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[§] = 10⁵ g/cm³

A User's Manual for grpfs/grafs

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Abstract

grpfs/grafs numerically solves the free boundary problem that describes internal structure of general relativistic polytropic/adiabatic fluid spheres with a non-zero cosmological constant.

This document serves as an introductory manual for the grpfs/grafs package only. Neither plotting programs used for creating graphical presentation, nor the package noutils used for postprocessing the output are covered—in this point the user is referred to the documentation cited hereafter.

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1 Introduction

grpfs/grafs is a command-line driven program that calculates the internal structure of general relativistic polytropic/adiabatic fluid spheres (from which the program name has been derived). Based on Tooper's papers [Tooper, 1964, Tooper, 1965], it implements the effect of a non-zero cosmological constant as a contribution to the physical facet and more reliable and effective numerical methods as a contribution to the computational facet of the problem.

Whether the program deals with polytropic or adiabatic case, depends on the way it is invoked. When invoked as grpfs, the polytropic case will be solved, when invoked as grafs, the adiabatic case will be solved. Because both cases are similar from the computational point of view, we will not distinguish between them if not stated otherwise, calling the program simply grpfs throughout the manual. The same applies for whole the package as well. The differences between grpfs and grafs are listed in Appendix A.

Together with standard Unix¹ utilities like *awk* or *sed* and with its redirection/piping facility, the program is pretty flexible. For these purposes one can use the noutils package [Ref].

Section 2 gives some information on how the program can be obtained, no warranty and installation guidelines follow in Sections 3 and 4. Although it is beyond the scope of this manual to deal with the underlying principles of the problem, one can find it useful to get some idea about the mathematics and numerics the program has to cope with. This is also necessary for fully appreciating all the features and helpful when trying to fix problems. Therefore, the basic underlying math is briefly discussed in Section 5. The description of usage includes Section 6 on basic tasks and Section 7 on dealing with advanced features. Section 8 gives an idea how to customize the program to match the user's needs; Section 9 mentions some potentially buggy aspects. Finally, Appendix A lists the differences between grpfs and grafs, and Appendix B presents the full program code.

2 Obtaining grpfs

The package grpfs can be obtained from the Computational Physics Laboratory homepage http://uf.fpf.slu.cz/cpl/. Since the source code contain routines from [Press et al., 1997] (see Appendix B for more info) that are subject to copyright to Numerical Recipes Software that forbids to distribute them on a free basis, the site is password-protected. In order to obtain the password, please contact the author at Email address Stanislav.Hledik@fpf.slu.cz.

^{1.} Unix is a registered trademark of Unix Systems Laboratories.

The source code can be modified and distributed without prior author's permission, with the exception of the module grp_node.c (containing routines from [Press et al., 1997]) that can solely be used for studying purposes.

The preferred platform for the noutils package is a GNU Unix system. However, this package is fully portable on most Unix platforms.

3 No Warranty

The software described in this manual has no warranty, it is provided 'as is.' It is your responsibility to validate the behavior of the program and its ability to perform the task required using the source code provided.

4 Installation

All sources are distributed packed in the tarball grpfs.tar.gz. Download that file and save it in a working directory of your choice. Once downloaded, cd to this directory and use the command (the leading \$ represents the shell prompt hereafter)

\$ gzip -dc grpfs | tar xvvf - ; cd grpfs
(On a GNU Unix system, the simpler command

\$ tar xzvvf grpfs ; cd grpfs

also works.) This unpacks all files into the directory grpfs and moves you to it. Then edit subsection 'Installation' of the 'Customization section' in the file Makefile to match your needs. Unless you are a superuser, you will probably want to install into your ~/bin directory; in such case there is no need to customize Makefile at all. Superusers should change the destination directory to another one, preferably to /usr/local/bin.

Once Makefile is customized, simply type

\$ make

to compile, and

\$ make install

to install the package. (The first command may be omitted, is this case it is performed automatically.)

In order to get rid of backup, object, log and similar files, use

\$ make clean

The command

\$ make veryclean

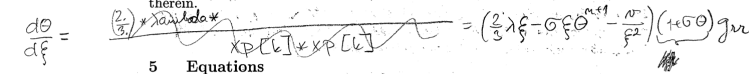
1+60 = PTORH1(Y1,S) = (1+6)+(Y1) $MOG = NTORH1(N_1S,Y1) = ((m)+(S)+(Y1))$ TOTAL WORLD = 1+00+m00 = 1+(m+1)60

> makes the same as the previous one in addition, it deletes the executable files (or symbolic links pointing at them) in package directory, leaving it in original state. To uninstall, run

\$ make uninstall

It is recommended to install the package noutils [Ref] as well if you have not installed it yet.

