The Weekly Progress

Since Nov. 10, 2018

In the autumn semester, hopefully I'll record my progress every Thursday morning

Before Nov. 10, 2018

- 1. Understand why X-ray cause profound effects to ISM
- 2. Aquire the X-ray spectra emitted the SMBH in the center of Milky Way Galaxy several mys ago
- 3. Decide to focus on a cold dense cloud 8 kpc from the SMBH, thus fixing the flux
- 4. Figure out how KROME works, especially its X-ray module (details on https://github.com/slow_divePTG/X-ray-chemistry/tree/master)
- 5. Be able to calculate primary X-ray ionizations according to a X-ray spectra and the cross sections of certain species.
- 6. Learn about the cosmic ray ionization

Nov. 10, 2018

- 1. Trace the most important reactions using the Jacobian Matrix pd(i,j)
 In krome_oed.f90, the jex function calculate the Jacobian Matrix.
- 2. Test a simple model but consider a mixture of H/He rather than $\text{Lext}(H)_2/\text{Lext}(He)$. J_{X21}=0.1 for 1 mys then turn off the X-ray radiation. Plot the abundance of $\text{Lext}(CH)_4$ with time.

For this molecular cloud test we choose the osu_01_2007 network and the initial conditions proposed by Wakelam & Herbst (2008): a constant temperature of \text{T} = 10\text{ K}, \ce{H2} density of 10^4\text{ cm}^{-3}, cosmic rays ionization rate of 1.3 \times 10^{-17}\text{ s}^{-1}, and a visual extinction of 10. The initial conditions of the species are listed in Table 7 and correspond to the EA2 model of Wakelam & Herbst (2008), an high-metal environment observed in the diffuse cloud \zeta Ophiuchi

```
Tgas = 1d1 !gas temperature (K)
xH = 2d4 !Hydrogen density

!user commons for opacity and CR rate
call krome_set_user_av(1d1) !opacity Av (#)E11.3,
call krome_set_user_crate(1.3d-17) !CR rate (1/s)
call krome_set_user_gas_dust_ratio(7.57d11) !gas/dust
```

```
call krome_init()
  j21xs = 0d0
  call krome set J21xray(j21xs)
  print *,"J21 Xray = ",j21xs
 x(:) = 1.d-20
  !initial densities (model EA2 Wakelam+Herbst 2008)
 x(KROME_idx_H2) = 0.5d0 * xH !In this case we actually use
x(KROME_idx_H) = 0.5d0 * xH, which is not true
  x(KROME idx He) = 9d-2 * xH
 x(KROME_idx_N) = 7.6d-5 * xH
 x(KROME idx O) = 2.56d-4 * xH
 x(KROME_idx_Cj) = 1.2d-4 * xH
 x(KROME_idx_Sj) = 1.5d-5 * xH
 x(KROME idx Sij) = 1.7d-6 * xH
 x(KROME_idx_Fej) = 2d-7 * xH
  x(KROME idx Naj) = 2d-7 * xH
 x(KROME_idx_Mgj) = 2.4d-6 * xH
 x(KROME_idx_Clj) = 1.8d-7 * xH
 x(KROME idx Pj) = 1.17d-7 * xH
  x(KROME_idx_Fj) = 1.8d-8 * xH
  !calculate elctrons (neutral cloud)
  x(KROME_idx_e) = krome_get_electrons(x(:))
```

Nov. 15, 2018

1. (From Prof. Du) The secondary ionization rate of molecular hydrogen can be approximately considered as 2 hydrogen atoms.

