

The background evolution of the universe

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Abstract. I set up the background cosmology of the universe.

1. Introduction

In this project I will follow the algorithm presented in Callin (2005)[1] for simulating the cosmic microwave background. This is part one of four for this project. The first part involves computing the expansion history of the universe, as well as looking at the evolution of the density of various matter and energy components. This sets up the background cosmology such that we in the next step can turn to perturbations. I have chosen to make this first part compatible with the inclusion of neutrinos. This can of course be removed by setting the neutrino density to zero and raising the radiation density accordingly.

To ease the development of this code, I have been provided with a skeleton code of the project. This code includes a variety of methods needed to solve the project. These methods will be explained as they are used throughout the project.

2. Equations

For the background cosmology I use the standard Friedmann-Lemaître-Robertson-Walker (FLRW) metric for flat space. This gives the line element.

$$\begin{aligned} ds^2 &= -dt^2 + a^2(t)(r^2(d\theta^2 + \sin^2\theta d\phi^2)) \\ &= a^2(\eta)(-d\eta^2 + r^2(d\theta^2 + \sin^2\theta d\phi^2)) \end{aligned} \quad (1)$$

where $a(t)$ is the scale factor, and η conformal time. I also introduce a parameter x defined as

$$x = \ln(a) \quad (2)$$

The redshift is defined as

$$1 + z = \frac{a_0}{a} \quad (3)$$

We assume that the universe consist of cold dark matter (CDM, m), baryons (b), radiation(r), neutrinos(ν), and a cosmological constant (Λ). With these components, the Hubble parameter H becomes

$$H = \frac{1}{a} \frac{da}{dt} = H_0 \sqrt{(\Omega_m + \Omega_b)a^{-3} + (\Omega_r + \Omega_\nu)a^{-4} + \Omega_\Lambda}. \quad (4)$$

I also introduce a scaled H , namely

$$\begin{aligned} \mathcal{H} &= \frac{1}{a} \frac{da}{d\eta} \equiv \frac{\dot{a}}{a} = aH \\ &= H_0 \sqrt{(\Omega_m + \Omega_b)a^{-1} + (\Omega_r + \Omega_\nu)a^{-2} + \Omega_\Lambda a^2}. \end{aligned} \quad (5)$$

Note that the dot means derivative with respect to conformal time. Why this is useful will become apparent later. We also want the derivative of this with respect to x .

$$\frac{d\mathcal{H}}{dx} = H_0 \frac{-(\Omega_m + \Omega_b)e^{-x} - 2(\Omega_r + \Omega_\nu)e^{-2x} + 2\Omega_\Lambda e^{2x}}{\sqrt{(\Omega_m + \Omega_b)e^{-x} + (\Omega_r + \Omega_\nu)e^{-2x} + \Omega_\Lambda e^{2x}}}. \quad (6)$$

Ω_x is the relative density of component x compared to the critical density ρ_c needed for the universe to be flat.

$$\Omega_x = \frac{\rho_x}{\rho_c} \quad (7)$$

$$\rho_c = \frac{3H^2}{8\pi G} \quad (8)$$

We want to keep track of the densities of each component.

$$\rho_m = \rho_{m,0}a^{-3} \quad (9)$$

$$\rho_b = \rho_{b,0}a^{-3} \quad (10)$$

$$\rho_r = \rho_{r,0}a^{-4} \quad (11)$$

$$\rho_\nu = \rho_{\nu,0}a^{-4} \quad (12)$$

$$\rho_\Lambda = \rho_{\Lambda,0} \quad (13)$$

We will also need to know the distance to the particle horizon at different times. This can be found by noting that

$$\frac{d\eta}{dt} = \frac{c}{a}$$

which can be rewritten such that

$$\frac{d\eta}{da} = \frac{c}{a\mathcal{H}} \quad (14)$$

There are two ways to solve this differential equation numerically. One can either integrate directly, or one can use an ordinary differential equation solver (ODE solver). Since the skeleton code contains an ODE solver I have chosen the latter. (See section 3 for more information).

3. Implementation

The programming language of choice is Fortran90. This is chosen because of its speed, and because the skeleton code provided was written in it.

The first step is to set up arrays for a , x , η , all five Ω_x , and ρ_x . We will also need arrays for ρ_c , H , and z .

