CO31: Structure of white dwarf stars

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Abstract

This will be the abstract of the report. It shall be written when the rest is nearly done. The goal of this experiment is to study the internal structure of white dwarf stars. They are one possible end state in the life cycle of stars. What is particular about white dwarf stars is that the matter they are composed of is different from the matter in a usual star similar to the Sun. While in a usual stable star the equilibrium is supported by fusion energy and the pressure of the plasma, a white dwarf star has exhausted its fuel, and the equilibrium is supported by electron degeneracy pressure, making the star extremely dense; for example, a white dwarf of 1 solar mass is about the size of Earth. In this report, we will first derive the coupled differential equations for mass and density in a spirit similar to Oxford Physics (2020) and Chandrasekhar (1984) in Section 2. Then, in Section 3, we develop the numerical method used to solve the coupled equations. Results and discussion take place in Section 4. Finally, we give a summary of the experiment and possible improvements in Section 5.

1 Introduction

The goal of this experiment is to study the internal structure of white dwarf stars. They are one possible end state in the life cycle of stars. What is particular about white dwarf stars is that the matter they are composed of is different from the matter in a usual star similar to the Sun. While in a usual stable star the equilibrium is largely supported by fusion energy and the pressure of the plasma, a white dwarf star has reached the end of the fusion processes, and the equilibrium is supported by electron degeneracy pressure, making the star extremely dense; for example, a white dwarf of 1 solar mass is about the size of Earth. In this report, we will first derive the coupled differential equations for mass and density in a spirit similar to Oxford Physics (2020) and Chandrasekhar (1984) in Section 2. Then, in Section 3, we develop the numerical method used to solve the coupled equations. Results and discussion take place in Section 4. Finally, we give a summary of the experiment and possible improvements in Section 5.

2 Theory of white dwarf stars

Stars are astronomical objects with ellipsoid shape made up of heated plasma. The force of gravity tries to compress the ball, holding the matter together, while the gaseous plasma opposes gravity with the pressure it exerts. When the two forces balance each other out, the star is in hydrostatic equilibrium. In usual stars, pressure is due to the gas and the radiation (which becomes important mostly in the largest non-degenerate stars). In white dwarf stars, however, the densities are far greater than the ones found in usual stars, and matter behaves differently. All electrons are no longer bound to atoms and are free to roam. The behaviour can approximately be modelled by a Fermi gas of electrons at zero Kelvin (i.e. fully degenerate state). Hence, the pressure ensuring the equilibrium is the degeneracy pressure of the electrons.

2.1 Equation of equilibrium

Assume the star is spherically symmetric, in equilibrium, non-rotating, and the effect of magnetic fields is negligible. Therefore, all properties depend only on the distance to the centre of the star, r. The gravitational force on a small volume of matter with area $\mathrm{d}A$ and radial height $\mathrm{d}r$ is:

$$F_{\rm G} = -\frac{Gm(r)\,\mathrm{d}m}{r^2},\tag{1}$$

where m(r) is the mass of the star contained up to r, $dm = dA dr \rho(r)$ is the mass of the small volume, and the density $\rho(r)$ was assumed constant (negative sign because it pulls towards the centre). The force due to the pressure is the difference of the forces at r and r + dr:

$$F_{\mathbf{P}} = (P(r) - P(r + \mathrm{d}r)) \,\mathrm{d}A. \tag{2}$$

For a star in equilibrium the two forces balance each other out, i.e. $F_{\rm G} + F_{\rm P} = 0$. After rearranging, we get:

$$\frac{\mathrm{d}P(r)}{\mathrm{d}r} = \frac{P\left(r + \mathrm{d}r\right) - P\left(r\right)}{\mathrm{d}r} = -\frac{G\rho(r)m(r)}{r^2}.$$
(3)

We can rewrite the hydrostatic equilibrium by using the chain rule:

$$\frac{\mathrm{d}P(r)}{\mathrm{d}r} = \frac{\mathrm{d}P}{\mathrm{d}\rho} \frac{\mathrm{d}\rho}{\mathrm{d}r},\tag{4}$$

so equation (3) becomes:

$$\frac{\mathrm{d}\rho}{\mathrm{d}r} = -\frac{\mathrm{d}\rho}{\mathrm{d}P} \frac{G\rho(r)m(r)}{r^2}.$$
 (5)

On the other hand, the equation for the inscribed mass is:

$$m(r) = \int_0^r dr' 4\pi^2 r' \rho(r') \quad \Rightarrow \quad \frac{dm}{dr} = 4\pi r^2 \rho(r) \tag{6}$$

We are now left with two coupled differential equations, (5) and (6). The last task before obtaining the full equation is to determine $\frac{dP}{d\rho}$, which depends on the equation of state.

2.2 Equation of state

We assume the star is made up primarily of heavy 56 Fe nuclei and their electrons. The nuclei carry most of the mass and little of the pressure, while the opposite is true for the electrons. Since *very high* densities are considered, we can approximate that the nuclei are stationary while the electrons move freely, not bound to any nuclei. A good model for the freely moving electron gas is the free Fermi gas at T = 0 K. This is the fully degenerate state, in which electrons fill all energy (momentum) levels up to the Fermi energy (momentum).

Fermions obey the Pauli exclusion principle, therefore each energy level is 2S+1 degenerate, where $S=\frac{1}{2}$ in the case of electrons. This degeneracy factor multiplies gives the density function:

$$g(k) = \frac{2S+1}{2\pi^2} V k^2, \tag{7}$$

where k is the wave vector of particles and V is the volume of the system. The number density of particles can therefore be written as:

$$n = \frac{N}{V} = \frac{1}{V} \int_0^{k_{\rm F}} dk \, g(k) = \int_0^{k_{\rm F}} dk \frac{2S+1}{2\pi^2} k^2.$$
 (8)

Here we integrate up to $k_{\rm F}$ where this is given by $p_{\rm F} = \hbar k_{\rm F}$ and $p_{\rm F}$ is the Fermi momentum. Rewrite the above equation in terms of p:

$$n = \int_0^{p_F} dp \frac{2S+1}{2\pi^2 \hbar^3} p^2 = \int_0^{p_F} dp \frac{8\pi}{h^3} p^2 = \int_0^{p_F} dp \, n(p) = \frac{m_e 8\pi}{h^3} p_F^3$$
 (9)

where we define n(p) dp as the number of electrons per unit volume with momentum between p and p + dp. Pressure, on the other hand, is given by the kinetic expression:

$$P = \frac{1}{3} \int_{0}^{p_{\rm F}} p v_p n(p) \, \mathrm{d}p. \tag{10}$$

At this point we need to differentiate between non-relativistic and relativistic cases - v_p has a different value in the two cases:

$$v_p = \begin{cases} \frac{p}{m_e} & \text{in the non-relativistic case} \\ \frac{pc^2}{\sqrt{p^2c^2 + m_e^2c^4}} & \text{in the relativistic case} \end{cases}$$
 (11)

Let us first work out the answer in the non-relativistic case. The integral becomes const $\times \int p^4 dp$, so the pressure becomes:

$$P_{\rm non} = \frac{8\pi}{15h^3 m_{\rm e}} p_{\rm F}^5. \tag{12}$$

For the relativistic case, rather than do the integral for P involving hyperbolic functions, differentiate (10) by parts:

$$\frac{\mathrm{d}P_{\rm rel}}{\mathrm{d}\rho} = \frac{\mathrm{d}P_{\rm rel}}{\mathrm{d}p_{\rm F}} \frac{\mathrm{d}p_{\rm F}}{\mathrm{d}\rho}, \quad \frac{\mathrm{d}P_{\rm rel}}{\mathrm{d}p_{\rm F}} = \frac{8\pi}{3h^3} \frac{\mathrm{d}}{\mathrm{d}p_{\rm F}} \int_0^{p_{\rm F}} \frac{p^4c^2}{\sqrt{p^2c^2 + m_{\rm e}^2c^4}} \,\mathrm{d}p = \frac{8\pi}{3h^3} \frac{p_{\rm F}^4c}{\sqrt{p_{\rm F}^2c^2 + m_{\rm e}^2c^4}}.$$
 (13)

We can collect the derivative of (12) and the result of (13):

$$\frac{\mathrm{d}P}{\mathrm{d}\rho} = \frac{\mathrm{d}p_{\mathrm{F}}}{\mathrm{d}\rho} \frac{8\pi}{3h^{3}m_{\mathrm{e}}} p_{\mathrm{F}}^{4} \times \begin{cases} 1 & \text{in the non-relativistic case} \\ \left(1 + \frac{p_{\mathrm{F}}^{2}}{m_{\mathrm{e}}^{2}c^{2}}\right)^{-1/2} & \text{in the relativistic case} \end{cases}$$
(14)

What is left is expressing $p_{\rm F}(\rho)$ to complete the equation of state in the form needed by (5). The relation between density and number density can be stated as $\rho = \mu_{\rm e} m_{\rm p} n$, where $\mu_{\rm e}$ is the mean molecular weight per electron and $m_{\rm p}$ is the mass of 1 proton. In our case, we can approximate $\mu_{\rm e} \approx 2$. Using this relation and (9), we arrive at:

$$p_{\rm F} = \left(\frac{h^3}{8\pi}n\right)^{1/3} = \left(\frac{h^3}{16\pi m_{\rm p}}\rho\right)^{1/3} \quad \Rightarrow \quad \frac{\mathrm{d}p_{\rm F}}{\mathrm{d}\rho} = \frac{1}{3}\left(\frac{h^3}{16\pi m_{\rm p}}\right)^{1/3}\rho^{-2/3} \tag{15}$$

2.3 The limiting mass

The limiting case of ultra-relativistic electrons can be obtained by setting $v_p = c$ (i.e. taking the limit $pc \gg m_e c^2$) in (10) to arrive at the equation of state:

$$P_{\text{ultra}} = \frac{p\pi}{3h^3} \int_0^{p_F} p^3 c \, dp = \frac{2\pi c}{3h^3} p_F^4.$$
 (16)

As noted in Chandrasekhar (1984), this leads to a specific equation for the mass with a well specified value $M_{\rm lim} = 5.76 \mu_{\rm e}^{-2} M_{\odot}$, where M_{\odot} is one solar mass; this limit is equivalent to infinite mean density and zero radius of the star. In the approximation made above the limiting mass is $M_{\rm lim} \approx 1.435 M_{\odot}$. This will be "experimentally" verified by the computations presented in Section 4. We can draw two important conclusions from this fact: first, there is an upper limit to masses of degenerate stars in the later stages of evolution; and second, we cannot predict the end evolutionary state of stars with mass $M > M_{\rm lim}$ in this physical framework (we expand more on this in Section 4).

