A. Introduction and overview

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GRTensorIII is a package for the calculation and manipulation of components of tensors and related objects. Rather than focus upon a specific type or method of calculation, the program has been designed to operate efficiently for a wide range of applications and allows the use of a number of different mathematical formalisms. Algorithms are optimized for the individual formalisms and transformations between formalisms has been made simple and intuitive. Additionally, the package allows for customization and expansion with the ability to define new objects, user-defined algorithms, and add-on libraries.

Regardless of the algorithm or formalism used, it is often the case that only certain simplifications applied at a crucial stages can make some problems tractable. For this reason, GRTensorIII has been designed to provide full control of the calculation path and of the simplifications to be performed at each stage. Once a calculation is completed, a variety of commands are available for the manipulation and simplification of results.

In designing the package, emphasis has also been placed on the interface, allowing simple user input, as well as presenting readable output. Metrics and basis vectors are easily defined and can be saved for later use. Calculations are specified in an intuitive manner using a minimal number of commands.

This booklet introduces a number of characteristics and features of the GRTensorIII computer algebra package, emphasizing its input and output facilities, the ability to define new tensor objects, and the ability to use and switch between a number of different formalisms (classical tensor methods, general bases, and null tetrads). This booklet itself is intended only as a brief overview. More detailed information regarding the commands introduced here can be found in Booklets B–F:

Booklet B: Specifying spacetimes

C: Calculating tensor components

D: Defining new tensors

E: Bases and tetrads

Together these articles are intended to serve as both a user's guide and command reference for the GRTensorIII package. Unfortunately because of this dual purpose, there are some shortfalls in their ability to perform each task: Someone looking for tutorial information might find themselves buried in uninteresting technical points, while experienced users will find some sections overly pedagogic.

It is not intended that the booklets be read linearly from start to finish. To begin, only the information from the first sections of the *Specifying spacetimes* and *Calculating tensor components* booklets are needed to perform some sophisticated calculations. Once familiarity with the interface is gained, the ability to define new tensors and build customized libraries, described in Booklet *D: Defining new tensors*, greatly increase the power of the package. For the first time user, probably the best way to approach the package is to study and execute some of the demonstrations available in the worksheets directory provided with the package, referring to these booklets and the online help as necessary.

Release 1.00: GRTensorIII is a adaptation of GRTensorII version 1.50 developed in 1994-1999. It is primarily a refactoring of internal data representation with some updates to the "modern" Maple environment.

- move all variables inside a scoped Maple package
- include all extension objects from GRII (basic etc.) directly in the main package

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- support Maple worksheet dialog based input
- remove some seldom used commands (grloaddef, grsavedef, grloadobj, grsaveobj)
- no changes have been made to the expressions used to define the standard tensor objects

The version of GRTensorIII described in these booklets runs within the Maple environment. It has been tested with Maple 2016 in command line and worksheet mode. We encourage users to check back periodically for new versions of software, documentation, and demonstrations, at http://gitlab.com/grtensor/grtensor. This represents the "official" version.

A copy of the GRTensorIII source code is available online at http://gitlab.com/grtensor/GRTensorIII. Merge requests will be considered.

1 Startup

The GRTensorIII package is distributed as Maple package. To start the GRTensorIII package, in a new Maple session ensure that the location of greensor/lib is in the Maple path. It can be added by:

```
libname := libname, "path";
```

The package is loaded in the standard manner:

```
with(grtensor);
```

Build-in help pages can be viewed via:

?grtensor

The examples described below are designed to be run in a single GRTensorIII session.

2 Calculating components

Tensors are specified by their name and index configuration, using the abbreviations dn and up to indicate covariant and contravariant indices, respectively. For instance, the Ricci tensor could be referenced

$$\texttt{R(dn,dn)} \quad (=R_{ab}), \qquad \texttt{R(up,dn)} \quad (=R^a_{\ b}), \qquad \texttt{R(bup,bdn)} \quad (=R^{(a)}_{\ \ (b)}),$$

where the last example uses the labels bup and bdn to reference the components of the Ricci tensor in terms of a basis rather than metric coordinates.