If you are going to install on a system without the make program (like, e.g., M\$ DOG), look into the Makefile and follow the compilation steps described therein.



The program grpfs starts its performance by numerical integration the initial value problem for the following set of six coupled first-order ordinary differential

value problem for the following set of six coupled first-order ordinary differential equations
$$\begin{cases} \frac{1}{2} \frac{1$$

where

 $\theta(\xi) = 0$

$$g_{rr}(\xi, v; n, \sigma, \lambda) \equiv \frac{1}{1 - 2\sigma(n+1)\left(\frac{v}{\xi} + \frac{1}{3}\lambda\xi^2\right)} \tag{7}$$

for six functions $\theta(\xi)$, $v(\xi)$, $\bar{\xi}(\xi)$, $e_0(\xi)$, $e_{0g}(\xi)$, and $z(\xi)$ subject to initial conditions

$$\theta(0) = 1$$
 $\bar{\xi}(0) = v(0) = e_0(0) = e_{0g}(0) = z(0) = 0$ (8)

on an a priori unknown interval $0 \le \xi \le \xi_1$ with the right end ξ_1 being the first Ptouln + vitorin1 positive root of the equation²

[4/()] (4) de [20m /30] (4) (4)

& The Colan Under certain circumstances, eg, when calculating dependences on o, 77, 18 necessary beep constant not ble central mass devery Se, but the constant of the eg. of state detunicisty of the leg. of state detunicisty of the bound of the bound of instead. Then the egs, (10)- (10) have to be rewitten in the four

The parameters n (polytropic index), σ (relativity parameter), λ (cosmological parameter) may take values from the regions $(0, \infty)$, $(0, \infty)$, and $(-\infty, \lambda_{\text{max}})$, respectively, where λ_{\max} depends on n and σ . (This dependence is beyond the scope of this manual, however.) The limit $\sigma \to 0$ corresponds to Newtonian fluid spheres.

It can be shown [Tooper, 1964, Tooper, 1965] that as far as the parameter λ is small enough to keep the function (7) positive, the function $\theta(\xi)$ is decreasing, dropping to its first zero at $\xi = \xi_1$. Consequently, further task is to find the root ξ_1 and the corresponding values of the remaining five functions $v(\xi_1)$, $\xi(\xi_1)$, $e_0(\xi_1)$, $e_{0g}(\xi_1)$, and $z(\xi_1)$. Finding the root ξ_1 requires, of course, some user-supplied first estimate ξ_0 ; if it turns out to be too large or too small, the integration must resume with a better first estimate ξ_0 .

All the quantities that the program calculates with are dimensionless. The true scaling can be obtained by multiplying the output with appropriate scaling factor for length, mass, energy, mass density, energy density, pressure, and velocity

$$\mathcal{L}^* = \sqrt{\frac{\sigma(n+1)c^2}{4\pi G\rho_c}} \qquad \left[10^4 \text{ km} = 10^9 \text{ cm}\right]$$
 (10)

$$\mathcal{M}^* = 4\pi \mathcal{L}^{*3} \rho_{\rm c} = \frac{c^2}{G} \sigma(n+1) \mathcal{L}^* \qquad \boxed{1 \, \mathcal{M}_{\odot} = 1.989 \times 10^{33} \, \text{g}}$$
(11)

$$\mathcal{E}^* = \mathcal{M}^* c^2 = \frac{c^4}{G} \sigma(n+1) \mathcal{L}^*$$

$$\mu^* = \rho_c \qquad \text{Model Model } \qquad \boxed{\frac{\mu_c}{(\sigma^4 \omega_m)^3}} = 1.989 \times 10^{\frac{1}{2}} \text{M}$$

$$\epsilon^* = \mu^* c^2 = \rho_c c^2 \qquad (1)$$

$$\epsilon^* = \mu^* c^2 = \rho_{\rm c} c^2 \tag{14}$$

$$\pi^* = \sigma \mu^* c^2 = \sigma \rho_{\rm c} c^2 \tag{15}$$

$$c^* = c = 2.99792458 \times 10^{10} \,\mathrm{cm/s}$$
 G [Migner et al. 1972])

where (see [Misner et al., 1973])

$$c^2 = 0.89875518 \times 10^{21} \,\mathrm{cm}^2/\mathrm{s}^2$$
 $\text{M}^2 \,\text{s}^2$ Lg (17)

$$\frac{c^2}{G} = 1.3469 \times 10^{28} \,\text{g/cm}$$
 $m^{2} \, \text{kg}^{-1} \, \text{kg}^{-2} = \frac{1.3469 \times 10^{28} \,\text{g/cm}}{m}$ (18)

$$\frac{c^4}{G} = 1.2106 \times 10^{49} \,\mathrm{g \, cm/s^2} \tag{19}$$

As one can easily see, to calculate true scales it is necessary to know, besides $n, \sigma, \text{ and } G/c^2$, the central mass density ρ_c or fle coulant K.

