plots of $\rho(r)$ and m(r) for the relativistic case

This time, our plots tell us that for a star with $\rho_c = 10^{14}$ kg m⁻³, it has overall radius $R \sim 6.4 \times 10^5$ m and total mass $M \sim 3.0 \times 10^{30}$ kg. These values are a few orders of magnitude smaller than what we obtained in the previous section. We also perform an identical analysis for M and R as a function of ρ_c in the same limits for the relativistic case. The new plots of M and R are shown in Figure 5.

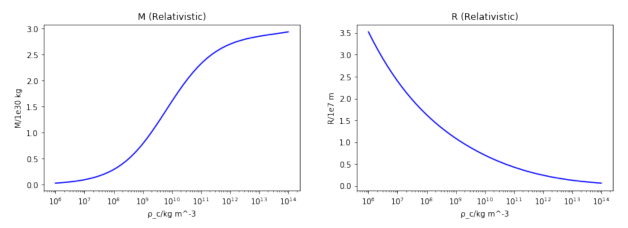


Figure 5: Plots of M and R for the relativistic case

This time, we observe a clear upper limit for M as ρ_c increases. This is consistent with the Chandrasekhar limit. If the core density is large enough such that the electron degeneracy pressure is insufficient to maintain hydrostatic equilibrium, the star collapses. This implies the existence of black holes and that no white dwarf can exist with a mass higher than this limit, which is the Chandrasekhar limit M_C . To observe this, we superpose and compare our plots.

4.3 Comparisons

We compare the plots of M and R against ρ_c for both cases as shown in Figure 6. Note that we use a log scale for M instead, due to the difference in order of magnitude.

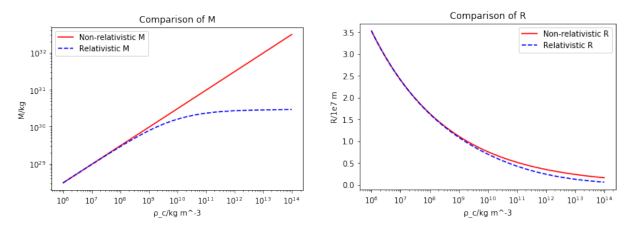


Figure 6: Comparisons of M and R

The theoretical Chandrasekhar mass is $M_C = 2.765 \times 10^{30}$ kg. By taking the last value of our rel_total_mass array, our computationally determined $M_C = 2.935 \times 10^{30}$ kg, which deviates from the theoretical value by roughly 6%.

We also plot M as a function of R for both comparisons, which also allow us to see the Chandrasekhar limit clearly.

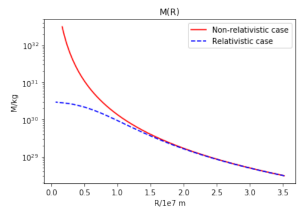


Figure 7: Comparison of M(R)

We first notice that for both cases, M is larger for smaller R, which represents a stronger gravitational pull due to the denser core and smaller electron degeneracy pressure. More importantly, we once again observe the Chandrasekhar limit for the relativistic case.

5 Conclusion

We have shown that the structure of white dwarf stars can be modelled using hydrostatic equilibrium and electron degeneracy pressure considerations. The two coupled first-order differential equations can be solved with the fourth order Runge-Kutta method. We then use our Runge-Kutta method solver for both the non-relativistic and relativistic cases, and note that the relativistic case enforces the Chandrasekhar

limit. We obtain this limit to be $M_C = 2.935 \times 10^{30}$ kg, with an error of around 6%. However, this error is large from an empirical perspective. We have introduced many sources of error, such as capping the order of the Runge-Kutta method, capping the step size, and allowing underflow. Nevertheless, the fourth order Runge-Kutta method we have employed is remarkably accurate, and our function can be modified to solve any number of coupled first-order ODEs.

