Numerical tools to validate stationary points of $SO(8)$-gauged $\mathcal{N} = 8$ $D = 4$ supergravity

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Abstract
Until recently, the preferred strategy to identify stationary points in the scalar potential of $SO(8)$-gauged $\mathcal{N} = 8$ supergravity in $D = 4$ has been to consider truncations of the potential to sub-manifolds of $E_{7(+7)}/SU(8)$ that are invariant under some postulated residual gauge group $G \subset SO(8)$. As powerful alternative strategies have been shown to exist that allow one to go far beyond what this method can achieve – and in particular have produced numerous solutions that break the $SO(8)$ gauge group to no continuous residual symmetry – independent verification of results becomes a problem due to both the complexity of the scalar potential and the large number of new solutions. This article introduces a conceptually simple self-contained piece of computer code that allows independent numerical validation of claims on the locations of newly discovered stationary points.

1 Introduction
Maximally extended ($\mathcal{N} = 8$) supergravity in four dimensions recently has seen a resurgence of interest, in particular due to speculations about its potential perturbative finiteness [1, 2, 3], as well as the AdS$_4$/CFT$_3$ correspondence in the context of ABJM/BLG theory [4] and also early ideas about an AdS/CMT correspondence [5, 6].

As is well known, $\mathcal{N} = 8$ supergravity allows the promotion of its 28 vector fields to nonabelian gauge fields. This was first demonstrated for the gauge group $SO(8)$ in [7]. As has been shown in [9, 10], there are other, more exotic, options as well; in particular, it is possible here to introduce noncompact $SO(p, 8-p)$.

In order to maintain supersymmetry, such a deformation of the model requires the introduction of a potential on the 70-dimensional manifold of scalars. This potential is known to have a number of nontrivial stationary points, some of which correspond to stable vacuum states with broken symmetry and usually some residual supersymmetry. Unfortunately, the analysis of these potentials is fairly involved, as they are most naturally regarded as functions on the coset manifold $E_{7(+7)}/SU(8)$, which is infeasible to parametrize analytically by introducing coordinates, as is readily seen by simple order-of-magnitude guesstimations on the number of expected combinations of trigonometric and hyperbolic function factors in a generalized Euler angle parametrization.

For a long time, the favored approach to nevertheless extract some information about stationary points was to consider restrictions of the potential to sub-manifolds of $E_{7(+7)}/SU(8)$ that are invariant under some subgroup $G$ of the gauge group, the rationale being that the stationarity condition on the restricted manifold readily is lifted to the full manifold, as the first order term in a perturbative expansion of the potential must be a $G$-invariant scalar. While further technical subtleties such as coordinate singularities might complicate the analysis, this has been shown to be a powerful technique that allowed the determination of all $SU(3)$-invariant stationary points [11], as well as a further stationary point with residual symmetry $SO(3) \times$
SO(3), cf. \cite{12}. No further solutions have been found until a new strategy has been proposed in \cite{13} that increased the number of known nontrivial solutions from six to twenty.

While this method, which is based on numerical techniques that were discovered about the same time \cite{14} as the scalar potentials of gauged maximal supergravity, allows a much deeper analysis and is expected to give rise to numerous other solutions – as it did \cite{15} for $SO(8) \times SO(8)$-gauged Chern-Simons $\mathcal{N} = 16$ supergravity in $D = 3$ \cite{16} – it raises two new issues: First, this is ‘only’ a numerical approach that provides evidence for new stationary points, but no stringent proof of their existence. And second, due to the large number of solutions and the analytic complexity of these potentials, checking these claims unfortunately amounts to a substantial task.

The first issue can be addressed in a semi-automatic way by employing another modern and fairly revolutionary algorithm – the the PSLQ ‘lattice shift’ algorithm to find integer relations between numbers given with high-precision \cite{17}. The basic principle is demonstrated in section 7 of \cite{18} and also briefly explained in section 3. As obtaining a very large number of valid digits is computationally expensive, the cut-off parameters for the PSLQ algorithm will generally require some manual tweaking, so striving for full automatization of this step, while possible, would probably not be a worthwhile objective. The post-processing of analytic conjectures obtained via the PSLQ algorithm, which amounts to establishing their analytic validity, should be fairly straightforward to automatize, however. The challenge is to analytically exponentiate exact expressions for Lie algebra elements using symbolic algebra (probably somewhat tailored to the algebraic task) and then symbolically verifying the stationarity condition.