There is another differential equation to be integrated,

$$\frac{\mathrm{d}\tilde{z}}{\mathrm{d}\xi} = (1 + \sigma\tilde{\theta})^n \left[2\sigma(n+1)g_{rr}(\xi_1, v_1; n, \sigma, \lambda)g_{rr}(\xi, v; n, \sigma, \lambda) \right]^{1/2}
\times Z^{1/2}(\xi, v; \xi_1, v_1; n, \sigma, \lambda)$$
(20)

where

$$Z(\mathbf{MMMS};n,\sigma,\lambda) = \left[1 + \sigma\left(\theta + (n+1)\xi\frac{\mathrm{d}\theta}{\mathrm{d}\xi}\right)\right]\left(\frac{v}{\xi} + \frac{1}{3}\lambda\xi^2\right)$$

$$-\xi \frac{\mathrm{d}\theta}{\mathrm{d}\xi} \left[1 + \sigma\theta + \frac{1}{2}\sigma(n+1)\xi \frac{\mathrm{d}\theta}{\mathrm{d}\xi} \right] \tag{21}$$

on an interval (unification of intervals) determined by condition of optical-geometry embeddability

$$Z(\mathcal{L}_{\mathbf{MMM}}, n, \sigma, \lambda) \ge 0$$
 (22)

with initial conditions

$$\tilde{z}(\xi_{i,1}) = 0
\tilde{z}(\xi_{i,2}) = \tilde{z}(\xi_{f,1})
\vdots
\tilde{z}(\xi_{i,n}) = \tilde{z}(\xi_{f,n-1})$$
(23)

where the indices 'i' and 'f' refer to initial and final boundary of the intervals in which (22) is satisfied.

Equation (20) has to be treated separately from the coupled set (1)–(6). The reason is based on the fact that the right hand side of Eq. (20) depends on the first zero point ξ_1 determined by Eq. (9) in a non-factorizable way, i.e., it cannot be decomposed as $s(\xi_1) \times (the \ rest \ of \ rhs \ (20) \ not \ containing \ \xi_1)$, which would allow appending the Eq. (20) to the set (1)–(6) without the constant factor $s(\xi_1)$ and scaling the results with $s(\xi_1)$ afterwards. As the numerical integration requires evaluation of the right-hand side at an arbitrary point, an interpolation between the tabulated values of the solution of Eqs (1)–(6) must take place.

The most straightforward way how to implement the restriction imposed by limit (22) is redefining the function (21) as

$$Z \leftarrow Z_{+} \equiv \begin{cases} Z & \text{if} \quad Z \ge 0\\ 0 & \text{if} \quad Z < 0 \end{cases} \tag{24}$$

which guarantees satisfying the initial conditions (22) automatically.

The optical geometry embedding diagram is then given parametrically in cylindrical coordinates $(\chi, \alpha, \tilde{z})$ as $(\chi(\xi), \tilde{z}(\xi))$ with the dimensionless radius ξ being the parameter, and

$$\chi(\xi) = \xi(-g_{tt})^{-1/2} = \xi(1+\sigma\theta)^{n+1} \left\{ 1 - 2\sigma(n+1) \left[\frac{v_1}{\xi_1} + \frac{1}{3}\lambda \xi_1^2 \right] \right\}^{-1/2} (25)$$

6 Basic running grpfs

This Section serves as the core tutorial for grpfs. We shall focus on running grpfs solely, with no cooperating utilities, as are, e.g., noutils [Ref]. This topic is covered in Section 7.

6.1 Invocation

The basic invocation of the program grpfs is

\$ grpfs \langle options \left[modifiers \right] \rangle\$

where recognized $\langle options \rangle$ are '-i', '-b', '-f', '-a', and '-x', legal [modifiers] for the '-a' options are 'K', 'B', 'R', and 'S'; the remaining options have no modifiers. Options can be given in arbitrary order and can be grouped. For example, the following commands lead to identical invocations:

- \$ grpfs -i -aK
- \$ grpfs -aK -i
- \$ grpfs -iaK
- \$ grpfs -aKi

However, notice that the modifier ('K' in the example) must immediately follow the option to be modified. Therefore,

\$ grpfs -aiK

is wrong, because the modifier 'K' is separated from the base option '-a' by the option '-i'. The options and modifiers will be fully described in the following Subsections in a free narrative way.