3 Numerical approach

To find the radius and mass of the star as a function of its central density, we will numerically integrate the coupled system. The distance r at which $\rho(r=R)\approx 0$ is where the star ends; the mass is therefore m(R) where m(r) is the inscribed mass, as defined in (6). Physically, we can consider solving the equation as an initial value problem. First, write the coupled system in vector form:

$$\mathbf{y} = \begin{pmatrix} \rho \\ m \end{pmatrix},\tag{17}$$

and the derivatives of its members with respect ro r given by (5) and (6). Then, for a given central density ρ_c the initial condition for the mass is specified by $m(\delta r) = \frac{4}{3}\pi(\delta r)^3\rho_c$ for some small distance from the centre δr : this is done so we can start the calculations (r=0 would give initial zero mass and the derivative is therefore zero). We will set it up the equation solver so that this δr is the step size in radial distance. The source code can be found in Appendix A.

3.1 Why integration?

Since we have an initial value problem, the underlying idea is the following: rewrite the derivatives as fractions of finite differences, and get $\Delta y = \frac{\mathrm{d} y(x)}{\mathrm{d} x} \Delta x$. This is the change in y when we step through by Δx . When the step is very small, the approximation is very good, i.e. $\lim_{\Delta x \to \mathrm{d} x} \Delta y = \mathrm{d} y$.

The basic idea boils down to the most elementary such method — Euler's method:

$$\mathbf{y}_{n+1} = \mathbf{y}_n + h\mathbf{f}(x_n, \mathbf{y}_n)$$

$$x_{n+1} = x_n + h,$$
(18)

where the vector function is just $\mathbf{f} = \frac{\mathrm{d}\mathbf{y}(x)}{\mathrm{d}x}$ i.e. the slope at x_{n} and $h = \Delta x$. This is a first order method: the local error is $\sim h^2$ and the global error therefore is $\sim h$. This method has only one computation of the derivative and is therefore very fast, but also very inaccurate.

A similar explicit second order method is Heun's method (trapezoid rule):

$$\mathbf{y}_{n+1} = \mathbf{y}_n + \frac{1}{2}h\Big(\mathbf{f}(x_n, \mathbf{y}_n) + \mathbf{f}(x_n + h, \mathbf{y}_n + \mathbf{f}(x_n, \mathbf{y}_n))\Big)$$

$$x_{n+1} = x_n + h,$$
(19)

Here the local error is $\sim h^3$ and the global error therefore is $\sim h^2$.

The method we will later use to compute properties of white dwarf stars is the 4th order Runge-Kutta (RK4) method. It features four evaluations of the derivative and a global error $\sim h^4$, offering a balance between computational time and accuracy. This method takes values for the slope at four different positions and estimates the step size by weighted average of these four steps. In equations:

$$\begin{aligned} & \boldsymbol{k_1} = h\boldsymbol{f}(x_{\mathrm{n}}, \boldsymbol{y}_{\mathrm{n}}) \\ & \boldsymbol{k_2} = h\boldsymbol{f}\left(x_{\mathrm{n}} + \frac{1}{2}h, \boldsymbol{y}_{\mathrm{n}} + \frac{1}{2}\boldsymbol{k_1}\right) \\ & \boldsymbol{k_3} = h\boldsymbol{f}\left(x_{\mathrm{n}} + \frac{1}{2}h, \boldsymbol{y}_{\mathrm{n}} + \frac{1}{2}\boldsymbol{k_2}\right) \\ & \boldsymbol{k_4} = h\boldsymbol{f}(x_{\mathrm{n}} + h, \boldsymbol{y}_{\mathrm{n}} + \boldsymbol{k_3}). \end{aligned}$$

Finally, the new position is calculated as:

$$y_{n+1} = y_n + \frac{1}{6}k_1 + \frac{1}{3}k_2 + \frac{1}{3}k_3 + \frac{1}{6}k_4.$$
 (20)

The RK methods family gives approximate solutions to Nth order by varying the weights in the k expressions and the final expression to match coefficients in the Taylor expansion.

We should note that the equation to be solved might prove difficult to solve, even using RK4. These equations are often called *stiff* and in constructing their solutions more precaution needs to be taken, for example by using an implicit or semi-implicit methods.

3.2 Comparison of methods

In this section we compare the stability/accuracy of three methods presented in Section 3.1. For the task, we will try to integrate a solution of the simple harmonic oscillator (SHO), namely:

$$\ddot{x} + \omega^2 x = 0$$
, with solution $x = x_0 \cos(\omega t)$ (21)

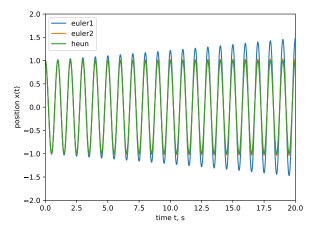
where we choose $x_0 = 1$ and $\omega = 2\pi$ so the period of oscillations is 1 s. From some initial runs integrating the white dwarf equation we are aware that the number of intervals relevant us is on the order of 10^6 so we choose to compare the two methods on this scale (more discussion on the topic in Section 3.4).

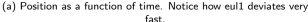
The runs will be split into two groups. The first group is with 10⁶ time intervals of length 1 ms for the three methods; denote the results from these runs by EULER1, HEUN, and RK4 respectively. The second group is 10⁷ time intervals of length 0.1 ms for the Euler method to cover the same total time interval; denote results by EULER2. Additionally, we will consider the energy in the system:

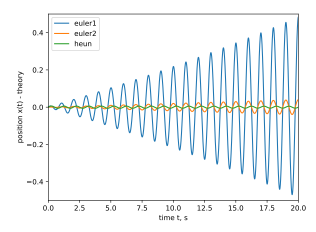
$$E = \frac{\dot{x}^2}{2} + \frac{\omega^2 x^2}{2}. (22)$$

When the method over/underestimates the correction, the effect will build up and change the total energy, which is equivalent to the change in amplitude.

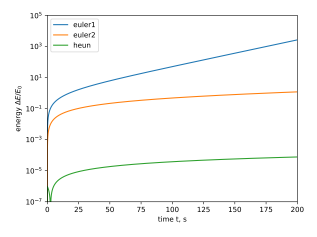
Results are presented in Figure 1. In 1a (first 20 periods of oscillation displayed), we see the calculated position by the three methods EULER1, EULER2, and HEUN. It is immediately obvious that EULER1 goes off in the solution pretty quickly. This is confirmed in 1b, where we can have a better look at how the numerical solution deviates from the actual one - EULER1 gains an additional half amplitude in 20 periods, while the other two methods oscillate about zero with HEUN having the smaller error. To compare for the whole time of integration and compare these three runs with RK4, plot the relative energy deviation compared to the theoretical value of $2E_0 = \omega^2 \times 1$ on a



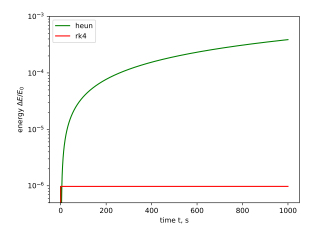




(b) Deviation from the theoretical solution.



(c) Energy deviation $\Delta E/E_0$ for first three methods.



(d) Energy deviation $\Delta E/E_0$ for heun and rk4.

Figure 1: Test runs of SHO integrated with different methods. Blue line is for Euler's method integrated with step $1\,\mathrm{ms}$, yellow line for Euler's method integrated with step $0.1\,\mathrm{ms}$, green line is for Heun's method with step $1\,\mathrm{ms}$, and red line is for RK4 method with step $1\,\mathrm{ms}$.

logarithmic scale. As it can be seen in 1c and 1d, the energy for the first three runs grows exponentially with time after some moment with different exponents, while the fourth run RK4 keeps almost a constant value of energy — the change in energy between $t_0 = 0$ s and $t_1 = 100$ s is:

$$\frac{\left|E(t_1) - E(t_0)\right|}{E(t_0)} \approx 9.755 \times 10^{-7}.$$
 (23)

With these results in mind, we make the obvious choice for a method and continue forwards using the Runge-Kutta 4th order method in all of the following chapters.

3.3 Dimensionless equation for the white dwarf

To calculate values with the computer, we will go to dimensionless quantities in the differential equation. Introduce the following substitutions:

$$\rho = \rho_{\rm c}\theta, \quad r = \ell\xi, \quad m = \rho_{\rm c}(1\,{\rm m}^3)\mu, \text{ therefore } \boldsymbol{y}(r) = \begin{pmatrix} \rho(r) \\ m(r) \end{pmatrix} \rightarrow \boldsymbol{\eta}(\xi) = \begin{pmatrix} \theta(\xi) \\ \mu(\xi) \end{pmatrix}.$$
(24)

We also specify the length ℓ by the equation $\frac{4}{3}\pi\ell^3=1\,\mathrm{m}^3$, which gives a value of $\ell\approx0.62\sim1\,\mathrm{m}$. This will be useful to make some cancellations in the coefficients in the final form of the differential equation for η . The initial

conditions therefore become $\eta(1) = (\theta(1), \mu(1\ell)) = (1.0, 1.0)$.