References

- [1] University of Oxford, Physics Practical Course Laboratory Scripts, CO31, 2018
- [2] Numerical Methods for Physicists, A. OHare, Oxford Physics, 2005.
- [3] B3 Astrophysics Lecture Series, P.Podsiadlowski, Oxford Physics, 2006
- [4] Asian Physics Olympiad 2012, Theory Question 2, 2012

Appendix A Python Code

This appendix outlines the Python code used for this practical. The original code was compiled as an ipynb in Jupyter Notebook.

```
# Author: Guo-Zheng Theodore Yoong, Date: 8/2/2019
1
2
3
   import numpy as np
   import matplotlib.pyplot as plt
   from astropy import constants as const
6
7
   def ode_solve_rk(f, g, y0, t):
8
9
       Solves the coupled ODEs, i.e. dx/dt = f(x,y,t) and dy/dt = g(x,y,t),
          using Runge Kutta algorithm.
10
11
       Input:
12
       f, g: functions that receives the current state (x and y) and the
          current position/time (t), and returns the derivative values of the
          state, dx/dt and dy/dt.
       y0: the initial state of the system, given in a column matrix of
13
          dimensions 2 x 1 (in this case mass and density).
       t: vector of positions/time steps with length N where the values of y
14
          will be returned.
15
16
       Output:
       y: 2 x N matrix that contains the values of y at every position/time
17
          step. Columns correspond to position/time and rows to the element of
          у.
18
       N = len(t) # Number of iterations
19
20
       h = (t[-1]-t[0])/(N-1) # Step size
21
       x_{curr}, y_{curr} = y0[0][0], y0[1][0] # Initialise current x and y
22
       y = y0 # Creates a copy of y0
23
       for i in range(0, N-1): # Python is zero-indexed
24
           # This loop performs the Runge-Kutta algorithm
25
           k1 = f(x_{curr}, y_{curr}, t[i])
```

```
26
           11 = g(x_curr, y_curr, t[i])
27
           k2 = f(x_{curr}+0.5*h*k1, y_{curr}+0.5*h*k1, t[i]+0.5*h)
28
           12 = g(x_curr + 0.5*h*k1, y_curr + 0.5*h*k1, t[i] + 0.5*h)
29
           k3 = f(x_{curr}+0.5*h*k2, y_{curr}+0.5*h*k2, t[i]+0.5*h)
30
           13 = g(x_curr + 0.5*h*k2, y_curr + 0.5*h*k2, t[i] + 0.5*h)
31
           k4 = f(x_{curr}+h*k2, y_{curr}+h*k2, t[i]+h)
32
           14 = g(x_curr + h*k2, y_curr + h*k2, t[i] + h)
33
34
           # Update next values of x_curr and y_curr
35
           x_{curr}, y_{curr} = x_{curr} + h*(k1+2*k2+2*k3+k4)/6, y_{curr} + h*(11+2*12)
               +2*13+14)/6
36
           y[0].append(x_curr)
37
           y[1].append(y_curr)
38
       return y
39
40 def get_nonrel_density(rho0, radius):
41
42
       Obtains the density, rho, as function of the radial distance, r, using
           the implemented ODE solver using the non relativistic equation.
43
44
       Input:
       rho0: the central density at r = 0.
45
46
       radius: the grid points of radial distance where the density is
           calculated in form of vector with N elements. Variable name changed
          for consistency.
47
48
       Output:
49
       density: a vector with N elements that contains the density at radial
           distance given in r. Variable name changed for consistency.
50
       mass: a vector with N elements containing the cumulative mass of the
          white dwarf from r=0 to the given radial distance in r.
       ,, ,, ,,
51
52
       # Initialising constants, all in S.I. units
53
       h = const.h.value # Planck's constant
54
       m_e = const.m_e.value # Mass of Electron
55
       m_p = const.m_p.value # Mass of Proton
56
       G = const.G.value # Gravitational constant
57
58
       k = (48*(np.pi**(2/3))*G*m_e*(m_p**(5/3)))/((h**2)*(2**(1/3))*(3**(2/3))
          ) # For calculation of drho_dr
59
60
       r0 = 10 # An initial radius close to 0 with the central density, but not
            0 so that 1/r^2 is defined
61