Three commands are used most frequently within a GRTensorIII session:

great calculates the components of tensors;

grdisplay displays the components;

gralter applies simplification routines to tensors;

Thus, to calculate the covariant Ricci tensor for the predefined Kerr metric, we might use the command sequence

```
> qload ( kerr ):
> grcalc ( R(dn,dn) ):
   Calculating g(dn,dn,pdn) for kerr ... Done.
   Calculating Chr(dn,dn,dn) for kerr ... Done.
   Calculating detg for kerr ... Done.
   Calculating g(up,up) for kerr ... Done.
   Calculating Chr(dn,dn,up) for kerr ... Done.
   Calculating R(dn,dn) for kerr ... Done.

> grdisplay ( R(dn,dn) ):
```

For the kerr spacetime:

$$R_{rr} = -a^{2} \left(r^{2} - r^{2} \sin(\theta)^{2} + 2 r^{2} \cos(\theta)^{4} - 3 r^{2} \cos(\theta)^{2} + 2 r^{2} \sin(\theta)^{2} \cos(\theta)^{2} + a^{2} \cos(\theta)^{4} - a^{2} \cos(\theta)^{2} + a^{2} \cos(\theta)^{2} \sin(\theta)^{2} \right) / \left(\left(-1 + \cos(\theta)^{2} \right) \left(r^{2} - 2 m r + a^{2} \right) \left(r^{2} + a^{2} \cos(\theta)^{2} \right)^{2} \right)$$

$$R_{\theta\theta} = \dots \text{ etc.}$$

Covariant Ricci

At this point, since no simplifications of the results have been applied, the components of R_{ab} are not obviously zero as we would expect. The following commands are needed:

```
> gralter ( R(dn,dn), trig ):
> grdisplay ( R(dn,dn) ):
```

For the kerr spacetime: $Covariant\ Ricci$ $R(dn,dn)=All\ components\ are\ zero$

The first command applies trigonometric simplification to the components of R(dn,dn). Upon using grdisplay(), we see that the the simplification has indeed reduced the components to zero.

Note that for a more complicated metric, it may be the case that it is better to carry out the computation in stages. That is, we can first calculate the Christoffel symbols and apply simplifications, then the Ricci tensor, and finally, for instance, the Ricci scalar.

A number of alternate simplification routines exist within gralter(), including trigonometric, radical, expansion and factorization techniques. The command grmap() can also be used to apply any Maple function to the components of a tensor.

The commands listed in this Section are described in more detail in Booklet *C: Calculating tensor components*. The calculation of objects in a coordinate basis is described in Booklet *E: Bases and tetrads*.

3 Input of spacetimes

Spacetimes are specified within GRTensorIII through use of the makeg() facility. They can be entered in the following forms:

Metric, g_{ab} A $n \times n$ symmetric matrix;

Line element An equation of the standard form $ds^2 = \dots$;

Non-holonomic basis A set of basis vectors (covariant or contravariant) with an inner product;

Null tetrad As described by the Newman-Penrose formalism.

As an illustrative (but computationally trivial) example, we consider the twisting Einstein-Maxwell solution described on p. 135 of Kramer, et al. [1],

$$kds^{2} = -2\left(x^{a}du - \frac{dy}{(a+1)x}\right)\left[dt + a(tdx + dy)/x + f(t)\left(x^{a}du - \frac{dy}{(a+1)x}\right)\right] + (dx^{2} + dy^{2})(t^{2} + 1)/2x^{2},$$

We can enter this metric directly in this form using makeg(), giving it the name 'sm':