For (6) the conversion is straightforward:

$$\frac{\rho_{\rm c} \,\mathrm{d}\mu}{\ell \,\mathrm{d}\xi} = 4\pi \ell^2 \xi^2 \rho_{\rm c}\theta \quad \Rightarrow \quad \frac{\mathrm{d}\mu}{\mathrm{d}\xi} = 3\xi^2 \theta. \tag{25}$$

For the θ equation we have to take into consideration the regime we operate in — whether we solve for a relativistic or non-relativistic equation of state. Firstly, write equations in (15) using the new coordinates we have introduced:

$$p_{\rm F} = \left(\frac{h^3 \rho_{\rm c}}{16\pi m_{\rm p}}\right)^{1/3} \theta^{1/3}, \quad \frac{\mathrm{d}p_{\rm F}}{\mathrm{d}\rho} = \frac{1}{3} \left(\frac{h^3}{16\pi m_{\rm p} \rho_{\rm c}^2}\right)^{1/3} \theta^{-2/3}. \tag{26}$$

Now substitute (14) in (5) using (26). After some algebra with the constants, the final result is the following:

$$\frac{\mathrm{d}\theta}{\mathrm{d}\xi} = K_1 \frac{\mu \theta^{1/3}}{\xi^2} \begin{cases} 1 & \text{in the non-relativistic case} \\ \left(1 + K_2 \theta^{2/3}\right)^{1/2} & \text{in the relativistic case} \end{cases}, \tag{27}$$

where the constants K_1 and K_2 are given by

$$K_1 = -2^{13/3} \pi G m_e m_p^{5/3} \rho_c^{1/3} h^{-2}, \qquad K_2 = \frac{1}{m_e^2 c^2} \left(\frac{3h^3 \rho_c}{16\pi m_p} \right)^{2/3}.$$
 (28)

We have completed the task of bringing the equation to normalized coordinates. This is the setup used in implementing the code for the white dwarf ODE wrapper presented in Appendix A.1.

3.4 Convergence and number of intervals

What is left is to choose the coarseness in the grid of ξ . We know that for some $\Delta \xi$ small enough, the answers will converge to \approx a single value. To find the number of intervals (NoI) at which the result converges, we make experimental runs at three different $\rho_c - 10^6$, 10^{10} , and $10^{14} \text{ kg m}^{-3}$ (this spans the range for which we run the experiment later), and test for a NoI in the range $\{a \times 10^b \mid a \in \{1, 2, 4, 8\}, b \in \{2, 3, \dots 8\}\}$. The maximal radius is set to 8.1×10^7 in units of ξ , since this turns out to be about the maximum physical radius a white dwarf can be for the range of ρ_c we test. Both the non-relativistic and relativistic cases are considered.

The results are presented in Figure 2. We use the relative error in computed value with respect to the converged value. The latter is estimated as the average of the last three values in the calculation, i.e. for NoI 2×10^8 , 4×10^8 , and 8×10^8 ; the result is M_{conv} , R_{conv} . The relative error is given by:

$$\frac{\Delta M}{M_{\rm conv}} = \frac{|M - M_{\rm conv}|}{M_{\rm conv}}, \quad \frac{\Delta R}{R_{\rm conv}} = \frac{|R - R_{\rm conv}|}{R_{\rm conv}}.$$
 (29)

What we aim for in our choice for the number of intervals is: on the one side, a low enough number so the computation does not take too long, and on the other side, a high enough number so an acceptable accuracy is achieved. From the logarithmic plots for the mass and the radius determination, we see that the computed value does not really converge to a specific point; it does, however, reduce its error exponentially fast. Let us choose a desired accuracy of below $10^{-5} = 0.001\%$, given that our model is not very accurate. From the logarithmic plots again we see that this is satisfied for NoI $\geq 2 \times 10^6$. Therefore, we can use this value as the NoI with a good compromise between speed and accuracy.

4 Results and discussion

We begin by choosing the parameters of the simulation. As discussed in Section 3.4, we use the RK4 method of integration. The range of integration in ξ will be $[1, 8.1 \times 10^7]$ with 2×10^6 intervals (we implement the white dwarf wrapper in this way to speed up the procedure, see Appendix A.1).

For the central density ρ_c we are tasked to study the range $10^6 < \rho_c < 5 \times 10^{14} \,\mathrm{kg} \,\mathrm{m}^{-3}$, so we set up to test using the following values: $\rho_c \in \{a \times 10^b \mid a = 1, 2, \dots, 9; b = 6, 7, \dots, 14\}$. The results are presented in Figures 3, 4, and 5. We have chosen logarithmic plots for the first two plots to better highlight the similar behaviour at low densities and the different behaviour at high densities.

In Figure 3 and 4 we can see that at central densities $\rho_c \approx 10^8$ the results for radius and mass start to deviate. The non-relativistic values follow an exponential relationship, as one might have predicted from (14) and (5). The relativistic case, on the other hand, exhibits a more interesting behaviour. At higher densities, the mass tends to a single maximum value — the *Chandrasekhar mass*, which can be seen in Figures 4 and 5. This implies that the theoretical model of a white dwarf cannot properly describe objects with higher mass.

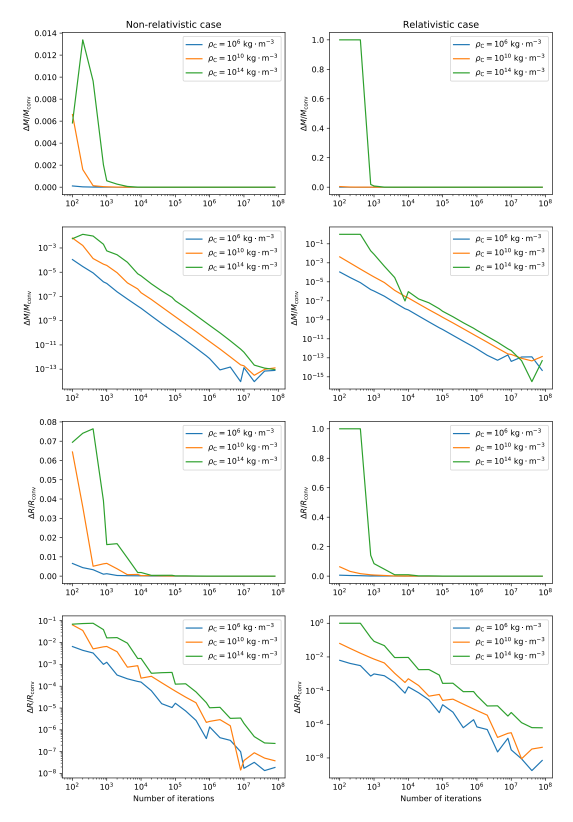


Figure 2: Plot of convergence for three central spanning the tested range of ρ_c in both non-relativistic (left column) and relativistic (right column) cases. As can be seen, convergence is slower in the relativistic case; after 10^6 intervals good accuracy is achieved in both cases for both mass and radius values. The values $M_{\rm conv}, R_{\rm conv}$ are computed as the average of the last three values; $\Delta M = |M - M_{\rm conv}|$ and $\Delta R = |R - R_{\rm conv}|$. Odd rows have a linear y-axis, even rows are the same plot with a logarithmic y-axis.

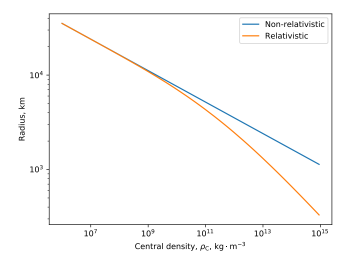


Figure 3: Radius of a white dwarf R in km as a function of central density ρ_c .

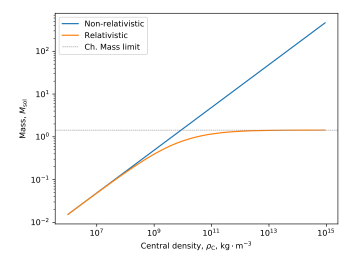


Figure 4: Mass of a white dwarf star M in solar masses as a function of central density ρ_c . Note the Chandrasekhar mass limit.

4.1 Compact objects

White dwarfs are part of the more general framework of compact objects as end stages in stellar evolution, which also include neutron stars and black holes. These objects are the final result in stellar evolution. In which one a star settles is almost completely determined by its initial mass during the stable stage on the Main sequence.

Low to medium mass stars $(0.6M_{\odot} \lesssim M \lesssim 8M_{\odot})$ usually evolve by hydrogen burning joined by helium burning in the red giant stage. Since their carbon cores are not massive enough to ignite, the stars contract again and the strong stellar wind expels all surrounding gas to form a planetary nebula with a hot central star, which in turn cools down to a white dwarf. These white dwarfs are usually carbon and oxygen-rich.

High mass stars $(M \gtrsim 8 M_{\odot})$ have massive enough cores to ignite further production of heavier elements via the alpha and triple alpha processes. After most of the fuel is burned through and a dense iron core has formed, they undergo a similar process of implosion in which the core rapidly compactifies (in some edge cases about $8 M_{\odot}$ only partial fusion of carbon occurs, they settle to a white dwarf state). In this case, however, the core mass has passed the (effective) Chandrasekhar limit and electron degeneracy pressure cannot support the equilibrium. Thus, the core collapses further to a neutron star or a black hole (the exact mass threshold for neutron stars is estimated to be between $2 M_{\odot}$ and $3 M_{\odot}$).

It is the discovery of this limit in Chandrasekhar (1931), however, that started the whole further theoretical study

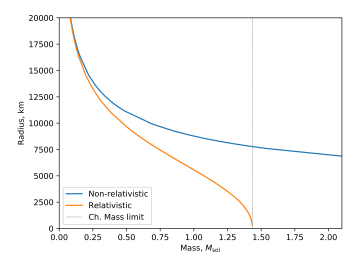


Figure 5: Radius of a white dwarf R in km as a function of the mass M in solar masses. Note the Chandrasekhar mass limit.

of the topic of compact objects. In the following year the neutron was discovered, and Landau published a paper which discussed a more precise calculation of the white dwarf mass limit and pondered the nature of heavier stars. This laid the groundwork for further discoveries in the 20th century.

4.2 Binary systems

Additionally, white dwarfs are often part of binary or multiple systems (about a third of all star systems are multiple systems). In the case the white dwarf is part of a very close binary, there is an accretion of mass to the surface of the white dwarf to a thin dense atmosphere layer atop the surface of the star. This atmosphere, usually comprised of hydrogen, is heated up by the star, which causes the ignition of runaway fusion. This is observed as a sudden flash of light - a *nova*.