If an unrecognized option or modifier is given, as in

\$ grpfs -iak

the program issues a message like

grpfs: k: unrecognized option, exiting to system

and exits. This example also illustrates that the options and modifiers are case-sensitive.

6.2 Input

Let us launch grpfs in the simplest way, with no option:

\$ grpfs

Program will start in interactive mode, displaying the following text:³

This is grpfs, Revision 1.6 <2002-02-08 19:53:11+01> by S. Hledik ODE solvers based on (C) Numerical Recipes Software.

Fixed stepsize 4th-ord. Runge-Kutta, double precision.

Please feed in the quantities asked for below:

Quantity Numer. type Input/Status Echo check

(r)hoc [10^5 g/cm^3] real,if 0:true

3. The revision number, date and time or other items in the banner line may differ from the actual program run.

rhock [witgemis,...] reel, if wood: bue

a ble Eos eousland k

g (which is also -)

The sunts of k

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The method that grpfs is going to exploit for numerical integration is fixed stepsize 4th-order Runge-Kutta. This is the default, but explicitly using of this method can be required by typing

\$ grpfs -f

where the option '-f' stands for 'fixed/stepsize.' Similarly, interactive run might be explicitly specified by the option \(-i' \), which is the default. Thus, invocation

\$ grpfs -if

works in exactly the same way as the previous ones. These options might seem to be superfluous, however, they can help the user to keep in mind what he/she is actually doing.

Now the program is expecting you to enter the central mass density ρ_c (which is intrinsically positive number) in the units indicated. If you enter zero (0, 0., .0, 0.0, .), the response looks like this:

(x)hoc [10^5 g/cm^3] real.if>0:true 0 rhook (- a-, a) real if x of her Dimensionless output opted no scales calculated Ignored --

Supplying zero for the central mass density (which cannot represent a valid value), the program is not able calculate the true scales (10)-(15); hence, both the output will be in dimensionless quantities and the scaling factors (10)-(15) will not be provided. If you enter negative central mass density (which also cannot represent a valid value), the program accepts the absolute value as the central mass density, and the minus sign is understood as a command 'do calculate and show the scaling factors (10)-(15), but do not multiply the output and frowell with them, leave the output dimensionless instead.' In such case, the response looks like this:

but seales provided Accepted | 1.000E+00 wdustood as Po as

Finally, if you enter a positive value, it will be accepted without change and dimensional output quantities will be multiplied by appropriate scale factors. Exactly my estimate the scale factors.

(r)hoc [10°5 g/cm°3] real,if>0:true 1.0 True scale on output opted Accepted (r) 1.000E+00

The second entry is the polytropic index $n \geq 0$. The input and grpfs's response looks like this:

(p)olytropic index real >= 0.0 1.0
Accepted 1.000

If you enter a value that is in conflict with the specification in the column 'Numer. type', the program will response (and the user may correct his/her mistake) like this:

(p)olytropic index real >= 0.0 -.2

Bad, retry:

1.0

Accepted 1,000

Similarly, entering a wrong value at any prompt, it will be rejected and the user will be prompted to provide a correct value. This protection might be bypassed by using the batchmode (see Subsection 7.1), but the results would be nonsensical.

Now, assume that we entered 0 at the first prompt. Then the rest of interaction grpfs—the user is self-explanatory, as the following example shows:

(s)igma	real >= 0.0	.2 Accepted	2.000E-01
(1) ambda		1.0E-34	
(f) inal xi guess	real > 0.0	4 Accepted	
# of s(t)eps	integer > 2	25 Accepted	25

Here we entered σ , the dimensionless 'cosmological parameter' $\lambda = \rho_{\rm vac}/\rho_{\rm c}$, the estimate of the first positive root of (9), and the number of steps. Remember: the points at which the functions $\theta(\xi)$, $v(\xi)$, $\bar{\xi}(\xi)$, $e_0(\xi)$, $e_{0g}(\xi)$, $z(\xi)$, and $\tilde{z}(\xi)$ will be evaluated, are enumerated $0, 1, \ldots, \langle number\ of\ steps \rangle$, their number being $\langle number\ of\ steps \rangle + 1$.