       m0 = (4/3) * rho0 * np.pi * (r0 * * 3) # The corresponding initial mass
62
63
       drho_dr = lambda rho, m, r: -k*m*(rho**(1/3))/(r**2) # Lambda function
           for drho_dr
64
       dm_dr = lambda rho, m, r: 4*np.pi*rho*(r**2) # Lambda function for dm_dr
65
66
       init_rho_and_mass = [[rho0], [m0]]
67
       return ode_solve_rk(drho_dr, dm_dr, init_rho_and_mass, radius) # Here,
```

```
[[rho],[mass]] is y, and radius is t.
68
69
   def get_rel_density(rho0, radius):
70
71
        Obtains the density, rho, as function of the radial distance, r, using
           the implemented ODE solver using the relativistic equation.
72
73
        Input and output is the same as the non-relativistic function.
74
75
        # Initialising constants, all in S.I. units
76
        h = const.h.value # Planck's constant
77
        m_e = const.m_e.value # Mass of Electron
        m_p = const.m_p.value # Mass of Proton
78
79
        G = const.G.value # Gravitational constant
80
        c = const.c.value # Speed of light
81
82
        # For calculation of drho_dr
83
        k = ((2**(7/3))*(3**(1/3))*(np.pi**(1/3))*G*(m_p**(4/3)))/((h**2)*c)
84
        a = (2**(8/3))*(np.pi**(2/3))*(m_e**2)*(m_p**(2/3))*(c**2)
85
        b = (3**(2/3))*(h**2)
86
        r0 = 10 \# An initial radius close to 0 with the central density, but not
87
            0 so that 1/r^2 is defined
88
        m0 = (4/3)*rho0*np.pi*(r0**3) # The corresponding initial mass
89
90
        drho_dr = lambda rho, m, r: -k*m*(rho**(1/3))*np.sqrt(a+b*(rho**(2/3)))
           /(r**2) # Lambda function for drho_dr
91
        dm_dr = lambda rho, m, r: 4*np.pi*rho*(r**2) # Lambda function for dm_dr
92
93
        init_rho_and_mass = [[rho0], [m0]]
94
        return ode_solve_rk(drho_dr, dm_dr, init_rho_and_mass, radius) # Here,
           [[rho],[mass]] is y, and radius is t.
95
96 # As stated in the script, 5
97 # We choose an initial rho0 of 10^14
   radius = np.linspace(10, 50000000, 10000) # We can filter off NaN Values
98
       later
99
   rho0 = 10**14
100
101
   nonrel_rho_and_mass = get_nonrel_density(rho0, radius) # Error will be
       raised; ignore
102 nonrel_density = nonrel_rho_and_mass[0]
103 nonrel_mass = nonrel_rho_and_mass[1]
104
105 rel_rho_and_mass = get_rel_density(rho0, radius) # Error will be raised;
       ignore
106 rel_density = rel_rho_and_mass[0]
107 \text{ rel_mass} = \text{rel_rho_and_mass[1]}
108
109 # Scaling for plotting purposes
110 nonrel_radius_scaled = list(map(lambda x: x/(10**6), radius))
```

```
nonrel_density_scaled = list(map(lambda x: x/(10**14), nonrel_density))
112 nonrel_mass_scaled = list(map(lambda x: x/(10**32), nonrel_mass))
113
114 rel_radius_scaled = list(map(lambda x: x/(10**5), radius))
115 rel_density_scaled = list(map(lambda x: x/(10**14), rel_density))
116 rel_mass_scaled = list(map(lambda x: x/(10**30), rel_mass))
117
118 plt.plot(nonrel_radius_scaled, nonrel_density_scaled, c='red', label="
       Density")
119 plt.xlabel('Radius/1e6 m')
120 plt.ylabel('Density/1e14 kg m^-3')
121 plt.title('rho(r) (Non-relativistic)')
122 plt.show()
123
124 plt.plot(nonrel_radius_scaled, nonrel_mass_scaled, c='red', label="Mass")
125 plt.xlabel('Radius/1e6 m')
126 plt.ylabel('Mass/1e32 kg')
127 plt.title('m(r) (Non-relativistic)')
128 \, plt.show()
129
130 plt.plot(rel_radius_scaled, rel_density_scaled, c='blue', label="Density")
131 plt.xlabel('Radius/1e5 m')
132 plt.ylabel('Density/1e14 kg m^-3')
133 plt.title('rho(r) (Relativistic)')
134 plt.show()
135
136 plt.plot(rel_radius_scaled, rel_mass_scaled, c='blue', label="Mass")
137 plt.xlabel('Radius/1e5 m')
138 plt.ylabel('Mass/1e30 kg')
139 plt.title('m(r) (Relativistic)')
140 \, plt.show()
141
142 samples = np.logspace(6,14) # Default no. of samples is 50
143
144 nonrel_overall_radius = []
145 nonrel_total_mass = []
146 for sample in samples:
147