If there is accretion from a more massive companion to a carbon-oxygen white dwarf (heavier elements support further collapse since more energy is needed for their ignition), this can cause its mass to increase beyond the Chandrasekhar limit and ignite carbon fusion in the centre of the white dwarf due to compressional heating. This results in the destruction of the star and is believed to be the physical model behind Type Ia *supernovae*.

This further highlights the critical mass limit of existence for white dwarfs. Without it, the two phenomena above would not be observed in nature.

5 Summary

this is a citation Sagert et al. (2005)

In this practical, we study the structure of white dwarf stars by numerically solving the equations of equilibrium. For a white dwarf star, the equilibrium is supported by the electron degeneracy pressure. We derive these equations in Section 2, differentiating between the cases of a non-relativistic and relativistic electrons. Additionally, we consider the ultrarelativistic limit and anticipate the result of calculations — a limit to white dwarf mass in the case of relativistic electrons.

We choose to solve these equations as an initial value problem, where the starting point is a small region at the centre of the star. In Section 3, we compare between the Euler, Heun, and Runge-Kutta 4th order (RK4) methods, which resulted in our choice to use the RK4 for solving the white dwarf equations. Then we introduce a transformation of variables to dimensionless quantities, after which we test the convergence of white dwarf mass and radius for a sample of central density values. We conclude that when the tested interval for radial distance in dimensionless units is $\xi \in [1, 8.1 \times 10^7]$ with 2×10^6 intervals we have a good compromise between accuracy and speed (this corresponds to physical dimensions of approximately $r \in [0.6 \text{ m}, 5 \times 10^7 \text{ m}]$ and a step size of $\Delta r = 25 \text{ m}$).

We present the solution to the white dwarf equation in Section 4. We note the difference of solutions between non-relativistic and relativistic equation of state description as the central density grows, seen in Figures 3 and 4. There we also discuss the implications of the existence of the *Chandrasekhar limit* and the place of white dwarfs in

the evolutionary track of stars as an end stage of medium mass stars $M \lesssim 8 M_{\odot}$, compared to other compact objects (neutron stars and black holes). Finally, we note some other astrophysical phenomena which occur due to the special properties of white dwarfs — novae and supernovae Type Ia.

While this project takes steps towards a numerical solution of a white dwarf equations, we have made some major assumptions along the way with regards to the physics. Among the effects to take into account are the following:

- The interaction of electrons with their surroundings: we can introduce magnetic fields and the electrostatic interaction in the force balance equations, which modifies the white dwarf equation.
- Chemical composition: since stars form an onion-like structure of elements in their cores, the mean molecular weight per electron μ_e varies slightly, which changes the constant factors in the equation for the white dwarf.
- Effects of General Relativity should also be taken into account. White dwarfs are compact objects, so as the density grows larger, the effects are more and more significant.
- We assumed that the star is in a complete stationary equilibrium state. However, we also have to consider the cases where the star might be rotating (a compact object can have a significant rotation speed due to conservation of angular momentum) or pulsating, which can disturb the conditions in the centre.

An introductory account of some of these effects can be found in e.g. Sagert et al. (2005), while a more standard and in depth textbook exposition to the topic can be found in Shapiro and Teukolsky (1991).

Additionally, there is room for improvement in the computational methods employed as well. The fairly simple RK4 method can be modified to have a varying step size that self-corrects depending on the derivative at a given point. That way the calculations can skip over a region with small a small gradient by increasing the step size, and decrease the step size to get a more accurate value for the change in a region with a big gradient. Another direction of improvement would be to change from RK4 to another, more sophisticated method that has better accuracy and faster computational time like the Bulirsch-Stoer method. This would be especially helpful if the system of differential equations to be solved was much more complex in its behaviour. A good introductory text on the subject is Press et al. (2007), Chapter 17.

References

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Press, W. H., Teukolsky, S. A., Vetterling, W. T. and Flannery, B. P. (2007), Numerical recipes: the art of scientific computing, third edition. edn, Cambridge University Press, Cambridge.

Sagert, I., Hempel, M., Greiner, C. and Schaffner-Bielich, J. (2005), 'Compact stars for undergraduates', Eur. J. Phys. 27:577-610, 2006.

Shapiro, S. L. and Teukolsky, S. A. (1991), Black Holes, White Dwarfs and Neutron: the physics of compact objects, VCH PUBN.

A Source code

The structure of the code has been guided by the guidance set in Oxford Physics (2020). Additionally, Press et al. (2007) has had some influence on general code structure. The documentation for the main ODE solver is in the Python code. A copy in an online repository can be found here.

A.1 Main ODE facilities

```
Implementation of a generic ODE integrator.
2
3
    import numpy as np
5
    from . import integrators
    class ODEinit(object):
8
9
        Base class for a general ODE to be numerically integrated. It provides
10
        facilities to compute a value of some linear system given the ODE form as
11
         y' = f(x, y) for some interval of values of x and initial conditions of
12
        the vector `y`.
13
14
        Parameters
15
        _____
16
        y0 : np.ndarray
17
            Initial conditions of the vector \dot{y} at position \dot{x} = \text{span}[0]
18
        deriv
            The function `f` in a functional form. Should accept two parameters at
20
            which to compute the derivative:
                x: float, the value of the independent variable;
22
                 y: np.ndarray, the value of the dependent variable vector `y`.
23
        span : np.ndarray
24
            The interval values over which to compute the value of the vector `y`.
25
26
        Attributes
         _____
28
        y0 : np.ndarray
29
            The initial state of the system. Passed at initialization.
30
        deriv
31
            The function `f` in a functional form. Passed at initialization.
32
        span : np.ndarray
33
            The interval of values to be integrated over. Passed at initialization.
            **Assumed to be equidistant!!**
35
        interval : float
36
            The size of the interval step; equal to `span[1] - span[0]`.
37
        yOut : np.ndarray
            The computed values of `y` after integration. Initialized as a matrix
39
            of zeros; if integration terminates before the end of `span`, the rest
40
            of the points are left as zeros.
        flagIntegrated : bool
            A flag to indicate whether the integration has been done (and `yOut`
43
            filled) to be used by methods depending on having computed values.
45
        Methods
46
        _____
47
        integrate
48
            Integrates the differential equation.
```

```
0.00
50
         def __init__(self, y0: np.ndarray, deriv, span: np.ndarray) -> None:
52
             self.y0 = y0
53
             self.deriv = deriv
             self.interval = span[1] - span[0]
55
             self.span = span
56
57
             # size of this is number of values to be computed x dimensions of y
             self.yOut = np.zeros((self.span.size, self.yO.size))
59
60
             # a flag taking note if the equation has been integrated
61
             self.flagIntegrated = False
63
         def integrate(self, integrator: int=3) -> None:
65
             Integrates the equation using the function of the derivative `deriv`.
66
             User has the option to choose an integrator from the library, the
67
             default is a RK4 integrator. As it computes, it fills the `yOut` with
             the computed values. Can terminate prematurely if the value passed to
69
             `deriv` is inappropriate; then the rest of the values are left as zero.
70
71
             Parameters
72
             _____
             integrator: {1, 2, 3}, optional
74
                 Choice of the integrator to be used given by the number:
75
                     1: the Euler first order method;
76
                     2: the Heun method, a second order method;
                     3: the Runge-Kutta 4th order method.
78
             See Also
80
             modules.integrators
82
83
             # choose an integrator
84
             self.integrator = integrators.index[integrator]
86
             # fill in the initial value of y
             self.yOut[0,:] = self.yO[:]
89
             #cycle does the integration
90
             for i in range(self.span.size-1):
91
                 # the try except handles the case where integration is impossible,
                 # i.e. when the value of an element of y is nonphysical. deriv
93
                 # decides when this is the case.
94
                 try:
95
                     # calculate the new value
                     adder = self.integrator(
97
                          self.span[i], self.yOut[i,:], self.deriv, self.interval)
                     # append new value
99
                     self.yOut[i+1,:] = adder
100
                 except ValueError:
101
                     # ends the integration if it is impossible to integrate - Value
102
                     # Error concerns the y-values
103
104
                     break
105
             # to assert that the ODE has been integrated in functions that use yOut.
106
```

```
self.flagIntegrated = True
107
108
     if __name__=="__main__":
109
         print(
110
             "This file contains declaration of a generic ODE class.")
111
     11 11 11
 1
     White dwarf class wrapper.
 2
 3
     import numpy as np
     from scipy import constants as cs # constants for the derivative equation
 5
     from . import derivatives, ode
 8
     class whiteDwarf(ode.ODEinit):
 9
10
         Provides a wrapper of the ODEinit class in the case of white dwarfs with
11
         some equation of state. Initializes an ODE with appropriate initial
12
         conditions while taking into account a normalization which leaves the ODE
13
         working with dimensionless quantities.
14
         Parameters
16
17
         rhoC : float
18
             The density of matter in the small region around the center. In units
19
             of kg * m^{-3}
20
         span : np.ndarray
             The range of values of r to be calculated at. In normalized units to
22
             cut down on computational time and improve calculation accuracy.
23
             Expects an array with first member = 1 for the starting central mass to
24
             be correctly defined. Unit of normalization given by 1 (see below).
25
         regime : {1, 2, 3}, optional
26
             The regime indicates the choice of equation of state -> derivative, by
27
             default 1 - non-relativistic.
28
29
         Attributes
         _____
31
         rhoC : float
32
             The density of matter in the small region around the center
33
         const1 : float
             Constant to be used in the derivative, depends on the `rhoC` value.
35
             Value after the normalization.
         const2 : float
37
             Second constant to be used in the derivative for the relativistic case,
38
             depends on `rhoC`. Value after the normalization.
39
         Radius : float
40
             Physical radius of the star. Has a value only after `getRadius()` has
41
             been called.
^{42}
         Mass : float
43
             Mass of the star. Has a value only after `getRadius()` has been called.
44
45
         Methods
46
47
         getRadius
48
             Returns the tuple `(Radius, Mass)` of the star if the equation has been
49
             integrated.
50
```

```
Background
52
         _____
         The differential equation is a coupled system of the density `rho` and mass
54
         `m` as functions of the distance from the centre `r`. These have some
55
         dimensions in powers of kg and m. We transform the variables with some
56
         scaling constants so that the ODE solver works with dimensionless
57
         variables. The following choice of constants is made:
58
             rho = rhoC * theta,
59
             m = rhoC * mu,
             r = 1 * ksi,
61
         where rhoC is given as input by the user and 1^3*4*pi/3 = 1. The choice
62
         of 1 gives it a value on the order of 1, and we will choose it as our step
63
         size in r i.e. the step size in ksi is just 1. This is a convenient choice
         for normalization.
65
         0.00
         def __init__(self, rhoC: float, span: np.ndarray, regime: int=1) -> None:
68
             self.rhoC = rhoC
69
             # vector to be given as initial conditions, [rho, mass]
             init\_condit = np.array([1.0, 1.0*(span[0]**3)])
71
             # constants in the derivative; computed here so that they are computed
73
             # only once rather than every time the derivative func is called
74
             self.const1 = -(2**(13.0/3))*cs.pi*cs.gravitational_constant *\
                 cs.electron_mass*(cs.proton_mass**(5.0/3)) * \
76
                 (rhoC**(1.0/3))*(cs.h**(-2.0))
77
             self.const2 = (3*rhoC/16/cs.pi/cs.proton_mass *
78
                             (cs.h/cs.c/cs.electron_mass)**3)**(2.0/3)
80
             # choose the equation of state and pass constants, prepares a function
             # to be passed to the constructor of ODEinit accepting (x, y) params
82
             if regime == 1:
                 func = lambda a, b: derivatives.nonRelativGas(a, b, self.const1)
84
             elif regime == 2:
85
                 func = lambda a, b: derivatives.relativGas(a, b, self.const1, self.const2)
86
             elif regime == 3:
                 func = lambda a, b: derivatives.ultraRel(a, b, self.const1)
88
89
             super(whiteDwarf, self).__init__(init_condit, func, span)
90
91
         def getRadiusMass(self) -> tuple:
92
93
             Finds the radius and mass of the white dwarf, and returns them. Takes
             into account the normalization of the ODE.
95
96
             The function scans `yOut[:,0]` column containing `rho(r)` values. We
97
             know that this is a decreasing function; the first instance where it
             has a value <= 0 it where the star "ends".
99
100
             Returns
101
             _____
102
             tuple
103
                 Returns a tuple `(Radius, Mass)`. Radius and Mass are calculated
                 from `yOut` with the appropriate normalization constants. For
105
                 Radius it is l = (3/(4*pi))^(1/3), result is in m. For Mass it is
106
                 `rhoC`, result is in kg.
107
108
```