In Maple worksheets the following prompts will appear in dialog boxes. In the example here we show the responses in command-line mode.

```
> makeg ( sm ):
  Makeg 2.0: GRTensorIII metric/basis entry utility
   To quit makeg, type 'exit' at any prompt.
  Do you wish to enter a 1) metric [g(dn,dn)],
                             2) line element [ds],
                             3) non-holonomic basis [e1...e4], or
                             4) null tetrad [1,n,m,mbar]?
> 2:
  Enter coordinates as a LIST (e.g. [r,theta,phi,t]):
> [ t, x, y, u ]:
   Enter the line element using 'd[coord]' to indicate differentials.
   (for example, r^2*(d[theta]^2 + sin(theta)^2*d[phi]^2)
   [Type 'exit' to quit makeg]
   ds^2 =
> 1/k*(-2*(x^a*d[u] - d[y]/((a+1)*x))*(d[t] + a*(t*d[x])
   + d[y])/x + f(t)*(x^a*d[u]-d[y]/((a+1)*x))) + (d[x]^2 + d[y])/((a+1)*x))
   d[y]^2(t^2+1)/(2*x^2):
```

Note that in the above expression the coordinate differentials are labeled using the variables d[coordinate], allowing the line element to be entered in exactly the form that it is given in

[1]. The output from makeg is:

$$\begin{split} ds^2 &= 2\frac{dt\ dy}{k(a+1)x} - 2\frac{x^a dt\ du}{k} + \left(\frac{1}{2}\frac{t^2}{kx^2} + \frac{1}{2}\frac{1}{kx^2}\right) dx^2 + 2\frac{a\ t\ dx\ dy}{k(a+1)x^2} \\ &- 2\frac{x^a a\ t\ dx\ du}{kx} + \left(2\frac{a}{k(a+1)x^2} - 2\frac{f(t)}{k(a+1)^2x^2} + \frac{1}{2}\frac{t^2}{kx^2} + \frac{1}{2}\frac{1}{kx^2}\right) dy^2 \\ &+ 2\left(-\frac{x^a a}{kx} + 2\frac{x^a f(t)}{k(a+1)x}\right) dy\ du - 2\frac{(x^a)^2 f(t) du^2}{k} \end{split}$$

makeg completed.

The parameter values can now be set for the particular Type-III case studied by Siklos and MacCallum [2] (referred to in [1] on p. 135):

$$> a := 1/2$$
: $k := -32/(78*lambda)$: $f(t) := (13*t^2+17)/32$:

For this metric, we calculate the (rather uninteresting, in this case) components of the Ricci tensor:

```
> grcalc ( R(dn,dn) ):
> grdisplay ( _ ):
```

For the sm metric:

$$R_{ab} = \begin{bmatrix} 0 & 0 & -\frac{13}{8}\frac{1}{x} & \frac{39}{16}\sqrt{x} \\ 0 & -\frac{39}{32}\frac{t^2+1}{x^2} & -\frac{13}{16}\frac{t}{x^2} & \frac{39}{32}\frac{t}{\sqrt{x}} \\ -\frac{13}{8}\frac{1}{x} & -\frac{13}{16}\frac{t}{x^2} & -\frac{65}{192}\frac{5+t^2}{x^2} & -\frac{13}{128}\frac{5+13t^2}{\sqrt{x}} \\ \frac{39}{16}\sqrt{x} & \frac{39}{32}\frac{t}{\sqrt{x}} & -\frac{13}{128}\frac{5+13t^2}{\sqrt{x}} & \frac{39}{256}x(13t^2+17) \end{bmatrix}$$

The specification of metrics and bases is described more fully in Booklet B: Specifying space-times. The grcalc(), gralter() and grdisplay() commands are the subject of Booklet C: Calculating tensor components.

4 The Newman-Penrose formalism

A more complicated problem is that of determining the Petrov type of this metric. In order to do this in GRTensorIII, the metric must be cast in the form of a Newman-Penrose (NP) null tetrad. Since no tetrad obviously presents itself from the form of the metric given above, we can use the command nptetrad() to generate one. This command constructs a set of null basis vectors satisfying the properties of a basis for the Newman-Penrose formalism.¹ Naturally, this tetrad may not be optimal for further calculation, but is often a good start to which further simplifications can be applied.

```
> nptetrad([u,t]):
```

The metric signature of the sm spacetime is +2. In order to create an NP-tetrad, the signature of g(dn,dn) will be changed

¹Since the -2 signature convention of the NP formalism conflicts with the +2 Landau-Lifschitz convention recommended for metrics in GRTensorIII, the construction of the tetrad also involves a signature change. This is carried out automatically if requested.

to -2.
Continue? (1=yes):
1:
The signature of the sm spacetime is now -2.