Related readings:

X-ray chemistry in the envelopes around young stellar objects, Stäuber 2005

X-Ray--irradiated Molecular Gas. I. Physical Processes and General Results, Maloney 1996

According to Maloney 1996, the main process in the X-ray (secondary) ionization of \ce{H2} is:

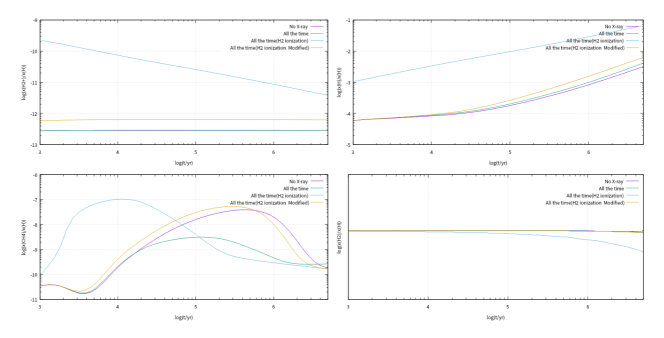
We can assume the rate coefficient is twice the ionization rates of \ce{H}, and thus include it in the network.

INCLUDE means when calculating the secondary ionization rates, we take the abundance of \ce{H}:

The modification of Fortran code in krome_subs.f90 is shown here

```
!H2 -> H2+ + E
```

Some results are:



Obviously the ionization of $\ensuremath{\mbox{\sc he}}\$ becomes more significant, and the profile of $\ensuremath{\mbox{\sc he}}\$ changes a lot. If we don't take n_t(\ce{H}) the effect is even more profound.

- (NOT AT PRESENT) For heavier atoms or other molecules, we can apply the approximation with the energy E fixed, neglecting the dependence of the cross sections on energy:
- 2. Species observants most interested in:

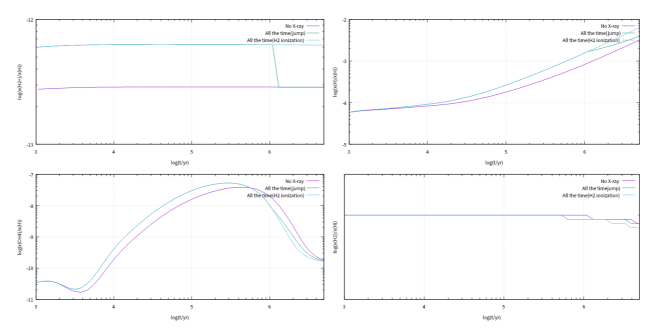
经常观测的分子包括\ce{CO、HCO+、N2H+、HCN}、各种碳氢化合物\ce{C2H、C3H3}等等。可以关注一下代表了电离度的电子丰度x_e,以及\ce{H3+、C+、OH+、H2O+}等离子的丰度;\ce{OH、CH、NH3、H2O}等也可以看看。还可以看更复杂一些的\ce{CH3CN、CH3OH}等

x_e is important because it decides the chemical composition and whether the heating process is sufficient

- 3. The simple \ce{H/He} model used before:
 - Abundance of \ce{CH4} too low
 - Not enough time to form \ce{H2}
 - Does not include \ce{H2} formation on grains

Therefore it is not reasonable enough, and I shall focus on the new \ce{H2/He} model to see what will happen if I turn off the X-ray after 1 mys.

Without $\ensuremath{\mbox{\sc Without }\mbox{\sc Without }\mbox{\sc$



Nov. 22, 2018

1. More details on secondary ionization

The secondary ionization rate for \ce{H} (Stäuber 2005, Maloney 1996,):

N_{sec}(E) is the number of secondary **ionizations** per unit energy produced by primary photoelectronsW(E) is the mean energy expended to produce an ion pair through rapid electron process

A BIG MISTAKE HERE LAST WEEK!