We then set the first x value to correspond to $a = 10^{-10}$, and the last to $a = 0$ (today). I've chosen to use 1000 points for these arrays and make the steps equal in the x array. After that one computes the a , and z values for each of the points in this x array. Because of this we now have linear steps in x . With that done, one computes H , \mathcal{H} , ρ_x , and Ω_x for each x value.

The last thing to find is $\eta(x)$. To do this we need to use an ODE solver on equation 14. For this we must have some initial value for the function. The equation we then want to solve is

$$\eta(a) = \int_0^a \frac{da'}{a' \mathcal{H}(a')} \quad (15)$$

This can be found by considering that $a\mathcal{H} \rightarrow H_0 \sqrt{\Omega_r + \Omega_\nu}$ as $a \rightarrow 0$. This gives

$$\eta(a) = \frac{a}{H_0 \sqrt{\Omega_r + \Omega_\nu}} \quad (16)$$

for small a . And thus one has an initial value.

Plugging this into the ODE solver returns $\eta(x)$ for all values in the x array. Thus completing all calculations needed for the first part of the project. The ODE solver uses the Bulirsch–Stoer algorithm. The steplength is set to be one hundredth the length between two neighbouring x values.

As preparation for the next part I also spline the resulting $\eta(x)$ values. These are then integrated and evaluated for a new set of x values not equal to but inside the x values used to make the spline. I have overplotted this in the plot for $\eta(x)$ and the functions overlap.

4. Results

The Ω_x values indicate the relative density of a given component compared to the critical density of the universe. The critical density being that which corresponds to the universe begin flat. Looking at the figure we see radiation dominating from the start together with neutrinos. This continues for some time until the dark matter component starts to rise. We see that this starts before the baryon component. This is good since that makes it possible for dark matter to form structures that baryons can later fall into. The radiation and neutrinos die out, and at a later point vacuum energy shoots up. At the end we end up with approximately 70% vacuum energy(dark energy?), 25% dark matter, and 5% baryons. This is exactly as it should be since that is what they were set to be at the present. See figure 1.

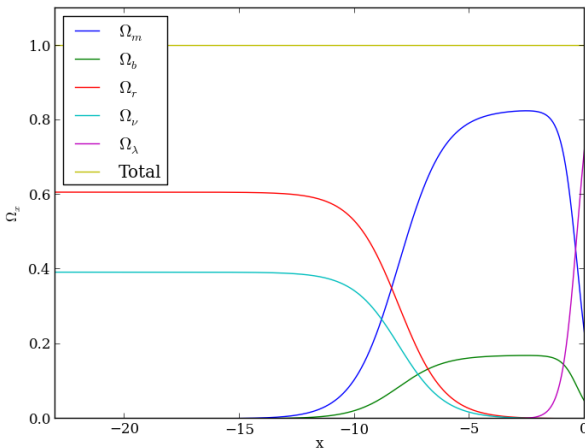


Fig. 1. The figure shows the evolution of the relative densities. As described they behave as they should.

Conformal time, denoted $\eta(x)$ measures the distance to the particle horizon for a given x value. A nice test of this is to insert

the x value of today. The answer should then be the radius of the observable universe. At the present this is measured to be approximately 14 billion parsecs (14Gpc)[2] The graph hits this value fairly well, indicating that everything is working properly so far. Note also that there are in fact two graphs in this figure, one which is calculated from the differential equation for η . And another one made by splining the first one and finding η values at arbitrary values between those of the first function.

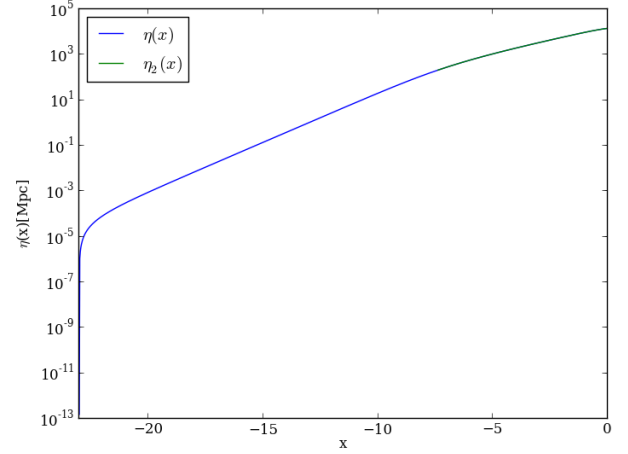


Fig. 2. The conformal time hits the value of today fairly accurately. The splined function hits the computed values so well that they overlap. Here $\eta(x)$ is the computed function, while $\eta_2(x)$ is the splined one.