67

104

```
109
             # find index of rho = 0 (changing signs)
             if self.flagIntegrated:
111
                  index = np.searchsorted(-self.yOut[:, 0], 0)
112
                  # index is the position where we have yOut[index,:] ~ [0,0]
113
114
                 1 = (3/(4*np.pi))**(1.0/3.0)
                                                                 # normalization const
115
                  self.Radius = (index-1)*self.interval*l
                                                                 # in m
116
                  self.Mass = self.yOut[index-1, 1]*self.rhoC # in kg*m^(-3)
                  return self.Radius, self.Mass
118
             else:
119
                  print("Integrate the eqn first")
120
                 return None, None
121
122
     if __name__=="__main__":
123
         print(
124
              "This file contains declaration of a white dwarf class.")
125
```

```
Integrators Library(`integrators.py`)
    _____
    This is a home to the different integrators used alongside the target of the
    exercise, namely Runge-Kutta 4th order (rk4) method. We implement two other
    methods to evaluate the advantages of the rk4 method, both in its accuracy and
    precision.
    Integration Methods implemented
10
11
12
        euler
                Euler method - 1st order.
13
        heun
                Heun method - a variant of a 2nd order.
14
        rk4
                Runge-Kutta 4th order method.
15
16
    Implementation notes
17
    All of the functions defined are to act on np.ndarray structures, therefore
19
    should be element-wise functions; all of the magic is done for us by numpy.
20
    The differential equation which they iterate is set up as dy/dx = f(x, y),
21
    where `y` and `f` are N-dimensional vectors. Hence, all of the functions
22
    receive four parameters:
23
        x : float, the point at which we start computing
25
        y : array_like, the values of the y-vector at x
        deriv : function, the derivative of y to be computed at some (x, y) coords
27
        h : float, the interval size
28
29
    All of the functions return the value of `y` for the next place in the
30
    integration range. How we determine its value, or more precisely the increment
31
    from the previous value, is the difference between these methods.
32
33
    Error handling is implemented in `rk4` (the one used for actual calculations),
34
    where evaluating the `deriv` function might not be possible.
35
36
    Choice of an integrator is aided by the dictionary `index`, mapping integer
37
    values to the integrator functions.
38
    0.01\,0
39
```

```
40
    import numpy as np
42
    def euler(x: float, y: np.ndarray, deriv, h: float) -> np.ndarray:
43
44
        Compute an iteration using Euler's method: a 1st order method.
45
46
        This function computes an iteration in the numerical integration of an ODE
47
        of the form dy/dx = f(x, y) using the first order Euler method:
        y(x + h) = y(x) + h*f(x, y(x)), where h is the size of the step in the
49
        x-coordinate.
50
51
        Parameters
52
        _____
53
        x : float
54
            Starting coordinate in x.
55
        y : ndarray
56
            Starting coordinate in y. In general can be an N-dimensional array.
57
        deriv : function
58
            A function which computes the derivative of the y-vector given the
59
            x and y coords.
60
        h : float
61
            The step size in x.
62
        Returns
64
        -----
65
        ndarray
66
            The returned value is the approximation of y(x + h) using the Euler
            algoritm.
68
        .....
        return y + h*deriv(x,y)
70
    def heun(x: float, y: np.ndarray, deriv, h: float) -> np.ndarray:
72
73
        Compute an iteration using Heun's method, a kind of 2nd order method.
74
75
        This function computes an iteration in the numerical integration of an ODE
76
        of the form dy/dx = f(x, y) using the second order Heun method:
77
        y(x + h) = y(x) + 0.5*h*(f(x, y(x)) + f(x + h, y + h*f(x, y))), where h
78
        is the size of the step in the x-coordinate. This is a predictor-corrector
79
        method, and uses the prediction of the slope at `x + h` to correct the
80
        final calculation of the slope to be added.
81
        Parameters
83
         -----
84
        x : float
85
            Starting coordinate in x.
        y : ndarray
87
            Starting coordinate in y. In general can be an N-dimensional array.
        deriv : function
89
            A function which computes the derivative of the y-vector given the
90
            x and y coords.
91
        h : float
            The step size in x.
93
        Returns
95
        _____
96
```

41

71

```
np.ndarray
97
             The returned value is the approximation of y(x + h) using the Heun
             algorithm.
99
         .....
100
         return y + 0.5*h*(deriv(x, y) + deriv(x + h, y + h*deriv(x,y)))
101
102
     def rk4(x: float, y: np.ndarray, deriv, h: float) -> np.ndarray:
103
104
         Compute an iteration using a 4th order Runge-Kutta method.
106
         This function computes an iteration in the numerical integration of an ODE
107
         of the form dy/dx = f(x, y) using a fourth order Runge-Kutta method.
108
109
         Parameters
110
         _____
111
         x : float
112
             Starting coordinate in x.
113
         y : ndarray
114
             Starting coordinate in y. In general can be an N-dimensional array.
115
         deriv : function
116
             A function which computes the derivative of the y-vector given the
117
             x and y coords.
118
         h : float
119
             The step size in x.
120
121
         Returns
122
123
         np.ndarray
124
             The returned value is the approximation of y(x + h) using the 4th order
125
             Runge-Kutta algorithm.
126
127
         Raises
128
129
         ValueError
130
             If an error is raised by the deriv function indicating impossible
131
             evaluation, raise an error to indicate the termination of integration.
132
133
134
         Notes
         ____
135
         Note that in the case deriv cannot be computed for any of the steps, the
136
         error is handled by raising a value error back to the integrator method of
137
         the ODE class where this is handled. Additionally, an improvement in the
138
         calculation order is implemented as per [1]_, where calculation cycle
139
         number is reduced, as well as memory usage.
140
141
         Sources
142
         ..[1] Press, William H., Saul A. Teukolsky, William T. Vetterling, and
144
         Brian P. Flannery. "Numerical Recipes : The Art of Scientific Computing."
145
         Third ed. Cambridge, 2007.
146
         11 11 11
147
         # TODO think of fix
148
         try:
149
             # USE OLD implementation, uses more cycles and memory (allegedly)
150
151
             k1 = h*deriv(x,y)
             k2 = h*deriv(x + 0.5*h, y + 0.5*k1)
152
             k3 = h*deriv(x + 0.5*h, y + 0.5*k2)
153
```

```
k4 = h*deriv(x + h, y + k3)
154
             # 'FASTER' implementation - produced an error while running tests on
156
             # the SHO for the report, revert to the more abstract version - some
157
             # cycles but sure it is correct
158
159
             # hh = 0.5*h
160
             # h6 = h/6.0
161
             # xh = x + hh
162
             # k1 = deriv(x, y)
163
             # yTemp = y + hh*k1
164
             # k2 = deriv(xh, yTemp)
165
             # yTemp = y + hh*k2
166
             # k3 = deriv(xh, yTemp)
167
             # yTemp = y + k3
168
             # k3 += k2
169
             # k2 = deriv(x+h, yTemp)
170
         except ValueError:
171
             raise ValueError("Invalid integration, terminate integration!")
172
173
         return y + (k1 + k4)/6.0 + (k2 + k3)/3.0
174
         # return y + h6*(k1 + k2 + 2*k3)
175
176
     # dictionary to choose integrator; for neater code in modules.ode
177
     index = {1: euler, 2: heun, 3: rk4}
178
179
     if __name__=="__main__":
180
         print(
181
             "This is a module home to several integrators to be used in solving \
182
     ODEs: Euler, Heun, Runge-Kutta4.")
183
```

```
Derivatives Library(`derivatives.py`)
2
3
4
    This is a home to several functions which give the derivative `f` in a system
    of linear ODEs y'(x) = f(x, y) in functional form.
6
7
    White Dwarf Functions
8
    nonRelativGas
                         Implements the eqn. of state for non-relativistic electrons
10
    relaitvGas
                         Implements the eqn. of state for relativistic electrons.
11
12
    Testing Functions
13
    ______
14
    SHO
                         Implements the Simple Harmonic Oscillator.
15
16
    Implementation Notes
17
18
    Choice of an equation of state for the white dwarf is aided by the dictionary
19
    `index`, mapping integer values to the integrator functions.
20
21
22
    import numpy as np
23
24
    def SHO(x: float, y: np.ndarray) -> np.ndarray:
25
        11 11 11
26
```