N_{sec}(E) is not the number of secondary electrons but the number of ion pairs produced by a secondary electron. **WE DON'T COUNT ELECTRONS HERE!**

More comments on the formula

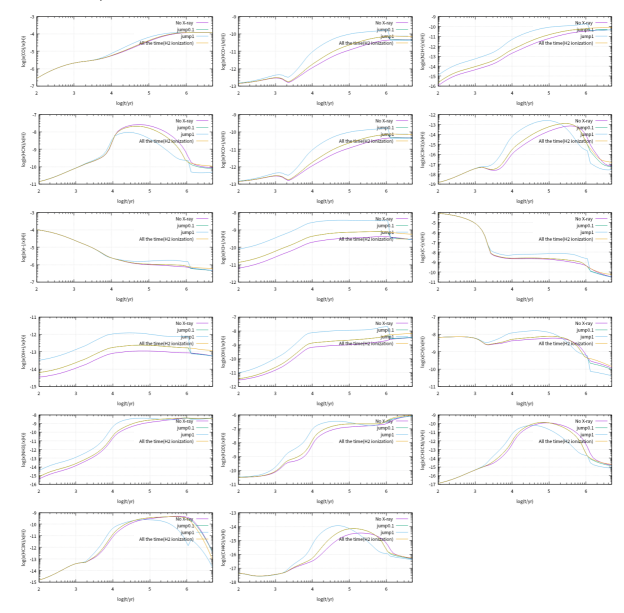
• W(E) is nearly independent of E if E>200\text{ eV} , which is easy to satisfied in our case

o N_{sec}(E,x_e) is dependent on x_e. When x_e>1\% , the energy in photoelectrons will be mainly lost to Coulomb Interaction with ambient thermal electrons. In our case, n_e/n_\ce{H}<10^{-4} . For weakly ionized gases, \approx40\% of the primary electron energy will be used for ionization (Krolik 1983, Shull 1985) , then for a secondary electron with energy 1\text{ keV} , the number of \ce{H} it can ionize is:

More precise calculation (Glassgold & Langer 1973, Dalgarno 1999, Glassgold 2012) shows that in $\ce{H2 + He}$ neutral gas, 47\% of energy goes to ionization, 47\% in $\ce{H2}$ and \ce{He} excitation, the rest goes for heating. N_{sec}(E)\approx27 and thus we can take W(E)\approx37\text{ eV}.

The formula I actually use

- The numbers of secondary ionization of \ce{H} and \ce{He} per primary ionization are taken from Shull 1985
- Can be rewritten like this:which average \phi over the X-ray spectrum F(E)
- 2. Test all the species listed last week



Nov. 29, 2018

 Recalculate the ionization rate using the primal formula rather than the simplified one and compare the results.

When we set n_\ce{H}=2\times10^4\text{ cm}^{-3},n_\ce{He}=2\times10^3\text{ cm}^{-3},J_{X21}=0.1,T=160\text{ K} , we get

Since we take $E_{\min}=2\text{keV}, E_{\max}=10\text{keV}$, and in our case, $F(E)\propto E^{-1.5}$, \sigma(E)\propto E^{-3}, \phi(E,x_e)\propto E, e^{-\tau(E)}\sim1, we have and

- 2. Why does KROME use a simplified formula to calculate the total ionization rate?
 - \zeta_p^\ce{H},\zeta_p^\ce{He} are stored in data-files rateH.dat and rateHe.dat (column density v.s. primary ionization rate)
 - To calculate \zeta^{(2)} the program needs to read x_e,n_\ce{H},n_\ce{He} for each calculation
 - By using the simplified formula, we may avoid integration

Here is part of the codes (krome subs.f90):

```
T = Tgas
   Trad = 2d4
   ncolH = num2col(n(idx H), n(:))
   ncolHe = num2col(n(idx_He),n(:)) !number density to column density
   logHe = log10(ncolHe) !take log
   logH = log10(ncolH)
   xe = min(n(idx_e) / (get_Hnuclei(n(:)) + 1d-40), 1d0)
   user xray H = fit anytab2D(user xray H anytabx(:), &
       user_xray_H_anytaby(:), &
        user_xray_H_anytabz(:,:), &
        user_xray_H_anytabxmul, &
        user_xray_H_anytabymul, &
        logH,logHe-logH)
        !linear interpolation to find the proper rate for the column
density
   phiH = .3908d0*(1e0-xe**.4092)**1.7592 * 327.832286034056d0 ! avoid
integration
   ratexH = 1d1**user xray H
   user_xray_He = fit_anytab2D(user_xray_He_anytabx(:), &
        user_xray_He_anytaby(:), &
       user_xray_He_anytabz(:,:), &
        user xray He anytabxmul, &
        user xray He anytabymul, &
        logH, logHe-logH)
   phiHe = .0554d0*(1d0-xe**.4614)**1.666 * 180.793458763612d0
```