        nonrel_rho_and_mass = get_nonrel_density(sample, radius) # Error will be
            raised; ignore
148
        nonrel_mass = nonrel_rho_and_mass[1]
149
        nonrel_mass = [val for val in nonrel_mass if val == val]
        lastcount = len(nonrel_mass)
150
151
        nonrel_total_mass.append(nonrel_mass[-1])
152
        nonrel_overall_radius.append(radius[lastcount -1])
153
154 nonrel_overall_radius_scaled = list(map(lambda x: x/(10**7),
       nonrel_overall_radius))
    nonrel_total_mass_scaled = list(map(lambda x: x/(10**32), nonrel_total_mass)
155
       )
156
157 plt.plot(samples, nonrel_overall_radius_scaled, c='red', label="R")
```

```
158 plt.xscale('log')
159 plt.xlabel('rho_c/kg m^-3')
160 plt.ylabel('R/1e7 m')
161 plt.title('R (Non-relativistic)')
162 plt.show()
163
164 plt.plot(samples, nonrel_total_mass_scaled, c='red', label="M")
165 plt.xscale('log')
166 plt.xlabel('rho_c/kg m^-3')
167 plt.ylabel('M/1e32 kg')
168 plt.title('M (Non-relativistic)')
169 plt.show()
170
171 rel_overall_radius = []
172 \text{ rel\_total\_mass} = []
173 for sample in samples:
        rel_rho_and_mass = get_rel_density(sample, radius) # Error will be
174
           raised; ignore
175
        rel_mass = rel_rho_and_mass[1]
176
        rel_mass = [val for val in rel_mass if val == val]
177
        lastcount = len(rel_mass)
178
        rel_total_mass.append(rel_mass[-1])
179
        rel_overall_radius.append(radius[lastcount-1])
180
181
   rel_overall_radius_scaled = list(map(lambda x: x/(10**7), rel_overall_radius
182 rel_total_mass_scaled = list(map(lambda x: x/(10**30), rel_total_mass))
183
184 plt.plot(samples, rel_overall_radius_scaled, c='blue', label="R")
185 plt.xscale('log')
186 plt.xlabel(' _c/kg m^-3')
187 plt.ylabel('R/1e7 m')
188 plt.title('R (Relativistic)')
189 plt.show()
190
191 plt.plot(samples, rel_total_mass_scaled, c='blue', label="M")
192 plt.xscale('log')
193 plt.xlabel(' _c/kg m^-3')
194 plt.ylabel('M/1e30 kg')
195 plt.title('M (Relativistic)')
196 \text{ plt.show()}
197
198 plt.plot(samples, nonrel_overall_radius_scaled, c='red', label="Non-
       relativistic R")
199 plt.plot(samples, rel_overall_radius_scaled, c='blue', label="Relativistic R
       ", linestyle='dashed') # Dashed line for relativistic case
200 plt.xscale('log')
201 plt.xlabel(' _c/kg m^-3')
202 \text{ plt.ylabel('R/1e7 m')}
203 plt.title('Comparison of R')
204 plt.legend(loc='upper right')
```

```
205 plt.show()
206
207 plt.plot(samples, nonrel_total_mass, c='red', label="Non-relativistic M")
208 plt.plot(samples, rel_total_mass, c='blue', label="Relativistic M",
       linestyle='dashed') # Dashed line for relativistic case
209 plt.xscale('log')
210 plt.yscale('log')
211 plt.xlabel(' _c/kg m^-3')
212 plt.ylabel('M/kg')
213 plt.title('Comparison of M')
214 plt.legend(loc='upper left')
215 plt.show()
216
217\, # The script requires R as a function of M, so this is included. However, as
        M as a function of R is more intuitive, the report focuses more on log M
       (R) instead
218 plt.plot(nonrel_total_mass, nonrel_overall_radius_scaled, c='red', label="
       Non-relativistic case")
219 plt.plot(rel_total_mass, rel_overall_radius_scaled, c='blue', label="
       Relativistic case", linestyle='dashed')
220 plt.xscale('log')
221 plt.xlabel('M/kg')
222 plt.ylabel('R/1e7 m')
223 \text{ plt.title('R(M)')}
224 plt.legend(loc='upper right')
225 plt.show()
226
227 plt.plot(nonrel_overall_radius_scaled, nonrel_total_mass, c='red', label="
       Non-relativistic case")
   plt.plot(rel_overall_radius_scaled, rel_total_mass, c='blue', label="
228
       Relativistic case", linestyle='dashed')
229 plt.yscale('log')
230 plt.ylabel('M/kg')
231 \text{ plt.xlabel('R/1e7 m')}
232 \text{ plt.title('M(R)')}
233 plt.legend(loc='upper right')
234 \, plt.show()
235
236 \text{ M_C} = \text{rel\_total\_mass[-1]}
237 print(M_C)
```