Function of the derivative for the simple harmonic oscillator written as a 27 system of linear ODEs. 29 Parameters 30 -----31 x : float 32 Value of the independent variable in the ODE. 33 y : np.ndarray 34 Value of the dependent variable in the ODE. Two element vector [z', z], where the SHO ODE is z''(x) + z(x) = 0. 36 37 Returns 38 np.ndarray 40 The value of the derivative at (x, y) 41 42 Notes 43 44 The equation z''(x) + z(x) = 0 is written using p = z' in the form: y' = [p, z]' = [-z, p]. The function returns the latter vector. 46 47 return np.array([-4*np.pi*np.pi*y[1], y[0]]) 48 49 def nonRelativGas(x: float, y: np.ndarray, const: float) -> np.ndarray: 50 51 Function for the coupled ODE using a non-relativistic equation of state. 52 53 Function of the derivative for the system of coupled equations 54 `y = [rho, m](x)` in the non-relativistic case. An additional argument 55 `const` is passed to save computational time - it is calculated only once in the ODE definition, and called as value to be used in the computations. 57 Parameters 59 60 x : float 61 Value of the independent variable in the ODE. y : np.ndarray 63 Value of the dependent variable in the ODE. *Two elements only.* 64 const: float 65 The scaling constant used in the definition. 66 67 Returns 68 _____ 69 np.ndarray 70 The value of the derivative at (x, y). Look at Notes for info on the 71 functional form. 72 Raises 74 ValueError 76 If the value y[0] < 0, the computation is discarded on physical 77 grounds - cannot have negative matter density. Raises an exception 78 propagated by the integrator to be handled in the .integration method. 80 81 Notes 82 The functional form is determined by the equation of state in the 83

```
non-relativistic case, where `pressure = someConst * rho^(2/3)`.The final
84
         form: [rho, m]' = [const1*m*rho^(1/3)/x^2, 3*x^2*rho].
86
87
         # need y[0] >= 0 to be exponentiated apart from physical grounds
88
         # this catches the moment y[0] becomes too small - only it is on the
89
         # computation of the following value
90
         if y[0] < 1e-10:
91
             raise ValueError
         return np.array([const*y[1]*(y[0]**(1.0/3))*(x**(-2.0)),
93
                              3*(x**2)*y[0]
94
95
     def relativGas(x: float, y: np.ndarray, const1: float ,const2: float) -> np.ndarray:
96
97
         Function for the coupled ODE using a relativistic equation of state.
99
         Function of the derivative for the system of coupled equations
100
         `y = [rho, m](x)` in the relativistic case. Two additional arguments
101
         `const1` and `const2` are passed to save on computational time - they are
102
         calculated only once in the ODE definition, and called as value to be used
103
         in the computations.
104
105
         Parameters
106
         _____
107
         x : float
108
             Value of the independent variable in the ODE.
109
         y : np.ndarray
110
             Value of the dependent variable in the ODE. *Two elements only.*
111
         const1: float
112
             One of the scaling constants, used in the derivative of rho.
113
         const2 : float
114
             The second scaling constant, used in the relativistic correction.
115
116
117
         Returns
118
         np.ndarray
119
             The value of the derivative at (x, y). Look at Notes for info on the
120
             functional form.
121
122
         Raises
123
124
         ValueError
125
             If the value y[0] < 0, the computation is discarded on physical
126
             grounds - cannot have negative matter density. Raises an exception
127
             propagated by the integrator to be handled in the .integration method.
128
129
         Notes
131
         The functional form is determined by the equation of state in the
132
         relativistic case, where `pressure = someConst*rho^(2/3)*relCorr`.The
133
         relativistic correction is `relCorr = sqrt(1+const2*rho^(2/3))^(-1)`.
134
         The final form: \[ \text{rho, m} \]' = [\text{const1*m*rho}(1/3)\text{*relCorr/x}^2, 3\text{*x}^2\text{*rho}] \].
135
         .....
136
137
138
         # need y[0] >= 0 to be exponentiated apart from physical grounds
         # this catches the moment y[0] becomes too small - only it is on the
139
         # computation of the following value
140
```

```
if y[0] < 1e-10:</pre>
141
             raise ValueError
143
         return np.array([const1*y[1]*(y[0]**(1.0/3))*(x**(-2.0))*
144
                                (1+const2*(y[0]**(2.0/3)))**(0.5), 3*(x**2)*y[0]])
145
146
     index = {1: nonRelativGas, 2: relativGas}
147
148
     if __name__=="__main__":
149
         print(
150
              "This is a home to several functions which give the derivative in a \
151
     system of linear ODEs.")
152
```

