SemiCore Code Workshop

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Let us know if you have questions, ideas, want to discuss or have discovered any bugs!

Physics behind SemiCore code: decaying dark matter (DDM)

$$\psi^* \to \psi + l$$

where $\psi^*=$ mother (mom), $\psi=$ daughter (dau), l= relativistic light particle \Rightarrow escaped

Parameters: $V_k = \text{recoil velocity of } \psi$, $\tau = \text{half life}$

Since
$$\frac{V_k}{c} pprox \frac{\Delta m}{m_{\psi^*}} \Rightarrow m_{mom} pprox m_{dau}$$
 and

$$\langle \boldsymbol{v_{dau}} \rangle = \langle \boldsymbol{v_{mom}} + V_k \boldsymbol{\hat{n}} \rangle \Rightarrow \langle \boldsymbol{v_{dau}} \rangle = \langle \boldsymbol{v_{mom}} \rangle$$

 $(\hat{m{n}} \text{ is random})$, energies and angular momenta can be related by:

$$E_{k,dau} = E_{k,mom} + \frac{1}{2}V_k^2$$

$$L_{dau} = L_{mom}$$

General information

2 scripts written in Julia (1.4.2): **main.jl** (parameters & algorithm), **myFunctions.jl** (functions used by main.jl)

- No package required
- Physics parameters: $\tau, V_k, M_{vir}, c, \Delta_{vir}^{-1}$
- Default units for constants and parameters:
 - Dimensionless "little h" 2 : h = 0.6727
 - Mass: $10^{10} M_{\odot} \ h^{-1}$
 - Length: $kpc h^{-1}$
 - Velocity: $km \ s^{-1}$
- GPE, K.E., L, etc are in terms of quantity per unit mass

 $^{{}^{1}\}rho_{avg,r=r_{vir}} = \Delta_{vir}\rho_{c}$

 $^{^{2}}H_{0} = 100 \ h \ kms^{-1} \ Mpc^{-1}$

Where to adjust constants and parameters: main.jl

```
const Mo = 1.98847e30 # Solar mass [kg]
const kpc = 3.08567758128e19 # Kiloparsec [m]
const h = 0.6727 # "little h" [dimensionless]. Defined by H = 100 * h [km s-1 Mpc-1]
const H = 0.1 # Hubble constant at present [h km s-1 kpc-1]
const G = 43007.1 # Gravitational constant [1e-10 kpc Mo-1 (km s-1)-2]
const rho c = 3 * H ^ 2 / (8 * pi * G) # Critical density [Mo kpc-3 h2]
tol ellipseGuess = 0.00001 / 100 # Tolerance for bisection method in ellipseRadii() [in the unit of sh
orderOfpolynomial = 14 # Order of polynomial for fitting res(x), 14 is recommended
M vir = 0.517 # [1e10 Mo h-1]
c = 21.6 # Concentration parameter of the NFW halo
rho avg = 103.4 * rho c # Average density within virial radius
firstShellThickness = 1e-4 # If interested range of radius starts from 1e-n, use 1e-(n-2) for good acc
shellThicknessFactor = 1.0032 # Thickness of each consecutive shell grows exponentially by this rate
extend factor = 4 # Maximum halo radius at initialization is set as R vir * extend factor. 4 is recomm
v k = 40 # Recoil velocity of new born daughter particles [km s-1]
t end = 14 # Ending time [Gvr]
numOfSteps = 15 # Total number of time evolution intervals
```

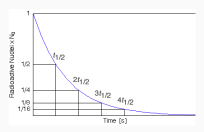
Time discretization

Mass m of mothers decays as: $m_{mom} = m_0 e^{-ln2\frac{t}{\tau}}$

Previous method: constant $\Delta t = \frac{t_end}{numOfSteps} \left(= \frac{Age\ of\ universe}{No.\ of\ time\ steps} \right)$

Now: set Δt for each step such that $\Delta m_{mom} = constant$

 More computationally cost-effective: calculate more frequently in the early phase during which more mass decays



Previous method: equal intervals in time (physicsabout.com)

Shell scheme

Spherical symmetry is assumed. The dark matter halo is partitioned into spherical shells, one enclosing another

Therefore, the main objects being manipulated in the code are arrays of shells named **X**shells_content:

- Particle type X: M (mom), D (dau), T (mom + dau)
- content: radii, mass, enclosedMass, GPE, L, totalE, etc

The ith item in any of such arrays corresponds to the **content** for that particle type \mathbf{X} in the ith shell. For example:

- $Dshells_mass[i]$: mass of daus in the ith shell
- ullet $Mshells_L[i]$: Angular momentum of moms in the ith shell

Initialization: Tshells_radii

At initialization, fixed positions and thicknesses of all shells are defined. $Tshells_radii$ stores all info specifying each shell and remains unchanged throughout the running of the code:

- \bullet $Tshells_radii[i, 1]$: inner radius of the ith shell
- $Tshells_radii[i, 2]$: outer radius
- $Tshells_radii[i, 3]$: shell radius = (inner + outer) / 2. It is generally used to represent the position of the ith shell when e.g. calculating GPE

For efficiency, thickness (outer - inner) of the ith shell is $\alpha \beta^{i-1}$

- $\bullet \ \alpha = firstShellThickness \ \text{and} \ \beta = shellThicknessFactor$
- \bullet Adjust α and β for more or fewer numOfShells

Initialization: Tshells_radii

All together, the shells span the halo space from r=0 (center) to $r=4R_{vir}$

 $Tshells_radii$ being unchanged means that masses falling outside of $4R_{vir}$ are neglected

• Discrepancy is negligible

Why $4R_{vir}$ (or larger; tunable via $extend_factor$) but not R_{vir} ?

- To trace mass contributions to the halo's center from more daughter particles
- More accurate GPE calculation

Initializing the $Xshells_mass$ arrays:

- NFW mass profile (analytic) $\Rightarrow Mshell_mass[i]$
- No daughter at initialization $\Rightarrow Dshell_mass[i] = 0$
- $\bullet \ Tshells_mass[i] = Mshells_mass[i] + Dshells_mass[i]$

Together with $Tshells_radii$, all useful quantities can be inferred

Therefore, the job of the code is to calculate the right amount of masses of moms and daus in each shell at the end of each decay time step

GPE: Tshells_GPE

At initialization:

• $\Phi_{NFW}(r)$ is used (analytic)

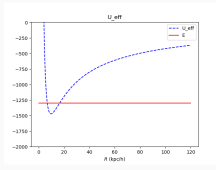
In other time steps:

•
$$\Phi(r) = -G\left[\frac{1}{r}\int_0^r dM(r') + \int_r^R \frac{dM(r')}{r'}\right]$$

- ullet Numerically, $\int \to \sum$ and $dM(r') \to Tshells_mass[i]$
- Shell radii are used, i.e. $r' = Tshells_radii[i, 3]$
- Mass in each shell is thought to concentrate at just beyond the shell radius
- ullet values at all shell radii are calculated and stored
 - $\Phi(r)$ at arbitrary r can be further obtained by interpolating these values (required in next step)

Roots finding: r_{max} , r_{min} of a daughter's orbit

Assume that every mother orbits circularly at the bottom of the effective potential $\Phi_{eff}(r)=\Phi(r)+\frac{L^2}{2r^2}$



Solve $\Phi_{eff}(r) = E_{k,dau}$ for r_{max} , r_{min} using bisection method

- ullet Increase accuracy by lowering error tolerance $tol_ellipseGuess$
- Escaped daughters $(E_{k,dau} \ge 0)$ are discarded

g-function: mass contribution from daughters

Recall: goal is to obtain resulting mass in each shell

• Calculate daughters' contribution to each shell via **g-function**

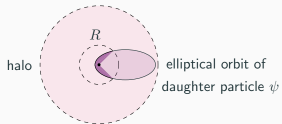
g-function: mass contribution from daughters

Let ψ be a daughter particle in an elliptical orbit and M(R) is its mass contribution to the region enclosed by r=R

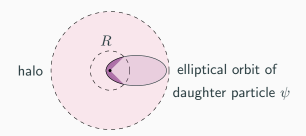
$$M(R) = \sum m_{\psi} g_{\psi}(R)$$

The **g-function** $g_{\psi}(R)$ is the ratio defined by $\Delta t_{\psi}(R)$ [the duration in which ψ orbits within R] to T_{ψ} [ψ 's orbital period]:

$$g_{\psi}(R) = \Delta t_{\psi}(R)/T_{\psi}$$



g-function: mass contribution from daughters



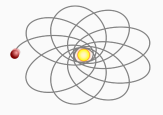
$$g_{\psi}(R) = \Delta t_{\psi}(R)/T_{\psi}$$