Dec. 5, 2018

- 1. Try to repeat some results in former papers...but pretty difficult to do
 - Krolik 1983 did not consider the X-ray ionization of \ce{H2}

Dec. 12 review: Actually, \ce{H2} ionization is considered with an approximate model (Brown and Gould's 1970)

- Those who have considered the x-ray ionization of \ce{H2} usually have too complex physical problems and models, which means they are not easy to fully understand and repeat within hours
- The model we consider is simple and does not include heating/cooling process and totally ignore UV-photon process
- 2. Add molecular hydrogen formation process to the network (from react_primordial, primordial rates from several authors mostly based on Abel et al. 1997 + Grassi et al. 2014)

Dec. 12, 2018

1. Reread and try to repeat the results in Krolik 1983.

A Review on Krolik 1983

- Modify the codes
 - Initial abundances

- Longer time period (steady-state solutions) \Delta t=10^8\text{ yrs}
 (Strange tendency appears when \Delta t>10^8\text{ yrs}, abundances of some species changes rapidly)
- Set two branches for \ce{H2} ionization & dissociation
- Add \ce{H2} formation process on grains
- Step 1. Still use the power law spectrum,
 - It is the total ionization rate rather than the detail of the spectrum that is the most important to the chemistry
- 2. A new paper on the enhanced molecular ionization in the central molecular zone of our Galaxy

Molecular Ionization Rates and Ultracompact Dark Matter Minihalos, Silk, J. 2018, PRL

Dec. 20, 2018

- 1. Grain Process
 - It's important to find out how \ce{H2} molecules form on the grain surface as well as how KROME deal with the process
 - Theoretically, KROME uses
 where v_g is the gas thermal velocity, a_{ij} is the size of the grain, T_i is the dust temperature, \epsilon_j depends on the type of the dust and \alpha is the sticking coefficient
 - The rate is defined in the file krome_dust.f90

```
function krome_H2_dust(nndust,Tgas,Tdust,nH,H2_eps_f,myvgas)
   !H2 formed on dust (1/cm3/s)
   use krome constants
   use krome commons
   real*8::H2 dust, krome H2 dust, Tgas, Tdust(:)
   real*8::myvgas,H2_eps,nndust(:),nH,H2_eps_f
   integer::i
   H2 dust = 0.d0
   do i = 1,size(Tdust)
     H2_eps = H2_eps_f(Tgas, Tdust(i))
     H2 dust = H2 dust + 0.5d0 * nH * myvgas * nndust(i) &
          * krome_dust_asize2(i) &
          * pi * H2 eps * stick(Tgas, Tdust(i))
    end do
   krome_H2_dust = H2_dust
  end function krome H2 dust
```

as well as related functions. Again in krome_ode.f90 this rate is added to the total rate of \ce{H2} formation. The process is shown in kromeobj.py where

```
#add H2 formation on dust
if(self.useDustH2const):
   if("H"==specs[idnw].name): x += " - 2d0*nH2dust"
   if("H2"==specs[idnw].name): x += " + nH2dust"
```