The Hubble parameter H is depicted both as a function of x , and z . Whether this is good or not is hard to say for the first part. We can at least put some faith in its precision by the fact that it ends at $H_0 \approx 70 \text{ km s}^{-1} \text{ Mpc}^{-1}$, which is what H_0 was set to be.

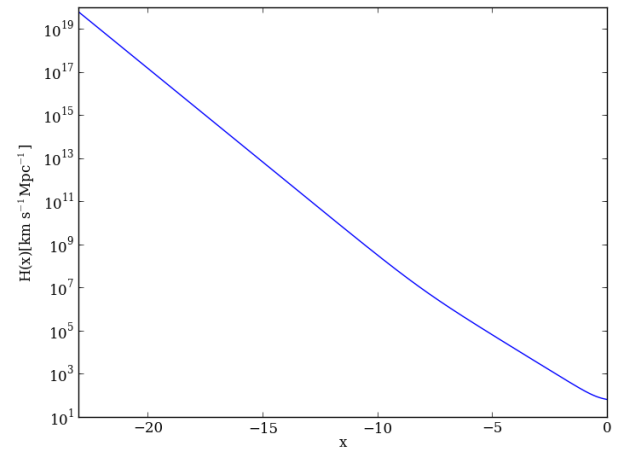


Fig. 3. The figure depicts the Hubble parameter as a function of x .

5. Conclusions

The background cosmology of the universe is done and everything has worked the way it was expected to. There are some values that I do not have an intuition for what should be. Namely the Hubble parameter at times before today. However, I expect

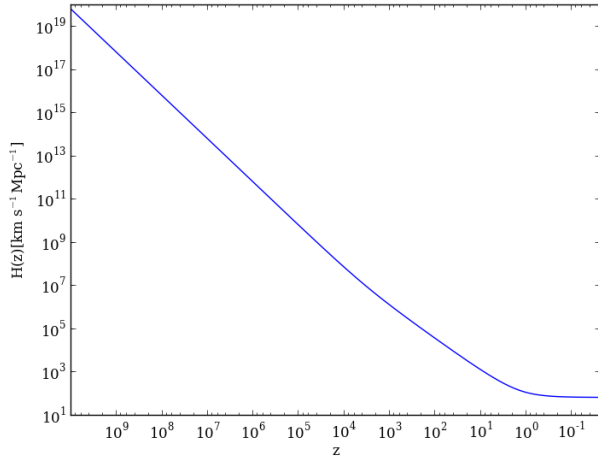


Fig. 4. The figure depicts the Hubble parameter as a function of redshift z . Note that z increases to the left, such that time goes towards the right.

these to be correct since everything else has given results generally considered to be correct, and the Hubble parameter is computed directly from these values. With this, the code is ready for the introduction of perturbations in part two.

6. References

- [1] P. Callin, astro-ph/0606683
- [2] I. Bars and J. Terning, Extra Dimensions in Space and Time, Springer 2010

7. Source code

The source code for the time_mod file is included for inspection. Note that the code makes use of several different files, one with various parameters, as well as the ODE solver, and the spline.

```
module time_mod
  use healpix_types
  use params
  use spline_1D_mod
  use ode_solver
  implicit none

  integer(i4b) :: n_t ! Number of x-values
  real(dp), allocatable, dimension(:) :: x_t ! Grid of relevant x-values
  real(dp), allocatable, dimension(:) :: a_t ! Grid of relevant a-values
  real(dp), allocatable, dimension(:) :: eta_t ! Grid of relevant eta-values

  integer(i4b) :: n_eta ! Number of eta grid points
  real(dp), allocatable, dimension(:) :: z_eta !Grid points for eta
  real(dp), allocatable, dimension(:) :: x_eta ! Grid points for eta
  real(dp), allocatable, dimension(:) :: a_eta ! Grid points for eta
  real(dp), allocatable, dimension(:) :: eta, eta2 ! Eta and eta'' at each grid point
  real(dp), allocatable, dimension(:) :: dydx

  real(dp) :: rho_m0 !matter density today
  real(dp) :: rho_b0 !baryong density today
  real(dp) :: rho_r0 !radiation density today
  real(dp) :: rho_nu0 !neutrino density today
  real(dp) :: rho_lambda0 !vacuum energy density today

  real(dp), allocatable, dimension(:) :: rho_m !matter density
  real(dp), allocatable, dimension(:) :: rho_b !baryong density
  real(dp), allocatable, dimension(:) :: rho_r !radiation density
  real(dp), allocatable, dimension(:) :: rho_nu !neutrino density
  real(dp), allocatable, dimension(:) :: rho_lambda !vacuum energy density

  real(dp), allocatable, dimension(:) :: Omega_mx !Relative densities
  real(dp), allocatable, dimension(:) :: Omega_bx
  real(dp), allocatable, dimension(:) :: Omega_rx
  real(dp), allocatable, dimension(:) :: Omega_nux
  real(dp), allocatable, dimension(:) :: Omega_lambdax

  real(dp), allocatable, dimension(:) :: H !Hubble constant as func of x
```

