

LECTURE 5: FUNDAMENTALS OF OBSERVATIONAL COSMOLOGY (I)

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In this lecture, we start covering basics of observational cosmology from analyzing the type Ia supernovae (SN Ia) data. As we discussed in the lecture, SN Ia are ‘standard candles’ of the Universe, which can be used to measure the luminosity distances. On the other hand, the redshifts of SN Ia can be independently measured by taking the spectra. Combining redshifts and distances, one can infer cosmological parameters. This is exactly what Adam Riess and Saul Perlmutter did in their Nobel-winning papers in 1998 [1, 2] (these papers have been cited over 10,000 times each)!

So the aim of this lecture and the next one is to reproduce the key results in the above-mentioned famous papers, *i.e.*, discover the cosmic acceleration using the SN Ia data!

The observed dataset of SN Ia is actually a list of redshifts z_i , the distance module μ_i , and a data covariance matrix \mathbf{C} . Note that,

$$(1) \quad \mu(z) \equiv m - M = 5 \log_{10} D_L(z) + 25$$

$$(2) \quad D_L = \begin{cases} (1+z) \frac{D_H}{\sqrt{\Omega_K}} \sinh \left(\sqrt{\Omega_K} \frac{D_C}{D_H} \right) & \Omega_K > 0; \\ (1+z) D_C & \Omega_K = 0; \\ (1+z) \frac{D_H}{\sqrt{-\Omega_K}} \sin \left(\sqrt{-\Omega_K} \frac{D_C}{D_H} \right) & \Omega_K < 0. \end{cases}$$

$$(3) \quad D_C = D_H \int_0^z \frac{dz'}{E(z')}$$

$$(4) \quad E(z) = \sqrt{\Omega_M(1+z)^3 + \Omega_\Lambda + (1 - \Omega_M - \Omega_\Lambda)(1+z)^2}$$

$$(5) \quad D_H = \frac{c}{H_0}$$

These equations are sufficient to allow for evaluating μ at arbitrary redshifts, given a set of $\{\Omega_M, \Omega_\Lambda, H_0\}$.

In class, we have written a Fortran code for this purpose, which is included in the Appendix. Note that the subroutine to perform the integral, `rombint`, is taken from Numerical Recipe [3], which is available online, <http://www.nrbook.com/a/bookfpdf.php>

REFERENCES

- [1] A. G. Riess *et al.* [Supernova Search Team], *Astron. J.* **116**, 1009 (1998) doi:10.1086/300499 [astro-ph/9805201].
- [2] S. Perlmutter *et al.* [Supernova Cosmology Project Collaboration], *Astrophys. J.* **517**, 565 (1999) doi:10.1086/307221 [astro-ph/9812133].
- [3] W. H. Press, S. A. Teukolsky, W. T. Vetterling and B. P. Flannery, “Numerical Recipes in FORTRAN: The Art of Scientific Computing.”

APPENDIX A. THE FORTRAN 90 CODE FOR COMPUTING THE LUMINOSITY DISTANCE

```
module precision
```

```
  integer , parameter :: dl = kind(1.d0)
end module precision
```

```
module cosmo
```

```
use precision
real(dl) :: om=0.3d0, ol=0.7d0, ok, H0=70.d0
real(dl) :: c =3.d5
real(dl) :: DH
end module cosmo
```

```
program test
```

```
use precision
use cosmo
implicit none
```

```
integer i
```

```
integer , parameter :: n=580
```

```
real(dl) :: z_dat(n), mu_dat(n), invcov(n,n)
```

```
character :: dummyc
```

```
real(dl) :: dummyr
```

```
real(dl) :: z
```

```
real(dl),external :: E2, Einv, DC, DA, mu
```

```
open(unit=50, file=’sn_z_mu_dmu_plow_union2.1.txt’)
```

```

do i=1, n
  read(50,*) dummyc, z_dat(i), mu_dat(i), dummyr, dummyr
  !write(*,*) z_dat(i), mu(i)
end do

close(50)

open(unit=50, file='sn_wmat_nosys_union2.1.txt ')

do i=1, n
  read(50,*) invcov(i,:)
  !write(*,*) invcov(i,i)
end do

close(50)

open(unit=50, file='test_mu.dat ')
do i=1, n
  write(50,*) z_dat(i), mu_dat(i), 1.d0/sqrt(invcov(i,i))
end do
close(50)

ok = 1.d0-om-ol

DH=c/H0

z=0.5d0
write(*, '(5e15.6) ') z, E2(z), Einv(z), DC(z), DA(z), mu(z)

end program test

function E2(z)
use precision
use cosmo
implicit none

real(dl) :: z, E2