2. Slightly modify the X-ray ionization

- In Krolik 1983, there are two branches for molecular Hydrogen ionization (See X-ray module in the <u>review</u>)
- Modify the krome_subs.f90

```
!H2 -> H+ + H + E
k(36) = rateEvaluateOnce(36)
!H2 -> H2+ + E
k(38) = rateEvaluateOnce(38)
!H -> H+ + E
k(4452) = small + ((ratexH &
    * (1d0+phiH) + n(idx He)&
    /(n(idx_H)+2*n(idx_H2)+1d-40) * ratexHe * phiH)&
    * J21xray)
!1keV cross section ratio
k(14) = k(14) + 150*k(4452) !C
k(22) = k(22) + 400*k(4452) !0
k(20) = k(20) + 280*k(4452)!N
k(19) = k(19) + 7*k(4452) !Mg
k(36) = k(36) + 2*k(4452)*0.2
k(38) = k(38) + 2*k(4452)*0.8
!Many molecular species are also included
!HE -> HE+ + E
    k(4453) = small + ((ratexHe &
        * (1d0+phiHe) + (n(idx_H)+2*n(idx_H2))&
        /(n(idx_He)+1d-40) * ratexH * phiHe)&
        * J21xray)
```

3. Repeat Krolik's results

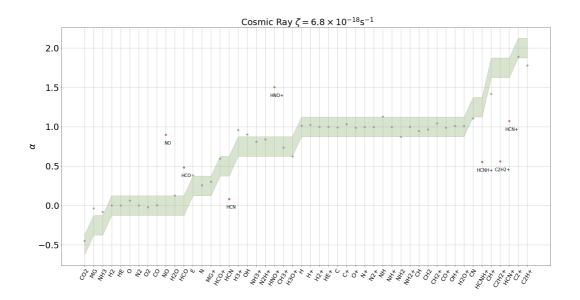
- Cosmic ray
 - well matched, except for several species such as \ce{HNO+} which has a larger \alpha.
 - The difference in \alpha between stronger and weaker ionization rates is larger.

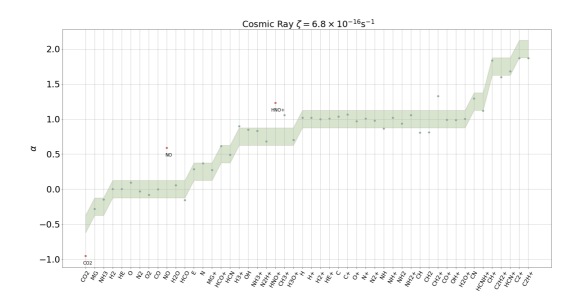
o X-ray

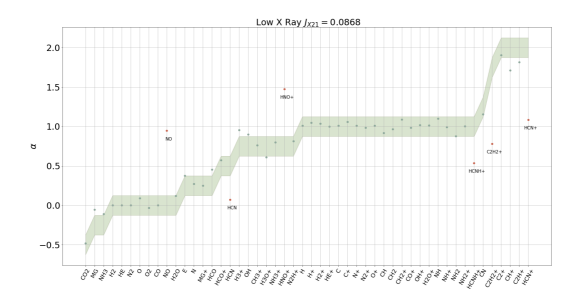
- For lower ionization rate (Jx21=0.08 and 8), the result is not far from Krolik's.
- For high ionization rate (Jx21=80), a strange problem occurs at about 20 million yrs, when the abundances of many species and reaction rates change by a few order of magnitude 'in a sudden'. Perhaps because of some calculation errors in the program?
- There seems not to be a state of chemical equilibrium, the abundance of molecular Hydrogen continue to go down.
- 4. Ameliorate the program to trace the important species