If a non-zero (either positive or negative) has been entered at the first prompt, the prompting for the 'cosmological parameter' and the response that follows will be modified like this:

(1|L)ambda [1|cm^-2] real | *real 1.0E-34 Accepted (1) 1.000E-34

That means if an 'ordinary' number is entered as in the example above, the programs accepts it as 'cosmological parameter' λ and, for the user's information,

will calculate the cosmological constant

$$\Lambda = \frac{8\pi G}{c^2} \rho_{\rm c} \lambda \tag{26}$$

in units of cm⁻². On the other hand, if the number entered is preceded by a star '*' (no space between), the number will be accepted as the cosmological constant Λ (in units of cm⁻²) and the 'cosmological parameter' λ will be calculated for the program's (and user's, of course) needs:

Notice that if ρ_c is not given, the relationship (26) is lacking any sense, which grpfs is aware of.

Now let us try another invocation (the option '-i' may be omitted):

The program writes out the following banner

This is grpfs, Revision 1.6 <2002-02-08 19:53:11+01> by S. Hledik ODE solvers based on (C) Numerical Recipes Software.

Adaptive stepsize 5th-ord. Cash-Karp Runge-Kutta, double precision. Please feed in the quantities asked for below:

indicating that now we have opted the adaptive stepsize 5th-order Cash-Karp Runge-Kutta method (see [Press et al., 1997]). The 'K' modifier stands for 'Kutta' and is the default modifier, i.e., launching

behaves as if the 'K' modifier were present. However, it is reasonable to specify the modifier explicitly.

The '-a' option is of higher precedence than '-f' option, i.e., if both these options would be given in any order, the program would behave as if only '-a' were specified.

The prompts and behavior up to the final ξ guess are identical, then the prompting and responses become different, as shown in the following example:

en e	(a)ccuracy		.000001 Accepted	1.0E-06	alk mark sav
	1st stepsize (g)uess	0.0 <real< th=""><th>.1 Accepted</th><th>1.0E-01</th><th></th></real<>	.1 Accepted	1.0E-01	
kmax	ma(x) points to save	int 1, or >3	25 Accepted	25	
dysav	(m)in sav. interval	real,if<0: f/m	-25 Acc 4.0/-25	1.6E-01	

Steps = ponts - 1

Steps = ponts - 1

Luax 3:

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Luax 6:

L

The user is asked for to supply the accuracy desired, then the first stepsize estimate, and maximum number of points (not steps) to save, and the minimum saving interval. If the last value is negative, the absolute value of (final ξ guess)/(minimum saving interval) will be used instead.

Replacing the 'K' modifier with 'B', 'R', or 'S' indicates using the adaptive stepsize Bulirsch–Stoer, 4th-order Rosenbrock, and Bulirsch–Stoer semi-implicit midpoint rule method, respectively (see [Press et al., 1997]). The modifier precedence can be symbolically expressed as K < B < R < S. The structure of input remains exactly the same for all these modifiers.

6.3 Output

General description of the output format can be found in [Ref]. Here we only summarize its main features and give more detailed account of some extras thrown in particularly for grpfs.

The output is separated into commentary part and the output data itself. Each line of the former part is preceded by comment sign '#', while the latter part (tabulated output data) are formatted in c space-separated columns.

The commentary part comprises the following sections.

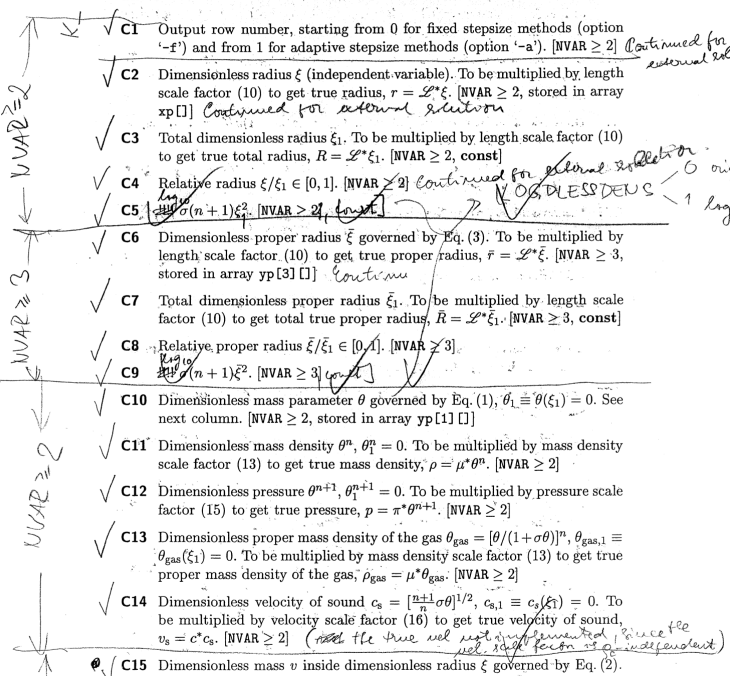
- #Mm This line at the very beginning of the output marks the method used. The digit m takes value 0 for fixed stepsize method (option 'f' or none) and 1 for adaptive stepsize methods (option 'a' with no modifier or with modifiers 'K', 'B', 'R', or 'S').
- #ID_CHART This sections accommodates various technical details concerning date and time, means of invocation, method used, host name and IP address, code version, compiler options and many others. the purpose of this section is to identify the run as exactly as possible. Because the actual contents may change, we shall not give it here explicitly. This section is omitted unless preprocessor option '-DVERBOSE' is set.
- #INPUT In this section, the input data are echoed. Notice that the numbers are commented out by strings '#!' instead of '#'; see subsection 7.1 and manual [Ref] for explanation.
- #COMPUTATION LOG This section gives information about solving ODE's and auxiliary numerical routines (such as interpolation) calls. This section is omitted unless preprocessor option '-DVERBOSE' is set.