Null tetrad (covariant components)

$$l_a = \left[\begin{array}{ccc} -\frac{9}{4} \lambda \sqrt{x} & -\frac{9}{8} \frac{\lambda t}{\sqrt{x}} & \frac{3}{64} \frac{\lambda \left(-7+13 \, t^2\right)}{\sqrt{x}} & -\frac{9}{128} \lambda x \left(13 t^2+17\right) \end{array} \right]$$

$$m_a = \left[\begin{array}{ccc} 0 & 0 & -\frac{2}{3} \frac{1}{x^{3/2}} & 1 \end{array} \right]$$

$$m_a = \left[\begin{array}{ccc} 0 & \frac{3/4 \, i \left(t^2+1\right) \lambda}{x \sqrt{-(t^2+1) \lambda}} 3/4 \, \frac{\left(t^2+1\right) \lambda}{x \sqrt{-(t^2+1) \lambda}} & 0 \end{array} \right]$$

$$mbar_a = \left[\begin{array}{ccc} 0 & \frac{-3/4 \, i \left(t^2+1\right) \lambda}{x \sqrt{-(t^2+1) \lambda}} 3/4 \, \frac{\left(t^2+1\right) \lambda}{x \sqrt{-(t^2+1) \lambda}} & 0 \end{array} \right]$$

The null tetrad has been stored as e(bdn,dn).

From this point we can proceed to determine the Petrov type of our example metric by first calculating the spin coefficients (NPSpin and their conjugates NPSpinbar) and the Weyl scalars (WeylSc) [3].

```
> grcalc (NPSpin, NPSpinbar):

> grcalc (WeylSc):

> gralter (_, factor):

> grdisplay (_):

\Psi 0 = -\frac{5}{64} \frac{Ix(13t^2 + 10It + 11)}{(t+I)(-t+I)^2}
\Psi 1 = \frac{-5/8i\sqrt{x}(t+i)\lambda^2(-t+i)^2}{(-(t^2+1)\lambda)^{5/2}}
\Psi 2 = 0
\Psi 3 = 0
\Psi 4 = 0
```

Note that the calculation of NPSpin and NPSpinbar did not need to be specified in a separate step, since this the calculation of these objects implicit in the calculation of WeylSc. However, in general a good rule of thumb is to calculate these quantities first and simplify them fully (not required in this case) before proceeding to the curvature scalars.

```
> grcalc ( Petrov ):
> grdisplay ( _ ):
```

```
For the sm spacetime:
Petrov Type = III (or simpler)
```

Note the caveat 'or simpler' in the final line of output. The calculation of the Petrov type involves determining whether a number of intermediate quantities can be evaluated to zero [4]. It is sometimes the case that complicated ways of expressing zero do not, in fact, reduce to zero within the computer algebra system. In such cases it will return a Petrov type which is unnecessarily specialized. To determine the reliability of the computer's evaluation, the PetrovReport() command is provided, which details the intermediate calculations performed in the determination of the Petrov type.

> PetrovReport():

```
The conclusion 'Petrov type = III (or simpler)' for the sm metric
was based on the following results:
Weyl scalar Psi0 could not be evaluated to zero.
Weyl scalar Psi1 could not be evaluated to zero.
Weyl scalar Psi2 = 0
Weyl scalar Psi3 = 0
Weyl scalar Psi4 = 0
---> Therefore the metric is Petrov III (or simpler).
```

We see from the previous output that the indeed the calculations have been carried out correctly (all of the terms which the computer believes to be non-zero are actually non-zero).

The use of null tetrads and general bases is treated in more detail in Booklet E: Bases and tetrads.