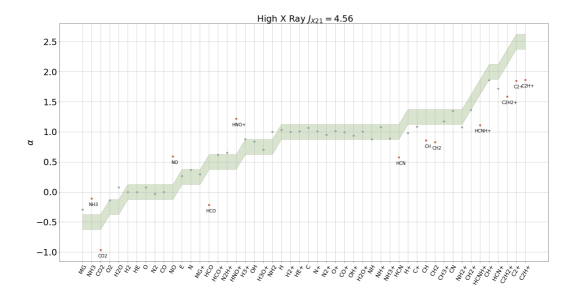
Dec. 27, 2018

1. Plot and compare the \alpha of Krolik's and the calculation results









Jan. 3, 2019

1. Find the internal time-step of KROME to see whether it may cause instability of the calculation.

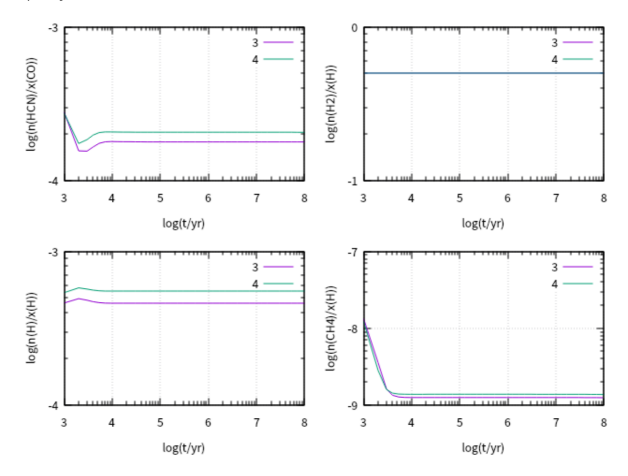
In the introduction of KROME, it goes like this

Note that the internal time-step of Krome is chosen according to the ODE system and the stiffness of the set of equations is evaluated and the user has no need to provide it, since it is automatically determined by the solver using the values of relative and absolute tolerance employed.

Do I have to find out how the solver would choose the value?

Jan. 16, 2019

- 1. Realize that there is a mistake in the rate coefficient of \ce{H}. The result is that the number of each atom is not conserved so there would not be any equilibrium state for the system. After fixing the bug, the system reaches equilibrium within as most several million years whatever the initial condition is, and we can re-plot the diagram.
 - In this case the step-length should be okay, and there seems no threshold of abundances. For safety I set a lower bound of 1d-99*nH for each species.
- 2. For high ionization rate situation, the result is far from Krolik's though a steady equilibrium is quickly reached.



Jan. 24, 2019

- 1. Recheck the X-ray ionization mechanism in the present code and compare the mechanism with those in present papers.
- 2. Move on.

Feb. 19, 2019

1. Consider electron cross section ratios in the calculation of secondary ionization of other species according to Maloney+ 1996

In krome subs.f90

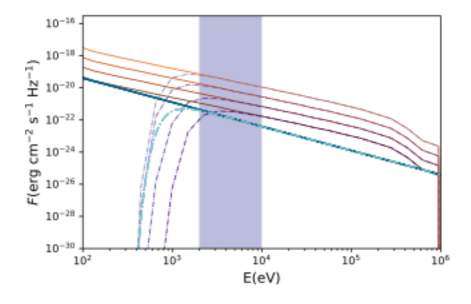
```
! 50eV cross section ratio
k(14) = k(14) + 3.1136*k(4452) !C
k(22) = k(22) + 1.5957*k(4452) !0
k(20) = k(20) + 2.0029*k(4452)!N
k(19) = k(19) + 13.1726*k(4452) !Mg
k(36) = k(36) + 2*k(4452)*0.05
k(38) = k(38) + 2*k(4452)*0.95
!C2 \rightarrow C + C
k(26) = k(26) + 3.1136*2*k(4452)
!CN \rightarrow C + N
k(30) = k(30) + (2.0029+3.1136)*k(4452)
!CO -> C + O
k(31) = k(31) + (3.1136+1.5957)*k(4452)*0.8
!CO -> CO+ + E
k(32) = k(32) + (3.1136+1.5957)*k(4452)*0.2
!NH -> N + H
k(45) = k(45) + (1+2.0029)*k(4452)
!NO -> N + O
k(46) = k(46) + (2.0029+1.5957)*k(4452)*0.8
!NO -> NO+ + E
k(47) = k(47) + (2.0029+1.5957)*k(4452)*0.2
102 -> 0 + 0
k(49) = k(49) + 1.5957*2*k(4452)*0.8
!02 -> 02+ + E
k(50) = k(50) + 1.5957*2*k(4452)*0.2
!H2O -> OH + H
k(72) = k(72) + (1.5957+2)*k(4452)
!HCN -> CN + H
k(75) = k(75) + (1+3.1136+2.0029)*k(4452)
!HCO -> CO + H
k(76) = k(76) + (1+3.1136+1.5957)*k(4452)*0.8
!HCO -> HCO+ + E
k(77) = k(77) + (1+3.1136+1.5957)*k(4452)*0.2
```