#OUTPUT ℓ This string starts tabulated output consisting of ℓ rows of data.

The sections described above are separated by single comment lines containing the comment sign '#' only.

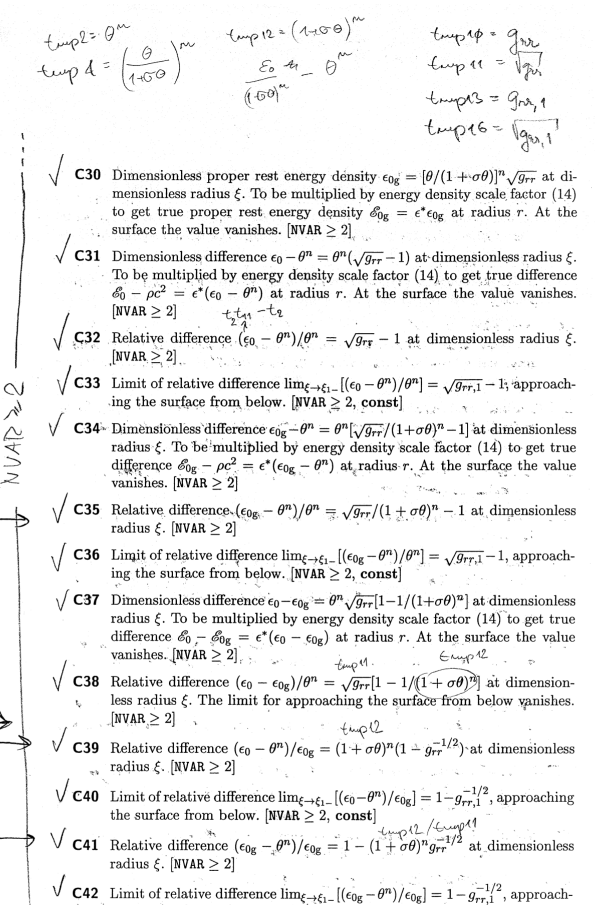
The columns of tabulated output are described in the following. The first column shows the output row number. The term 'dimensionless' refers to quantities that can be scaled by the scaling factors (10)–(16), while the term 'relative' refers

either to quantities that range from 0 through 1 since they are divided by the corresponding value on the surface or total quantity (at the point ξ_1), or to quantities related to another quantities (typically energies or energy densities). The value of a language C symbolic constant NVAR shows the number of Eqs (1)–(6) (NVAR = 2, 3, ..., 6) and (20) (NVAR = 7) that have to be taken into account for the particular column to be computed (it is padded with zero otherwise). The boldface **const** reminds that all rows are of constant value.



 ξ . To be multiplied by energy density scale factor (14) to get true proper energy density $\mathscr{E}_0 = \epsilon^* \epsilon_0$ at radius r. At the surface the value vanishes.