- $r_{max} \leq R \Rightarrow$ entire ellipse inside $R \Rightarrow g_{\psi}(R) = 1$
- $r_{min} > R \Rightarrow$ entire ellipse outside $R \Rightarrow g_{\psi}(R) = 0$
- $r_{min} \le R < r_{max} \Rightarrow g_{\psi}(R) = \Delta t_{\psi}(R)/T_{\psi}$

g-function: the accurate picture and its general formula

However, daughters do NOT travel in closed ellipses

Not a point mass field \Rightarrow not exactly a conic section



(Wikipedia)

 $g_{\psi}(R)$ must be evaluated via this **general formula**:

$$\Delta t_{\psi}(R)/T_{\psi} = \int_{r_{min}}^{R} \frac{dr}{\sqrt{E - \Phi_{eff}(r)}} / \int_{r_{min}}^{r_{max}} \frac{dr}{\sqrt{E - \Phi_{eff}(r)}}$$

g-function: integration method

Define the normalization: $x = \frac{r - r_{min}}{r_{max} - r_{min}}$

$$\begin{array}{l} \text{Modify the integrand} \ \frac{1}{\sqrt{E-\Phi_{eff}(r)}} = \\ \frac{1}{\sqrt{E-\Phi_{eff}(r)}} + \frac{1}{\sqrt{\Phi'_{eff}(x_{max})(x_{max}-x)}} + \frac{1}{\sqrt{\Phi'_{eff}(x_{min})(x-x_{min})}} - \\ \frac{1}{\sqrt{\Phi'_{eff}(x_{max})(x_{max}-x)}} - \frac{1}{\sqrt{\Phi'_{eff}(x_{min})(x-x_{min})}} \end{array}$$

$$\begin{array}{l} \text{Define } res(x) = \\ \frac{1}{\sqrt{E - \Phi_{eff}(r)}} - \frac{1}{\sqrt{\Phi'_{eff}(x_{max})(x_{max} - x)}} - \frac{1}{\sqrt{\Phi'_{eff}(x_{min})(x - x_{min})}} \end{array}$$

Employ polynomial approximation:

$$res(x) = a + bx + cx^{2} + dx^{3} + \dots + C_{N-1}x^{N-1}$$

How: take N points to set up a system of linear equations and solve for coefficients by Gaussian elimination (order tunable via orderOfpolynomial)

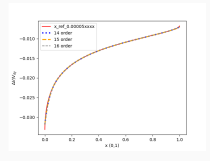
g-function: code for polynomial approximation of res(x)

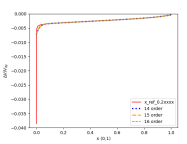
 $N = \mathsf{number} \; \mathsf{of} \; \mathsf{points} = orderOfpolynomial + 1$

```
Use the set of points x=[x_1,x_2,...,x_N] where x_1=0.001,\ x_2=\frac{0.998}{N-1},\ x_3=\frac{2(0.998)}{N-1},...,\ x_N=0.999
```

```
Number of point = orderOfpolynomial + 1
dx = 0.998 / (Number of point - 1)
equation matrix = zeros(Number of point, Number of point + 1)
 or i in 1:Number of point
   x = 0.001 + (i - 1) * dx
    for j in 1:Number_of point
        equation matrix[i,j] = (x-x a)^{(j-1)}
    equation matrix[i,end] = res(x)
soln = zeros(Number of point)
gaussian elimination!(equation matrix)
gauss jordan elimination!(equation matrix)
soln = back substitution(equation matrix)
```

g-function: plots of res(x) and polynomial approximations





- 14-th order polynomial is smooth and good enough
- Higher order (e.g. 21-st order) approximations give fluctuating curves, not recommended

g-function: integration method

Recall: modified integrand

$$\frac{1}{\sqrt{E-\Phi_{eff}(r)}} = res(x) - \frac{1}{\sqrt{\Phi'_{eff}(x_{max})(x_{max}-x)}} - \frac{1}{\sqrt{\Phi'_{eff}(x_{min})(x-x_{min})}}$$