contains

```
subroutine initialize_time_mod
  implicit none

  integer(i4b) :: i, n, n1, n2
  real(dp) :: z_start_rec, z_end_rec, z_0, x_start_rec, x_end_rec, x_0
  real(dp) :: dx, x_eta1, x_eta2, a_init, h1, eta_init, a_end, rho_crit0, rho_crit
  real(dp) :: eps, hmin, yp1, ypn

  ! Define two epochs, 1) during and 2) after recombination.
  n1 = 200 ! Number of grid points during recombination
  n2 = 300 ! Number of grid points after recombination
  n_t = n1 + n2 ! Total number of grid points

  z_start_rec = 1630.4d0 ! Redshift of start of recombination
  z_end_rec = 614.2d0 ! Redshift of end of recombination
  z_0 = 0.d0 ! Redshift today
```

```

x_start_rec = -log(1.d0 + z_start_rec) ! x of start of recombination
x_end_rec   = -log(1.d0 + z_end_rec)   ! x of end of recombination
x_0         = 0.d0                     ! x today

n_eta       = 1000                     ! Number of eta grid points (for spline)
a_init      = 1.d-10                   ! Start value of a for eta evaluation
a_end       = 1.d0
x_eta1      = log(a_init)               ! Start value of x for eta evaluation
x_eta2      = 0.d0                     ! End value of x for eta evaluation
eta_init    = a_init/(H_0*sqrt(Omega_r+Omega_nu))

eps = 1.d-10
hmin = 0.d0

! Task: Fill in x and a grids ( These will be used in later milestones)
allocate(x_t(n_t))

do i = 0,n1-1 ! Fill interval during recombination
  x_t(i+1) = x_start_rec + i*(x_end_rec-x_start_rec)/(n1-1)
end do

do i = 1,n2 !Fill from end of recomb to today
  x_t(n1+i) = x_end_rec + (i)*(x_0-x_end_rec)/(n2)
end do

!write(*,*) x_t !print x_t to terminal

allocate(a_t(n_t+1))
a_t = exp(x_t) !fill the a grid using the x grid

!write(*,*) a_t !print a_t to terminal

!Allocate and fill a,x, and z arrays
allocate(a_eta(n_eta))
allocate(x_eta(n_eta))
allocate(z_eta(n_eta))

x_eta(1) = x_eta1
do i = 1,n_eta-1
  x_eta(i+1) = x_eta1 + i*(x_eta2-x_eta1)/(n_eta-1)
end do

a_eta = exp(x_eta)
z_eta = 1.d0/a_eta -1.d0

!write(*,*) z_eta
!write(*,*) size(z_eta)
!print *, "x"
!write(*,*) x_eta(1)
!write(*,*) x_eta(-1)
!print *, "a"
!write(*,*) a_eta(1)
!write(*,*) a_eta(-1)
!print *, "z"
!write(*,*) z_eta(1)
!write(*,*) z_eta(-1)

!Calculate the various densities for each scale factor
rho_crit0 = 3.d0*H_0**2.d0/(8.d0*pi*G_grav)
rho_m0    = Omega_m      *rho_crit0
rho_b0    = Omega_b      *rho_crit0

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```

rho_r0    = Omega_r      *rho_crit0
rho_nu0   = Omega_nu     *rho_crit0
rho_lambda0 = Omega_lambda*rho_crit0

allocate(rho_m(n_eta))
allocate(rho_b(n_eta))
allocate(rho_r(n_eta))
allocate(rho_nu(n_eta))
allocate(rho_lambda(n_eta))

allocate(Omega_mx(n_eta))
allocate(Omega_bx(n_eta))
allocate(Omega_rx(n_eta))
allocate(Omega_nux(n_eta))
allocate(Omega_lambdax(n_eta))
allocate(H(n_eta))

do i=1,n_eta+1
H(i) = get_H(x_eta(i))
Omega_mx(i) = Omega_m  *H_0**2.d0/H(i)**2.d0 *a_eta(i)**-3.d0
Omega_bx(i) = Omega_b  *H_0**2.d0/H(i)**2.d0 *a_eta(i)**-3.d0
Omega_rx(i) = Omega_r  *H_0**2.d0/H(i)**2.d0 *a_eta(i)**-4.d0
Omega_nux(i) = Omega_nu *H_0**2.d0/H(i)**2.d0 *a_eta(i)**-4.d0
Omega_lambdax(i) = Omega_lambda *H_0**2.d0/H(i)**2.d0
end do
!End of density calculations

allocate(eta(n_eta+1))
eta(1) = eta_init !Start value of eta

h1 = abs(1.d-2*(a_eta(1)-a_eta(2))) !Defines the steplength
allocate(dydx(1))

do i =2,n_eta+1
eta(i) =eta(i-1)
    call odeint(eta(i:i),a_eta(i-1) ,a_eta(i), eps, h1, hmin, eta_derivs, bsstep, output)
end do
!write(*,*) eta !check that eta gives reasonable values

!Spline eta and place the second derivative of
!this function in eta2
allocate(eta2(n_eta+1))
yp1 = 1.d30
ypn = 1.d30
call spline(a_eta, eta, yp1, ypn, eta2)

allocate(eta_t(n_t+1))
do i=1,n_t+1
    eta_t(i) = get_eta(x_t(i))
end do

end subroutine initialize_time_mod

!Begin Stuff needed to make odeint work