 $[NVAR \ge 2]$



ing the surface from below. [NVAR ≥ 2 , const]

C43 Relative difference $(\epsilon_0 - \epsilon_{0g})/\epsilon_{0g} = (1 + \sigma\theta)^n - 1$ at dimensionless radius

14

original Con 16 och i restite
all: ell*
all: E* ξ . The limit for approaching the surface from below vanishes. [NVAR ≥ 2] Dimensionless proper energy e_0 inside dimensionless radius ξ governed by Eq. (4). To be multiplied by energy scale factor (12) to get true proper energy $E_0 = \mathscr{E}^* e_0$ inside radius r. [NVAR ≥ 4 , stored in array yp [4] []] Total dimensionless proper energy $e_{0,1} \equiv e_0(\xi_1)$. To be multiplied by energy scale factor (12) to get true total proper energy $E_{0,1} = \mathscr{E}^* e_{0,1}$. $[NVAR \ge 4, const]$ C46 Dimensionless gravitational potential energy $\omega \equiv e_0 - v$ inside dimensionless radius ξ . To be multiplied by energy scale factor (12) to get true gravitational potential energy $\Omega = E_0 - mc^2 = \mathscr{E}^*(e_0 - v)$ inside radius $r. [NVAR \ge 4]$ Total dimensionless gravitational potential energy $\omega_1 = e_{0,1} - v_1$. To be multiplied by energy scale factor (12) to get total true gravitational potential energy $\Omega_1 = E_{0,1} - Mc^2 = \mathscr{E}^*(e_{0,1} - v_1)$. [NVAR ≥ 4 , const] C48 Relative gravitational potential energy $\omega/v = e_0/v - 1$ inside dimensionless radius ξ . [NVAR ≥ 4] Total relative gravitational potential energy $\omega_1/v_1 = e_{0,1}/v_1 - 1$. [NVAR \geq 4, const C50 Dimensionless proper rest energy e_{0g} inside dimensionless radius ξ governed by Eq. (5). To be multiplied by energy scale factor (12) to get true proper rest energy $E_{0g} = \mathscr{E}^* e_{0g}$ inside radius r. [NVAR ≥ 5 , stored in array yp[5][[][] C51 Total dimensionless proper rest energy $e_{0g,1}$. To be multiplied by energy scale factor (12) to get true total proper rest energy $E_{0g,1} = \mathscr{E}^* e_{0g,1}$. $[NVAR \geq 5, const]$ C52 Dimensionless binding energy $e_b \equiv e_{0g} - v$ inside dimensionless radius ξ . To be multiplied by energy scale factor (12) to get true binding energy $E_{\rm b} = \mathscr{E}^* e_{\rm b}$ inside radius r. [NVAR ≥ 5] Total dimensionless binding energy $e_{\rm b,1}=e_{\rm 0g,1}-v_{\rm 1}$. To be multiplied by energy scale factor (12) to get true total binding energy $E_{b,1} = \mathscr{E}^* e_{b,1}$. $[NVAR \geq 5, const]$ C54 Relative binding energy $e_b/v = e_{0g}/v - 1$ inside dimensionless radius ξ . Caution: the limit for $\xi \to 0_+$ is $(1+\sigma)^{-n} - 1$. [NVAR ≥ 5] Total relative binding energy $e_{\rm b,1}/v_{\rm 1}$. [NVAR ≥ 5 , const] Relative binding energy ω/e_{0g} inside dimensionless radius ξ . [NVAR ≥ 5] **C57** Total relative binding energy $\omega_1/e_{0g,1}$. [NVAR ≥ 5 , const]

16

G=pe/Pge&c2

C73 Polytropic/adiabatic index n. [NVAR ≥ 2 , const]

 $\sqrt{\text{C74}^{\text{T}}}$ Relativity parameter $\sigma = p_{\text{c}}/\rho_{\text{c}}c^2$. !!!! For ADIABATIC [NVAR ≥ 2 , const]

 $\sqrt{\text{C75}^6}$ Cosmological parameter $\lambda = \rho_{\text{vac}}/\rho_{\text{c}}$. See also Eq. (26). [NVAR ≥ 2 , const]

C76 Length scale factor \mathcal{L}^* defined by Eq. (10). [NVAR ≥ 2 , const]

C77 Mass scale factor \mathcal{M}^* defined by Eq. (11). [NVAR ≥ 2 , const]

C78 Energy scale factor \mathscr{E}^* defined by Eq. (12). [NVAR ≥ 2 , const]

Mass density scale factor μ^* defined by Eq. (13). [NVAR \geq , 2, const]

C80 Energy density scale factor ϵ^* defined by Eq. (14). [NVAR ≥ 2 , const]

C81 Pressure scale factor π^* defined by Eq. (15). [NVAR ≥ 2 , const]

C82 Velocity scale factor et defined by Eq. (16). [NVAR 22, Const]

Examples

profes's output to the file output dat after an epsize method: here is an example how to send grpfs's output to the file output dat after an

interactive run using the fixed-stepsize method: \$ grpfs -if > output.dat

To append another output to a previous one, use

\$ grpfs -if >> output.dat

Advanced usage

This Section describes some features of grpfs that rely on Unix-like systems' capability of redirecting standard terminal input/output into a file or from a file and piping it between processes, as has been touched in the previous Section. Therefore, some basic knowledge of these features can be beneficial.