All 3 terms can now be directly integrated. In particular:

$$\int res(x)dx = ax + bx^2/2 + cx^3/3 + dx^4/4 + \dots + C_{N-1}x^N/N$$

Adiabatic expansion

After distributing the mass of newborn daughters, the halo's enclosed mass M(R) changes from $M_{t_{i-1}}$ to M_{t_i} . Consider a test particle orbiting circularly at R and the conservation of its angular momentum L:

$$v^{2} = \frac{GM}{R}, \ L = mRv \Rightarrow \frac{L^{2}}{Gm^{2}} = MR$$
$$\Rightarrow M_{t_{i}}R_{t_{i}} = M_{t_{i-1}}R_{t_{i-1}} \Rightarrow R_{t_{i}} = \frac{M_{t_{i-1}}(R_{t_{i-1}})}{M_{t_{i}}(R_{t_{i-1}})}R_{t_{i-1}}$$

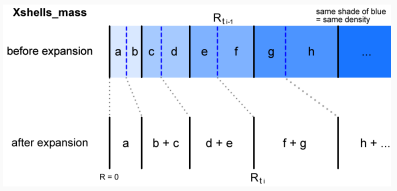
i.e. orbiting radius of the test particle evolves from ${\cal R}_{t_{i-1}}$ to ${\cal R}_{t_i}$

This adiabatic expansion of orbits is applied to both mothers $(Mshells_mass)$ and daughters $(Dshells_mass)$ which includes new daughters just born in the current time step)

Adiabatic expansion: implementation

```
# Adiabatic expansions (applied to both mothers and daughters)
Mshells_mass = adiabaticExpansion(Tshells_radii, Mshells_mass, Tshells_enclosedMass, Tshells_enclosedMass_updated)
Dshells_mass = adiabaticExpansion(Tshells_radii, Dshells_mass, Tshells_enclosedMass, Tshells_enclosedMass_updated)
```

What adiabaticExpansion() does:



- Positions and dimensions of shells stay the same (boundaries in black)
- Given R_{t_i} , $R_{t_{i-1}}$ is retrieved by interpolation
- Assuming uniform density within each shell, obtain mass to be moved

Adiabatic expansion: more details on implementation

A more detailed explanation:

- 1. Recall: positions and dimensions of shells ($Tshells_radii$) stay the same before and after expansion (boundaries in black). Goal is to figure out the amount of mass to be put in each shell after expansion
- 2. Using the set of boundaries (outer radii) as $\{R_{t_{i-1}}\}$, we use the relation $R_{t_i} = \frac{M_{t_{i-1}}(R_{t_{i-1}})}{M_{t_i}(R_{t_{i-1}})}R_{t_{i-1}}$ to compute the corresponding set of expanded radii $\{R_{t_i}\}$
- 3. Given an outer radius R_{t_i} of the expanded halo, retrieve the corresponding $R_{t_{i-1}}$ from the 2 sets of radii by interpolation (note: R=0 does not expand)
- 4. Assuming uniform density within each shell before expansion, we can deduce the right parts of mass to be put in each shell after expansion

Summary: algorithm of dark matter only SemiCore code

- Initialize NFW profile (mothers only + NO daughters)
- Calculate GPE
- For loop starts (t = 0)
 - ullet Mothers decay into daughters, solve for r_{max} and r_{min}
 - Calculate g-function to distribute daughters
 - Obtain updated enclosed mass profile
 - · Adiabatic expansion of mothers and daughters
 - Update GPE
- For loop ends ($t = t_end$)