```

```

subroutine eta_derivs(a, eta, dydx) !Define the derivative d/da(eta)
  use healpix_types
  implicit none
  real(dp),          intent(in)  :: a
  real(dp), dimension(:), intent(in) :: eta
  real(dp), dimension(:), intent(out) :: dydx
real(dp) :: H_p
real(dp) :: x
  x = log(a)
H_p = get_H_p(x)
  dydx = c/(a*H_p)
end subroutine eta_derivs

subroutine output(x, y)
  use healpix_types
  implicit none
  real(dp),          intent(in)  :: x
  real(dp), dimension(:), intent(in) :: y
end subroutine output
!End Stuff needed to make odeint work

! Task: Write a function that computes H at given x
function get_H(x)
  implicit none

  real(dp), intent(in) :: x
  real(dp)          :: get_H
  real(dp)  :: a
  a = exp(x)
  get_H = H_0*sqrt((Omega_b+Omega_m)*a**-3.d0 + (Omega_r+Omega_nu)*a**-4.d0 + Omega_lambda)
end function get_H

! Task: Write a function that computes H' = a*H at given x
function get_H_p(x)
  implicit none

  real(dp), intent(in) :: x
  real(dp)          :: get_H_p
  real(dp)  :: a
  a = exp(x)
  get_H_p = a*get_H(x)
end function get_H_p

! Task: Write a function that computes dH'/dx at given x
function get_dH_p(x)
  implicit none

  real(dp), intent(in) :: x
  real(dp)          :: get_dH_p
  get_dH_p = H_0/2.d0*1/sqrt((Omega_m+Omega_b)*exp(-x)+Omega_r*exp(-2.d0*x) &
+ Omega_lambda*exp(2.d0*x)) * (-(Omega_m+Omega_b)*exp(-x)-2.d0*Omega_r*exp(-2.d0*x) &
+ 2.d0*Omega_lambda*exp(2.d0*x))
end function get_dH_p

! Task: Write a function that computes eta(x), using the previously precomputed splined function
function get_eta(x_in)
  implicit none

  real(dp), intent(in) :: x_in
  real(dp)          :: get_eta
  real(dp)  :: a_in
  a_in = exp(x_in)

```

```
    get_eta = splint(a_eta, eta, eta2, a_in)
end function get_eta

end module time_mod
```