7.1 Feeding input from an output file

Although the title of this Subsection may sound rather weirdly, the output produced by grpfs can really be reused as the input. If you examined the output thoroughly, you must have noticed that certain comment lines start with strings '#!' instead of mere '#'. These metacomments precede exactly those numerical strings that the program requires as input. So, if you redirect the grpfs's standard input to a file written out beforehand and named, say, output.dat, like this,

\$ grpfs -b < output.dat</pre>

the program will skip all lines commented out with '#' but will load the numerical strings that follow the metacomment strings '#!'. Don't forget the '-b' option; although this omission has no influence on the function, the screen would be messed with nasty interactive prompts. Moreover, this option turns on the input data conformity check (see below in this Subsection), and turns off checking for meaningful values.

So, this capability makes it possible to rerun any calculation you have stored in a file. In fact, the output should exactly mimic the input (with the exception of some records in the ID_CHART section, of course), as you can convince yourself by giving the commands

- \$ grpfs -b < output.dat > output.new
- \$ diff output.dat output.new

The structure of grpfs's input differs for fixed stepsize and adaptive stepsize methods. That is why each output is labeled with a 'magic string' at the very beginning: the fixed stepsize method has magic string '#M0', while the adaptive stepsize ones have #M(method number), where (method number) is 1 for 5th-order Cash-Karp Runge-Kutta method, 2 for the Bulirsch-Stoer method, 3 for 4-th order Rosenbrock method, 4 for Bulirsch-Stoer semi-implicit midpoint rule method. You can only input data conforming with the method selected by the '-f' or '-a' option (in other words, choose '-f' or no option if the input is labeled '#M0' and '-a' if the input is labeled '#M1' through '#M4'), otherwise the program stops, complaining

grpfs: input dataset: mismatch between option/stdin, exiting to system
There should be emphasized that all the adaptive stepsize outputs are interchangeable if used as input, however, the output data may differ.

7.2 Filtering output through noutils

See the package noutils [Ref] for more info.

7.3 Examples

Here will be examples on using less, tee, etc.

8 Configuration

There are three preprocessor options in the Makefile 'Customization section,' subsection 'Preprocessor options.'

DOPT1 = -DDBL_PREC If set, grpfs will use double precision floating-point arithmetic—this is highly recommended since the single precision range is frequently not sufficient to hold values of cosmological constant and similar ones.

This option is set by default. Unless set, grpfs will use single precision floating-point arithmetic.

DOPT2 = -DVERBOSE If set, grpfs runs in a very verbose mode; this option is recommended and set by default.

DOPT1 = -DRAT_EXTR If set, grpfs will use rational extrapolation subroutine instead of the polynomial one. For experimental purposes and disabled by default.

It is also worth mentioning the preprocessor symbolic constant NVAR. Setting it to $1, 2, \ldots, 7$ one can control the number of the equations to solve (see also Section 6). This constant can be set in the header file grp.h.

There is a lot of customizable symbolic constants in grp.h, grp_io.h and grp_node.h. In this point, the user is recommended to study [Press et al., 1997] and to read the source codes.

When using package noutils [Ref] and dealing with constant combination of columns for a long time, it is worth setting and exporting the environmental variable PICKOLS. See [Ref] for detailed information.

9 Bugs

Please send bug reports to Stanislav.Hledik@fpf.slu.cz. Indicate clearly grpfs-bugs in the subject field. Another possibility is to send complaints to /dev/null.

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fluid spheres, 2 general relativistic, 2 adiabatic, 2 polytropic, 2 Newtonian, 4

A Differences between grpfs and grafs

To be finished!!!!

B Source code

The kernel of the system—the numerical routines that perform integration, interpolation and extrapolation—are adopted from [Press et al., 1997] and included in module grp_node.c. The program also utilizes the memory allocating and other utility routines from [Press et al., 1997] concentrated in the module grp_nutl.c, but the latter are freely distributable contrary to the former ones that are subject to copyright and cannot be placed in a freely accessible site.

The user can make some settings in the header file grp.h, especially in the section marked as 'customization section.' Pe = Kge 1+1 G = Pe => Ockge = Kgegetn 1/2 (4.7)

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