5 Defining new tensors

Finally, we describe a facility which exists within GRTensorIII for defining tensors which are not part of the standard object libraries. Here we define the Bel-Robinson tensor,

$$T_{cdef} := C_{acdb}C^{a}{}_{ef}{}^{b} + C^{*}_{acdb}C^{*a}{}_{ef}{}^{b}$$

as well as it's trace and its divergence using the command grdef():

```
> grdef ('T{(c d e f)} := C{a c d b}*C{^a e f^b}+ Cstar{a c d b}
    *Cstar{^a e f^b}'):
> grdef ('TT{(c d)} := T{^a a c d}'):
> grdef ('TC{(b c d)} := T{^a b c d ;a}'):
```

In the strings defining these tensors, contravariant indices are indicated by preceding them with a caret, '^', while covariant differentiation is indicated as usual, with a semi-colon, ';'. The braces on the left-hand side of the assignment in each command indicate that the newly defined tensors should be assumed to be symmetric under interchange of their indices. The calculation functions which are automatically generated by grdef() will take these symmetries into account to improve the efficiency of calculations.

We now load the Kerr-Newman metric and calculate these objects:

```
> qload(newkn):
```

Coordinates:

$$x^a = [r \ u \ \phi \ t]$$

Line element

$$\begin{split} ds^2 = & \frac{(r^2 + u^2)dr^2}{r^2 - 2mr + a^2 + Q^2} + \frac{(r^2 + u^2)du^2}{a^2 - u^2} + \frac{(a^2 - u^2)\left(r^2 + a^2 + \frac{(a^2 - u^2)(2mr - Q^2)}{r^2 + u^2}\right)d\phi^2}{a^2} \\ & - \frac{(a^2 - u^2)(2mr - Q^2)d\phi dt}{a(r^2 + u^2)} + \left(-1 + \frac{2mr - Q^2}{r^2 + u^2}\right)dt^2 \end{split}$$

 $Kerr-Newman \ Solution \ in \ Boyer-Lindquist \ coordinates \ (u=a*cos(\theta))$

> grcalc (T(dn, dn, dn, dn)):

We display a component (e.g. T_{rrrr}):

> grcomponent (T(dn, dn, dn, dn), [r, r, r, r]):

$$6\frac{m^2r^2-2mrQ^2+m^2u^2+Q^4}{(r^2+u^2)^2(r^2-2mr+a^2+Q^2)^2}$$

We now check that $T^a{}_{acd} = 0$ and that $\nabla_a T^a{}_{bcd} = 0$ in vacuum:

- > grcalc (TT(dn,dn)):
 > grdisplay(_):
- $TT(dn,dn) = All \ components \ are \ zero$

- > Q:=0:
- > grcalc (TC(dn,dn,dn)):
- > grdisplay(_):

$$TC(dn,dn,dn) = All \ components \ are \ zero$$

A more thorough discussion of the definition of tensors in GRTensorIII is to be found in Booklet D: Defining new tensors.

Conventions

GRTensorIII can perform tensor calculations in any number of dimensions and for symmetric metrics of arbitrary signature. Tensor indices generally take values from $1 \dots n$ in an n-dimensional spacetime.

In its definitions of curvature tensors, GRTensorIII follows the Landau-Lifshitz spacelike conventions. That is, the curvature (Riemann) tensor is defined by

$$R^{a}{}_{bcd} := \frac{\partial \Gamma^{a}_{bd}}{\partial x^{c}} - \frac{\partial \Gamma^{a}_{bc}}{\partial x^{d}} + \Gamma^{a}_{ec} \Gamma^{e}_{bd} - \Gamma^{a}_{ed} \Gamma^{e}_{bc},$$

the Ricci tensor by

$$R_{ab} := R^c{}_{acb},$$

and the Einstein tensor by

$$G_{ab} := R_{ab} - \frac{1}{2}Rg_{ab}.$$

Although spacetimes of any signature can be defined, for four dimensional Lorentzian metrics the signature +2 is recommended. Most standard curvature calculations will be carried out correctly regardless of the signature convention. However, some tensors (in particular the vector operators of Booklet C: Calculating tensor components) which depend on the normalization of timelike or spacelike vectors can be affected by the choice of signature. If a signature other than +2 is employed, users should carefully check tensor definitions for signature dependent terms.

The only deviation from this standard signature is in the specification of the Newman-Penrose formalism, which follows the original definitions of [3], and thus requires a -2 signature for spacetimes.

References

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