A.2 Code to run the experiment and testing procedures

```
"""main.py
1
    The main script in this experiment. It calculates the values of the white dwarf
    radius and mass for different central densities (initial conditions to solve
    the ODE) by using our purpose built library and outputs them to a .csv file.
5
    # parallel computing library
7
    from multiprocessing import Pool, cpu_count
    import numpy as np
10
    # for saving files to csv (easier than the numpy implementation)
11
    import pandas as pd
12
    # solar mass for output scale
13
    from astropy.constants import M_sun
14
    # for setup of integration grid
15
    from numpy.core.function_base import linspace
16
17
    # our white dwarf integrator module
18
    import modules
20
21
    # Initialize a white dwarf with non-relativistic equation of state
22
    def starInitNonRelativ(rhoC):
        """This function initializes a white dwarf with a non-relativistic equation
24
        of state and calculates its radius and mass, then returns the values.
26
        Parameters
27
        _____
28
        rhoC : float
29
            The density of the center in kg*m^(-3), used to set the initial
30
            conditions for integration.
31
32
        Returns
33
        list
35
            Returns a three element list of the density in in kg*m^(-3), the radius
            in km/1000, and the mass in solar mass in this order.
37
38
        # Integration grid in normalized units, optimal step size determined on
39
        # convergence ground for more info look at convergence.py
        span = linspace(1,8.1e7,num=2000000)
41
```

```
# Initialize the white dwarf and integrate
42
        star = modules.whiteDwarf(rhoC, span, 1)
        star.integrate(3)
44
45
        radius, mass = star.getRadiusMass()
46
        # return the three values rho in kg*m^(-3), radius in km/1000, mass in solar mass
47
        return [rhoC, radius/1000, mass/M_sun.value]
48
49
    # Initialize a white dwarf with relativistic equation of state
    def starInitRelativ(rhoC):
51
        """This function initializes a white dwarf with a relativistic equation of
52
        state and calculates its radius and mass, then returns the values.
53
        Parameters
55
        rhoC : float
57
            The density of the center in kg*m^(-3), used to set the initial
58
            conditions for integration.
59
        Returns
61
        _____
62
        list
63
            Returns a three element list of the density in in kg*m^(-3), the radius
64
            in km/1000, and the mass in solar mass in this order.
65
66
        # Integration grid, optimal step size determined on convergence ground
67
        # for more info look at convergence.py
68
        span = linspace(1,8.1e7,num=2000000)
69
        # Initialize the white dwarf and integrate
70
        star = modules.whiteDwarf(rhoC, span, 2)
71
        star.integrate(3)
72
        radius, mass = star.getRadiusMass()
74
        # return the three values rho in kg*m^(-3), radius in km/1000, mass in solar mass
75
        return [rhoC, radius/1000, mass/M_sun.value]
76
78
79
    def main():
        # the rho values to test in the range, change the list accordingly
80
        rhoVal = np.array([float(a)*10**b for b in range(6,15) for a in range(1,10)])
81
82
        # Implements parallel computation of the ODEs using multiple processes.
83
        # Computation is done over the range of central density values rhoVal.
85
        # Computation for non-relativistic gas. This process sets up the
86
        # multiprocessing and cleans up after the calculations are over.
        with Pool(cpu_count(), maxtasksperchild=1) as executor:
            # The executor.map(f, s) applies the function f to each member of the
89
            # iterable s and collects the results in a list following a First In
            # First Out procedure, so the results collected are ordered by the
91
            # order in which they were in s, i.e. by central density values.
92
            resNonRelativ = executor.map(starInitNonRelativ, rhoVal)
93
        # Save results for non-relativistic case
95
        dfNonRel = pd.DataFrame(resNonRelativ, columns=["rhoC", "radiusKM", "massSolMass"])
        dfNonRel.to_csv("nonRelativRes.csv", index=None)
97
98
```

```
# Computation for non-relativistic gas. This process sets up the
99
         # multiprocessing and cleans up after the calculations are over.
100
         with Pool(cpu_count()-6, maxtasksperchild=1) as executor:
101
             # The executor.map(f, s) applies the function f to each member of the
102
             # iterable s and collects the results in a list following a First In
103
             # First Out procedure, so the results collected are ordered by the
104
             # order in which they were in s, i.e. by central density values.
105
             resRelativ = executor.map(starInitRelativ, rhoVal)
106
         # Save results for relativistic case
108
         dfRel = pd.DataFrame(resRelativ, columns=["rhoC", "radiusKM", "massSolMass"])
109
         dfRel.to_csv("relativRes.csv", index=None)
110
111
112
     if __name__ == "__main__":
113
         main()
114
```

```
"""convergence.py
1
    This is an auxiliary script, which helps determine the convergence of
2
    computing white dwarf parameters and in turn judge the needed coarseness of
3
    the integration space. Many of the functions here are direct copies of the ones
    in main.py; look there for more explanation. Outputs .csv files of calculations.
5
    import math
7
    from multiprocessing import Pool, cpu_count
    # for saving files to csv (easier than the numpy implementation)
10
    import pandas as pd
11
    # solar mass for output scale
12
    from astropy.constants import M_sun
13
    # for setup of integration grid
14
    from numpy.core.function_base import linspace
15
16
    # our library
17
    import modules
18
20
    # Direct copy of the method in main.py, only added a variable for the number of
21
    # intervals in the space (since we are probing it).
22
    def starInitNonRelativ(myTuple):
23
        rhoC, n = myTuple
24
        span = linspace(1,8.1e7,num=n)
25
        star = modules.whiteDwarf(rhoC, span, 1)
26
        star.integrate(3)
28
        radius, mass = star.getRadiusMass()
29
        # return the three values No of iter, radius in km/1000, mass in solar mass
30
        return [n, radius/1000, mass/M_sun.value]
31
32
    # Direct copy of the method in main.py, only added a variable for the number of
33
    # intervals in the space (since we are probing it).
34
    def starInitRelativ(myTuple):
35
        rhoC, n = myTuple
36
        span = linspace(1,8.1e7,num=n)
37
        star = modules.whiteDwarf(rhoC, span, 2)
38
        star.integrate(3)
39
```

```
radius, mass = star.getRadiusMass()
41
        # return the three values No of iter, radius in km/1000, mass in solar mass
        return [n, radius/1000, mass/M_sun.value]
43
    numberOfIterations = [int(float(a)*10**b) for b in range(2,8) for a in [1,2,4,8]]
45
46
    # wrapped all actions in a main function because of the multiprocessing
47
    def main():
48
        # Loop the three densities - two at the ends of the interval and one at the middle
        # multiprocessing copied from main.py
50
        for density in [1e6, 1e10, 1e14]:
51
            # quick hack to prepare an iterable for the multiprocessing pool
52
            # function, since our function needs two parameters
            poolIterable = [(density, n) for n in numberOfIterations]
54
            with Pool(cpu_count(), maxtasksperchild=1) as executor:
                # The executor.map(f, s) applies the function f to each member of the
56
                # iterable s and collects the results in a list following a First In
                # First Out procedure, so the results collected are ordered by the
58
                # order in which they were in s, i.e. by central density values.
                resNonRelativ = executor.map(starInitNonRelativ, poolIterable)
60
61
            # Save results for non-relativistic case
62
            dfNonRel = pd.DataFrame(resNonRelativ, columns=["NoIter", "radiusKM", "massSolMass"])
63
            dfNonRel.to_csv(
                "convergenceE{}nonRelativ.csv".format(int(math.log10(density))),
65
                index=None
66
            )
67
            with Pool(cpu_count()-6, maxtasksperchild=1) as executor:
69
                # The executor.map(f, s) applies the function f to each member of the
                # iterable s and collects the results in a list following a First In
71
                # First Out procedure, so the results collected are ordered by the
                # order in which they were in s, i.e. by central density values.
73
                resRelativ = executor.map(starInitRelativ, poolIterable)
75
            # Record results to csv file
            dfRelativ = pd.DataFrame(resRelativ, columns=["NoIter", "radiusKM", "massSolMass"])
77
            dfRelativ.to_csv(
                "convergenceE{}Relativ.csv".format(int(math.log10(density))),
79
                index=None
80
            )
81
82
    if __name__ == '__main__':
84
        main()
85
```

```
# Test the error with the integrators and plot residuals
# NOTE we need some amount of iterations comparable to the white dwarf to
# compate the methods i.e. 1e6
from math import cos, sin

import matplotlib as mpl
import matplotlib.pyplot as plt
import numpy as np
import pandas as pd
from numpy.core.function_base import linspace
```

```
import modules
12
    from modules.derivatives import SHO
13
14
15
    def plots():
16
        # load data
17
        eul1= pd.read_csv("method1.csv")
18
        heun= pd.read_csv("method2.csv")
19
        rk4= pd.read_csv("method3.csv")
20
        eul2= pd.read_csv("method4.csv")
21
22
        # theoretical data
23
        x0 = 1.0
        step = 0.001
25
        span = np.array(step*linspace(1,1000000,num=1000000))
        # for eul1, heun, rk4
27
        theory = np.concatenate(([1], x0*np.cos(2*np.pi*span)))
28
29
        # 2*energy in systems/m
        c = 4*np.pi*np.pi
31
        enEul1 = eul1["xdot"]**2 + c*eul1["x"]**2
32
        enEul2 = eul2["xdot"]**2 + c*eul2["x"]**2
33
        enHeun = heun["xdot"]**2 + c*heun["x"]**2
34
        enRK4 = rk4["xdot"]**2 + c*rk4["x"]**2
35
36
        ## plot 1 - oscill of eul1, eul2, heun
37
        plt.figure(1)
38
        plt.plot(eul1.index/1000, eul1["x"])
39
        plt.plot(eul2.index/1000, eul2["x"])
40
        plt.plot(heun.index/1000, heun["x"])
41
        #axis setup
42
        plt.xlim(0, 20)
        plt.xlabel("time t, s")
44
45
        plt.ylim(-2,2)
        plt.ylabel("position x(t)")
46
        plt.legend(["euler1", "euler2", "heun"])
        plt.show
48
        plt.savefig("eul12+heun.pdf")
49
50
        ## plot2 - (oscill - theory) of eul1, eul2, heun
51
        plt.figure(2)
52
        plt.plot(eul1.index/1000, eul1["x"] - theory)
53
        plt.plot(eul2.index/1000, eul2["x"] - theory)
        plt.plot(heun.index/1000, heun["x"] - theory)
55
        #axis setup
56
        plt.xlim(0, 20)
57
        plt.xlabel("time t, s")
        plt.ylim(-0.5,0.5)
59
        plt.ylabel("position x(t) - theory")
        plt.legend(["euler1", "euler2", "heun"], loc="upper left")
61
        plt.show
62
        plt.savefig("eul12+heun-theory.pdf")
63
        ## plot3 - energy of eul1, eul2, heun
65
        plt.figure(3)
        plt.plot(enEul1.index/1000, abs(enEul1-c)/c)
67
        plt.plot(enEul2.index/1000, abs(enEul2-c)/c)
68
```

```
plt.plot(enHeun.index/1000, abs(enHeun-c)/c)
69
         #axis setup
         plt.xlim(0, 200)