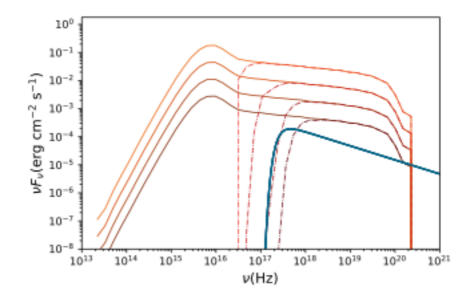
Mar. 2, 2019

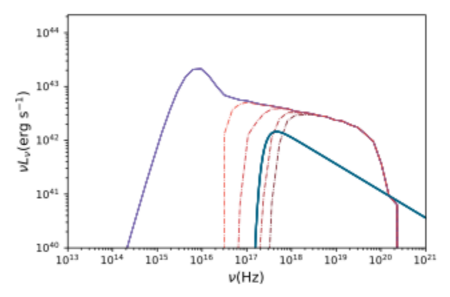
1. Include X-ray heating and \ce{H2} cooling processes to the code, still with bugs.

Mar. 14, 2019

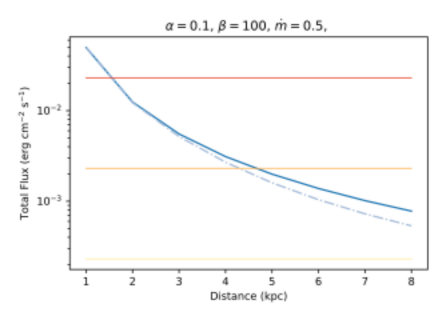
- Got Guangshuai's spectra, writing pipelines to plot and calculate X-ray spectra in different distances from Galactic center and different X-ray models (different \alpha,\ \beta,\ \dot{m})
- 2. Plot several figures
 - The standard (Latif+ 2015) X-ray spectra in comparison with Guangshuai's data (both with and without absorption)
 - J_{X21}=0.1,1,10
 - \alpha=0.1,\beta=100,\dot m=0.5 in difference distances (1, 2, 4, 8 kpc)



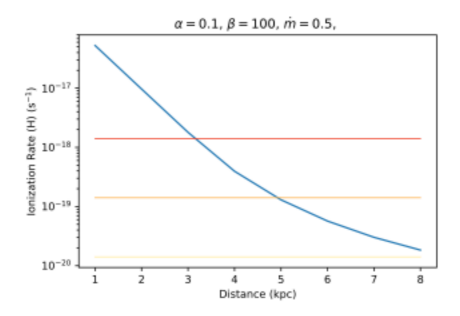


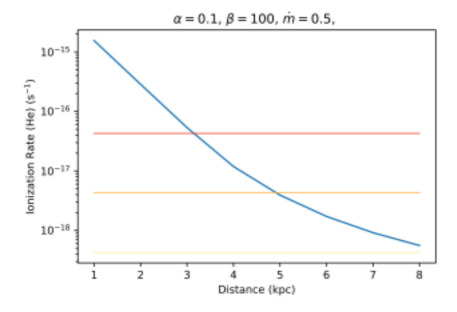


• The total flux in different distances as well as using the standard spectrum



• The comparison in ionization rates





To do list

- 1. Find more physical details of ISM closer to the galactic center with higher temperature and higher degree of ionization.
- 2. Figure out the extra process we need to consider (such as heating when the degree of ionization is high).