Code explained: for loop

```
# Calculate L and total E of mother from the total mass distribution

Mshells_L = L(Tshells_radii, Tshells_enclosedMass, 6)

Mshells_LotalE_afterDecay = totalE_afterDecay(Tshells_radii, Tshells_GPE, Mshells_L, v_k)

# Solve equation to get ellipse

Mshells_ellipseRadii = ellipseRadii(Mshells_L, Mshells_totalE_afterDecay, Tshells_radii, Tshells_GPE, tol_ellipseGuess)

# Decay the mothers in the shells, distribute the new daughters

Mshells_mass, Dshells_decayedMass = updateShellsMass(Tshells_radii, Mshells_ellipseRadii, Mshells_mass, p_undecayed,Mshells_mass, Dshells_decayedMass = updateShellsMass(Tshells_radii, Mshells_ellipseRadii, Mshells_mass, p_undecayed,Mshells_mass, p_undecayed,Mshells_mass, p_undecayed,Mshells_mass(Tshells_mass, Dshells_decayedMass = updateShellsMass(Tshells_radii, Mshells_ellipseRadii, Mshells_mass, p_undecayed,Mshells_mass(Tshells_mass, Dshells_decayedMass, p_undecayed,Mshells_mass, p_undecayed,Mshells_mass(Tshells_mass, Dshells_decayedMass, p_undecayed,Mshells_mass, p_undecayed,Mshells_mass
```

- 1. Obtain angular momenta L of mothers
- 2. Calculate total energy of newborn daughters by

$$E_{k,dau} = E_{k,mom} + \frac{1}{2}m_{dau}V_k^2$$

- 3. Solve for r_{max} and r_{min}
- 4. Decaying of mothers and distribution of daughters
 - $Mshells_mass$: decay by the ratio $=e^{\frac{-ln(2)}{\tau}t_i}/e^{\frac{-ln(2)}{\tau}t_{i-1}}$
 - Dshells_decayedMass: distributed newborn daughters using g-function

Code explained: for loop

```
Dshells_mass = Dshells_mass + Dshells_decayedMass # Combine new daughters with old daughters

Tshells_mass_updated = Mshells_mass + Dshells_mass # Total distribution

Tshells_enclosedMass_updated = enclosedMass(Tshells_madi), Tshells_mass_updated) # Total enclosed mass

Tshells_GPE_updated = GPE(Tshells_madii, Tshells_mass_updated, Tshells_enclosedMass_updated) # GPE for this is
```

- 1. Add up accumulated (from previous steps) and newborn daus
- 2. Add up mothers and daughters
- 3. Obtain updated enclosed mass array
- 4. Obtain GPE (for record keeping, not useful)

- 1. Adiabatic expansion of mothers
- 2. Adiabatic expansion of daughters
- 3. Add up mothers and daughters
- 4. Obtain updated enclosed mass array
- 5. Obtain updated GPE to be used in next loop

Output

The code creates a **folder** ("dmOnly") under the same directory as itself, and a **subfolder** named according to the parameters of that particular run (" M_{vir} - V_k - τ -numOfShells-numOfSteps")

Inside the subfolder:

- "params.txt" stores ALL constants and parameters necessary for reproducing the same result
- 2. "stepResults.txt" stores time used in each time step
- 3. Various files from each time step labeled accordingly
- 4. A folder ("results") containing result files from the end step

Output



Output: result files

A typical result file (e.g. "T_result.txt"):

T_result.txt - Notepad							
File Edit Format View Help							
0.0 9.833987177753629e-5 4.9169935888768145e-5 1.8142100166245178e-14 0.004554178388645854 1.8142100166245178e-14 0.004554178388645854 0.0028167547849814777							
3,833987177753628e-5	0,0001986465409906233	0.00014849320638407978	1.2253445570579064e-13	0.0042471511436860876	1,406765558720358e-13	0.0042844008460740015	0.00551874642213
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1,0029910020349033273	0.003149855567779134 0.003311192550912253	0.0032305240593456935	2.2663516154777238e-11 2.4985427323586124e-11	0.001208908444457603 0.0011806108480026196	2.1717756610436886e-10	0.0014681657540418888 0.0014281474319556688	0.05122025052002
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),004167588773220607	0.004349280420462556	0.0042584345968415815	4.179343974889498e-11	0.0010092469584741704	4.238268300989786e-10	0.0012298371489923511	0.06473745132508
),004349280420462555	0.004534605900649342	0.004441943160555949	4.5239206247342504e-11	0.0009843759412859697	4.690660363463211e-10	0.001200954993849061	0.06669868920507
),004534605900649343	0.004723637890439866	0.0046291218955446045	4,8738369172286264e-11	0.0009573444939632106	5,178044055186073e-10	0.0011728631432417558	0.06866171567971
).004723637890439866	0.0049164505200262	0.004820044205233033	5.288341930053881e-11	0.0009393214396204417	5.706878248191461e-10	0.0011464496337134408	0.07065510508873
),0049164505200262	0.0051131194022042605	0.00501478496111523	5,742667845188189e-11	0.0009238636816175711	6,28114503271028e-10	0.0011217405215164836	0.07268528921374
).00511311940220426	0.005313721662025882	0.005213420532115071	6.219112321318264e-11	0.000907577288502409	6.903056264842107e-10	0.001098389564712284	0.07474659007747
),0053137216620258824	0.005518335967043936	0.00541602881453491	6.663091581943427e-11	0.0008833164963308757	7.56936542303645e-10	0.0010753416069040362	0.07680610586167
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).0059399232811031336	0.006157061618502733	0.0060484924498029335	8.349682603206445e-11	0.0008363396159290906	9.897172363978459e-10	0.0010122831892153117	0.08314552814027
).006157061618502734	0.006378542722650325	0.00626780217057653	8.971694095615585e-11	0.0008204500849275294	1.0794341773540017e-9	0.0009929860543481607	0.08531147039811
).006378542722650323	0.006604453448880866	0.006491498085765594	9.553723296305864e-11	0.0007985318572583536	1.1749714103170604e-9	0.0009737064341773474	0.08747124084305
SARUSSSFFESTFFURGROUD L	0.000834883380838033	0.006719667919258445	1 02410827435005184 10	0 0007832474221078700	1 27730123775215554 0	0.000025025250377603	0.05981.052559080.0