```

```
E2 = om*(1.d0+z)**3.d0+ol+(1.d0-om-ol)*(1+z)**2.d0
```

```
end function E2
```

```
function Einv(z)
use precision
use cosmo
implicit none
```

```
real(dl) :: z, Einv
real(dl), external :: E2
```

```
Einv = 1.d0/sqrt(E2(z))
```

```
end function Einv
```

```
function DC(z)
use precision
use cosmo
implicit none
```

```
real(dl), external :: rombint, Einv
real(dl), parameter :: tol=1.d-4
real(dl) :: DC, z
```

```
DC = rombint(Einv,0.d0,z,tol)
DC = DC*DH
```

```
end function DC
```

```
function DA(z)
use precision
use cosmo
implicit none
```

```
real(dl) :: DA, z
real(dl), external :: DC
```

```
if(ok>0) then
```

```
DA = DH/(1.d0+z)/sqrt(ok)*sinh(sqrt(ok)*DC(z)/DH)
```

```
else if(ok<0) then
```

```
DA = DH/(1.d0+z)/sqrt(abs(ok))*sin(sqrt(abs(ok))*DC(z)/DH)
```

```
else
```

```
DA = DC(z)/(1.d0+z)
```

```
end if
```

```
end function DA
```

```
function Ldis(z)
```

```
use precision
```

```
use cosmo
```

```
implicit none
```

```
real(dl) :: Ldis, z
```

```
real(dl), external :: DA
```

```
Ldis = DA(z)*(1.d0+z)**2
```

```
end function Ldis
```

```
function mu(z)
```

```
use precision
```

```
use cosmo
```

```
implicit none
```

```
real(dl) :: mu, z
```

```
real(dl), external :: Ldis
```

```
mu=5.d0*log10(Ldis(z))+25.d0
```

```
end function mu
```

```

function rombint(f,a,b,tol)
  use Precision
  ! Rombint returns the integral from a to b of using Romberg integration.
  ! The method converges provided that f(x) is continuous in (a,b).
  ! f must be real(dl) and must be declared external in the calling
  ! routine. tol indicates the desired relative accuracy in the integral.
  !

  implicit none
  integer, parameter :: MAXITER=20
  integer, parameter :: MAXJ=5
  dimension g(MAXJ+1)
  real(dl) f
  external f
  real(dl) :: rombint
  real(dl), intent(in) :: a,b,tol
  integer :: nint, i, k, jmax, j
  real(dl) :: h, gmax, error, g, g0, g1, fourj

  h=0.5d0*(b-a)
  gmax=h*(f(a)+f(b))
  g(1)=gmax
  nint=1
  error=1.0d20
  i=0
10    i=i+1
      if (i.gt.MAXITER.or.(i.gt.5.and.abs(error).lt.tol)) &
        go to 40
  ! Calculate next trapezoidal rule approximation to integral.
  g0=0._dl
  do 20 k=1,nint
    g0=g0+f(a+(k+k-1)*h)
20    continue
  g0=0.5d0*g(1)+h*g0
  h=0.5d0*h
  nint=nint+nint
  jmax=min(i,MAXJ)
  fourj=1._dl
  do 30 j=1,jmax

```

```

! Use Richardson extrapolation.
    fourj=4._dl*fourj
    g1=g0+(g0-g(j))/(fourj-1._dl)
    g(j)=g0
    g0=g1
30  continue
    if (abs(g0).gt.tol) then
        error=1._dl-gmax/g0
    else
        error=gmax
    end if
    gmax=g0
    g(jmax+1)=g0
go to 10
40  rombint=g0
    if (i.gt.MAXITER.and.abs(error).gt.tol) then
        write(*,*) 'Warning: _Rombint_failed_to_converge; _'
        write (*,*) 'integral, _error, _tol:', rombint, error, tol
    end if

end function rombint

```