71
         plt.xlabel("time t, s")
         plt.ylim(1e-7,100000)
73
         plt.yscale('log')
74
         plt.ylabel("energy $\Delta E/E_0$")
75
         plt.legend(["euler1", "euler2", "heun"], loc="upper left")
76
         plt.show
         plt.savefig("eul12+heun+energy.pdf")
78
79
         ## plot4 - energy of heun, rk4
80
         plt.figure(4)
         plt.plot(enHeun.index/1000, abs(enHeun-c)/c, color='green')
82
         plt.plot(enRK4.index/1000, abs(enRK4-c)/c, color='red')
         #axis setup
         plt.xlabel("time t, s")
85
         plt.ylim(5e-7,1e-3)
86
         plt.yscale('log')
         plt.ylabel("energy $\Delta E/E_0$")
88
         plt.legend(["heun", "rk4"], loc="upper left")
89
         plt.show
90
         plt.savefig("heunRK4+energy.pdf")
91
92
93
     def main():
94
         # initial parameters of the SHO
95
         x0 = 1.0
         step = 0.001
97
         y0 = x0*np.array([-sin(step), cos(step)])
         span = step*linspace(1,1000000,num=1000000)
99
         # do integration for the three methods
101
         for integ in [1,2,3]:
102
             ode = modules.ODEinit(y0, SHO, span)
103
             ode.integrate(integ)
104
             res = np.vstack((np.array([0,1]), ode.yOut))
105
             dfOut = pd.DataFrame(res, columns=["xdot", "x"])
106
             dfOut.to_csv("method{}.csv".format(integ), index=None)
107
108
109
         # additional integration for the euler method with a step 1/10th of original
         span2 = 0.1*step*linspace(1,10000000,num=10000000)
110
         ode = modules.ODEinit(y0, SHO, span2)
111
         ode.integrate(1)
112
         res = np.vstack((np.array([0,1]), ode.yOut))
113
         # keep the data only at the same moments as the other ones
114
         dfOut = pd.DataFrame(res[::10], columns=["xdot", "x"])
         dfOut.to_csv("method4.csv", index=None)
116
117
         plots()
118
119
120
     if __name__=='__main__':
121
         main()
122
```

```
# mainPlots.py
# Plots of data for convergence and white dwarf star parameters.
```

```
from matplotlib import colors
    import matplotlib.pyplot as plt
    import numpy as np
5
    import pandas as pd
    def convergencePlots():
9
        # load in data
10
        E6nonRelativ = pd.read_csv("convergenceE6nonRelativ.csv")
11
        E6Relativ = pd.read_csv("convergenceE6Relativ.csv")
12
        E10nonRelativ = pd.read_csv("convergenceE10nonRelativ.csv")
13
        E10Relativ = pd.read_csv("convergenceE10Relativ.csv")
14
        E14nonRelativ = pd.read_csv("convergenceE14nonRelativ.csv")
        E14Relativ = pd.read_csv("convergenceE14Relativ.csv")
16
        # subplot setup, tighter spacing
18
        f, axs = plt.subplots(4,2,figsize=(11.2,17.15),constrained_layout=False)
19
20
        # mass, non-relativistic
        # values to which result converges
22
        e6convM = sum(E6nonRelativ["massSolMass"][-3:])/3
23
        e10convM = sum(E10nonRelativ["massSolMass"][-3:])/3
24
        e14convM = sum(E14nonRelativ["massSolMass"][-3:])/3
25
26
        # plots of delta m / m_conv
27
        plt.subplot(4,2,1)
28
        plt.title("Non-relativistic case")
29
        plt.plot(E6nonRelativ["NoIter"], abs(E6nonRelativ["massSolMass"]-e6convM)/e6convM)
30
        plt.plot(E10nonRelativ["NoIter"], abs(E10nonRelativ["massSolMass"]-e10convM)/e10convM)
31
        plt.plot(E14nonRelativ["NoIter"], abs(E14nonRelativ["massSolMass"]-e14convM)/e14convM)
32
        plt.xscale("log")
33
        plt.ylabel("$\Delta M/M_\mathrm{conv}$")
        plt.legend(["$\\rho_\mathrm{C} = 10^6$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
35
                36
                \label{local_condition} $$ \ \cdot \mathbf{0}^{14}\  \ \cdot \mathbf{mathrm{m}^{-3}}"],
37
                loc="upper right")
        # second fig
39
        plt.subplot(4,2,3)
        plt.plot(E6nonRelativ["NoIter"], abs(E6nonRelativ["massSolMass"]-e6convM)/e6convM)
41
        plt.plot(E10nonRelativ["NoIter"], abs(E10nonRelativ["massSolMass"]-e10convM)/e10convM)
42
        plt.plot(E14nonRelativ["NoIter"], abs(E14nonRelativ["massSolMass"]-e14convM)/e14convM)
43
        plt.xscale("log")
44
        plt.yscale("log")
45
        plt.ylabel("$\Delta M/M_\mathrm{conv}$")
46
        plt.legend(["$\\rho_\mathrm{C} = 10^6$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
47
                48
                \label{local_condition} $$ \ \cdot \mathbf{0}^{14}\  \ \cdot \mathbf{mathrm{m}^{-3}}"],
                loc="upper right")
50
        # mass, relativistic
52
        # converged to values
53
        e6RconvM = sum(E6Relativ["massSolMass"][-3:])/3
54
        e10RconvM = sum(E10Relativ["massSolMass"][-3:])/3
        e14RconvM = sum(E14Relativ["massSolMass"][-3:])/3
56
        # plots of delta m / m_conv
58
        plt.subplot(4,2,2)
59
```

```
plt.title("Relativistic case")
60
        plt.plot(E6Relativ["NoIter"], abs(E6Relativ["massSolMass"]-e6RconvM)/e6RconvM)
        plt.plot(E10Relativ["NoIter"], abs(E10Relativ["massSolMass"]-e10RconvM)/e10RconvM)
62
        plt.plot(E14Relativ["NoIter"], abs(E14Relativ["massSolMass"]-e14RconvM)/e14RconvM)
63
        plt.xscale("log")
        plt.ylabel("$\Delta M/M_\mathrm{conv}$")
65
        plt.legend(["$\\rho_\mathrm{C} = 10^6$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
66
                "$\\rho_\mathrm{C} = 10^{10}$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
67
                "$\\rho_\mathrm{C} = 10^{14}$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$"],
                loc="upper right")
69
        # log plot
70
        plt.subplot(4,2,4)
71
        plt.plot(E6Relativ["NoIter"], abs(E6Relativ["massSolMass"]-e6RconvM)/e6RconvM)
        plt.plot(E10Relativ["NoIter"], abs(E10Relativ["massSolMass"]-e10RconvM)/e10RconvM)
73
        plt.plot(E14Relativ["NoIter"], abs(E14Relativ["massSolMass"]-e14RconvM)/e14RconvM)
        plt.xscale("log")
75
        plt.yscale("log")
76
        plt.ylabel("$\Delta M/M_\mathrm{conv}$")
77
        plt.legend(["$\\rho_\mathrm{C} = 10^6$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
                "$\\rho_\mathrm{C} = 10^{10}$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
79
                "$\\rho_\mathrm{C} = 10^{14}$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$"],
80
                loc="upper right")
81
82
        # radius, non relativistic
        # converged to values
        e6convR = sum(E6nonRelativ["radiusKM"][-3:])/3
85
        e10convR = sum(E10nonRelativ["radiusKM"][-3:])/3
86
        e14convR = sum(E14nonRelativ["radiusKM"][-3:])/3
88
        # plots of delta m / m_conv
        plt.subplot(425)
90
        plt.plot(E6nonRelativ["NoIter"], abs(E6nonRelativ["radiusKM"]-e6convR)/e6convR)
        plt.plot(E10nonRelativ["NoIter"], abs(E10nonRelativ["radiusKM"]-e10convR)/e10convR)
92
        plt.plot(E14nonRelativ["NoIter"], abs(E14nonRelativ["radiusKM"]-e14convR)/e14convR)
93
        plt.xscale("log")
94
        plt.ylabel("$\Delta R/R_\mathrm{conv}$")
        plt.legend(["$\\rho = 10^6$ \mathbf{\xi} \cdot \mathbf{m}^{-3}$",
96
                       "$\\rho_\mathrm{C} = 10^{10}$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
                       loc="upper right")
        # log plot
100
        plt.subplot(427)
101
        plt.plot(E6nonRelativ["NoIter"], abs(E6nonRelativ["radiusKM"]-e6convR)/e6convR)
102
        plt.plot(E10nonRelativ["NoIter"], abs(E10nonRelativ["radiusKM"]-e10convR)/e10convR)
103
        plt.plot(E14nonRelativ["NoIter"], abs(E14nonRelativ["radiusKM"]-e14convR)/e14convR)
104
        plt.xscale("log")
105
        plt.yscale("log")
        plt.xlabel("Number of iterations")
107
        plt.ylabel("$\Delta R/R_\mathrm{conv}$")
108
        plt.legend(["$\\rho_\mathrm{C} = 10^6$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
109
                       110
                       111
                       loc="upper right")
113
        # radius, relativistic
114
        # converged to values
115
        e6RconvR = sum(E6Relativ["radiusKM"][-3:])/3
116
```

```
e10RconvR = sum(E10Relativ["radiusKM"][-3:])/3
117
         e14RconvR = sum(E14Relativ["radiusKM"][-3:])/3
119
         # plots of delta m / m_conv
120
         plt.subplot(426)
121
         plt.plot(E6Relativ["NoIter"], abs(E6Relativ["radiusKM"]-e6RconvR)/e6RconvR)
122
         plt.plot(E10Relativ["NoIter"], abs(E10Relativ["radiusKM"]-e10RconvR)/e10RconvR)
123
         plt.plot(E14Relativ["NoIter"], abs(E14Relativ["radiusKM"]-e14RconvR)/e14RconvR)
124
         plt.xscale("log")
         plt.ylabel("$\Delta R/R_\mathrm{conv}$")
126
         plt.legend(["$\\rho_\mathrm{C} = 10^6$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
127
                         "$\\rho_\mathrm{C} = 10^{10}$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
128
                         loc="upper right")
130
         plt.subplot(428)
131
         plt.plot(E6Relativ["NoIter"], abs(E6Relativ["radiusKM"]-e6RconvR)/e6RconvR)
132
         plt.plot(E10Relativ["NoIter"], abs(E10Relativ["radiusKM"]-e10RconvR)/e10RconvR)
133
         plt.plot(E14Relativ["NoIter"], abs(E14Relativ["radiusKM"]-e14RconvR)/e14RconvR)
134
         plt.xscale("log")
135
         plt.yscale("log")
136
         plt.xlabel("Number of iterations")
137
         plt.ylabel("$\Delta R/R_\mathrm{conv}$")
138
         plt.legend(["$\\rho_\mathrm{C} = 10^6$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
139
                         "$\\rho_\mathrm{C} = 10^{10}$ $\mathrm{kg} \cdot \mathrm{m}^{-3}$",
                         "^{-3}" \\rho_\mathrm{C} = 10^{14}\$ \mathrm{kg} \cdot \mathrm{m}^{-3}\$"],
141
                         loc="upper right")
142
143
         # save fig, clip white margins to fit on report page with caption
144
         plt.savefig("convergencePlot.pdf", bbox_inches="tight")
145
146
     def whiteDwarfPlots():
147
         # load data
         nonRres = pd.read_csv("nonRelativRes.csv")
149
         Rres = pd.read_csv("relativRes.csv")
150
151
         # fig 1 radius(rhoc)
         plt.figure()
153
         plt.plot(nonRres["rhoC"], nonRres["radiusKM"])
154
         plt.plot(Rres["rhoC"], Rres["radiusKM"])
155
         plt.xscale("log")
156
         plt.yscale("log")
157
         plt.xlabel("Central density, $\\rho_\mathrm{C}$, $\mathrm{kg} \cdot \mathrm{m}^{-3}$")
158
         plt.ylabel("Radius, km")
         plt.legend(["Non-relativistic", "Relativistic"])
160
         plt.savefig("radius-rhoC.pdf")
161
162
         # mass(rhoC)
         plt.figure()
164
         plt.plot(nonRres["rhoC"], nonRres["massSolMass"])
165
         plt.plot(Rres["rhoC"], Rres["massSolMass"])
166
         plt.axhline(1.434, c="grey", ls='--', lw=0.5)
167
         plt.xscale("log")
168
         plt.yscale("log")
169
         plt.xlabel("Central density, $\\rho_\mathrm{C}$, $\mathrm{kg} \cdot \mathrm{m}^{-3}$")
170
         plt.ylabel("Mass, $M_\mathrm{sol}$")
         plt.legend(["Non-relativistic", "Relativistic", "Ch. Mass limit"])
172
         plt.savefig("mass-rhoC.pdf")
173
```

```
174
         # radius(mass)
175
         plt.figure()
176
         plt.plot(nonRres["massSolMass"], nonRres["radiusKM"])
177
         plt.plot(Rres["massSolMass"], Rres["radiusKM"])
178
         plt.axvline(1.434, c="grey", ls='--', lw=0.5)
179
         plt.xlim(0, 2.1)
180
         plt.ylim(0, 20000)
181
         plt.xlabel("Mass, $M_\mathrm{sol}$")
         plt.ylabel("Radius, km")
183
         plt.legend(["Non-relativistic", "Relativistic", "Ch. Mass limit"])
184
         plt.savefig("radius-mass.pdf")
185
     def main():
187
         convergencePlots()
188
         whiteDwarfPlots()
189
190
     if __name__ == "__main__":
191
         main()
```