Output: result files

Many (= numOfShells) rows and 8 columns (from left to right):

- 1. Inner radius $[kpc \ h^{-1}]$
- 2. Outer radius $[kpc \ h^{-1}]$
- 3. Shell radius = (inner + outer) / 2 [$kpc h^{-1}$]
- 4. Shell mass $[10^{10}~M_{\odot}~h^{-1}]$
- 5. Shell density = shell mass / shell volume $[10^{10}~M_{\odot}~kpc^{-3}~h^2]$
- 6. Enclosed mass (enclosed by outer radius) $[10^{10}~M_{\odot}~h^{-1}]$
- 7. Average density = enclosed mass / enclosed volume [$10^{10}~M_{\odot}~kpc^{-3}~h^2$]
- 8. Circular velocity $[km \ s^{-1}]$

Output: use of shell radii

Different quantities may be defined using **different shell radii** (inner, outer or shell). For example, the circular velocity is defined

by:
$$shells_Vcir[i] = \sqrt{G \frac{shells_enclosedMass[i]}{Tshells_radii[i,\ 2]}}$$

For consistency, plot:

- $\bullet \ shells_Vcir[i] \ \mathsf{vs.} \ Tshells_radii[i,2] \ \mathsf{(outer\ radius)} \\$
- \bullet $shells_mass$ vs. $Tshells_radii[i,3]$ (shell radius)
- etc

Adding particle types

For example, baryons such as stars ($Sshells_mass$) and gas ($Gshells_mass$) may be easily added to the halo sharing with the dark matter particles the same scheme of shells ($Tshells_radii$)

Particle types can then be freely programmed to interact and evolve in various ways

It is reminded that masses from the additional particle types must be added to the total mass profile, such as $Tshells_enclosedMass$ for $Tshells_GPE$ calculations

Acknowledgements

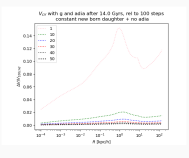
This version of the SemiCore code was developed under the close supervision of all members of the DDM group at the Department of Physics at the Chinese University of Hong Kong.

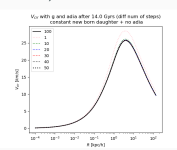
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Appendix: convergence on numOfSteps (without adiabatic expansion)

Without adiabatic expansion

$$(M_{vir} = 0.517, c = 21.6, \Delta_{vir} = 103.4)$$
:





• With kick-off effect only: $numOfSteps > 10 \Rightarrow good$ convergence

Appendix: convergence on numOfSteps (with adiabatic expansion)

Convergence cannot be tested with adiabatic expansion turned on

- $numOfSteps \uparrow \Rightarrow \Delta t \downarrow$
- Physically, there is insufficient time for particles to reach expanded radii, especially for particles in the outer region
- In our code, all particles undergo adiabatic expansion completely

Recommended numOfSteps = 15