The second issue is addressed by this article, which provides computer code to numerically validate claims about new stationary points of $SO(8)$-gauged $\mathcal{N} = 8$ supergravity with little effort. As it is expected that many more solutions and maybe also other, even more powerful methods to analyze such scalar potentials could be discovered in the future, having a self-contained dedicated tool for such a task – as is presented in this article – seems to make sense.

2 The Code

Given the main objective of validation of stationary points in the scalar potential of $SO(8)$-gauged $\mathcal{N} = 8$ supergravity, computer code that calculates this potential should be clean, compact, self-contained, and easily verifiable. This is best achieved by an independent re-implementation of numerical code that emphasizes simplicity and verifiability (i.e. a close match between mathematical formulae and code) over algorithmic tricks to achieve high performance and also is independent of the codebase that was used to find the new solutions presented in \cite{13}.

2.1 Design Choices

The problem of numerically computing the supergravity scalar potential – while involving some intricate definitions – only requires arithmetics, a small bit of combinatorics, and as the only tricky step, exponentiation of a complex matrix. Therefore, pretty much any programming language could be used as an implementation basis. While problems of such a nature that do not require symbolic transformations often are addressed with MATLAB \cite{19}, or its free alternative GNU Octave \cite{20}, Python \cite{21} has been chosen for this work, for the following reasons:

\footnote{That codebase is at the time of this writing a complex multi-language mix of LISP and Python modules that mostly deal with problems other than validating results and are not even available on all computing platforms (in particular Microsoft Windows). It will be provided after major clean-up work to make it easy to install also on non-Unix platforms.}
• Python allows a transcription of the required combinatorics to computer code
  with considerably less overhead than many other languages (but not as effec-
  tively as Lisp or Scheme).

• Python is free (in contradistinction to commercial computer algebra packages)
  and readily available across a broad range of hardware and operating system
  platforms.

• Python syntax is very simple and conceptually sufficiently similar to other
  programming languages popular with casual programmers that it can be rea-
  sonably expected to be comprehensible to a broader audience than many other
  languages.

• Python provides a command prompt that allows interactive evaluation of code.

• Python itself has become somewhat popular for many scientific problems.

A major architectural drawback of the code presented here is that, as it de-
liberately is kept algorithmically simple, it neither uses sophisticated combinatorics
nor efficient black-box approaches (such as relational database algorithms for sparse
tensors, as the LambdaTensor \cite{22} package does) to keep the computational effort
 to a minimum. However, for the purpose of validating results, speed is not an
important concern.

2.2 Installation and Usage

The codebase utilizes the Python modules \texttt{numpy} for (non-sparse) tensor numerics
as well as \texttt{scipy} (Scientific Python) for matrix inversion and exponentiation, and
hence requires both Python as well as these extension modules to be installed. Most
free Unix distributions are strongly modularized and provide the corresponding
packages; for other platforms such as Microsoft Windows, convenient pre-bundled
Python packages that contain these modules are available. Installation of Interactive
Python (\texttt{ipython}) also is highly recommended.

Once Python is installed and the source that accompanies this article is down-
loaded from \url{http://arxiv.org/e-print/1007.0600} and unpacked, the functions
described below can be made available by importing the corresponding module into
Python. The most important constant certainly is the dictionary \texttt{vacua_v70} from
\texttt{e7\_vacua.py} – which contains all the known stationary points. The dictionary keys
are strings of the form ‘#0’, ‘#1’, ‘#2’, etc. that correspond to the numbers used
in the tables in \cite{13}. All Python functions come with interactive online help. A
transcript of an interactive Python session is given in figure \ref{fig:ipython).

For some investigations, scripting is preferable to interactive usage. This is most
easily done by adding new Python files to the code directory; alternatively, one can
extend the PYTHONPATH environment variable to tell Python where to look for mod-
ules and put additional code into an arbitrary directory. The file \texttt{e7\_example.py}
which also is listed in figure \ref{fig:example} shows scripting use. It is executed as ‘\texttt{python
e7\_example.py}’.

2.3 Modules and Functions

The code itself is modularized and consists of these components:

2.3.1 The tensor\_io.py module

This component defines functions that read and write numerical (higher-rank) ten-
sors in a well-defined simple sparse textual data format that is also human-readable.
other codebases. It is generally expected that programs which manipulate tensors
might want to save these to files, both for persistent data storage and data exchange.
The full format definition, which may be adopted by other programs, is given in the
$ ipython$

In [1]: from e7_analysis import *
   # (This takes a while, as definitions have to be computed)

In [2]: from e7_vacua import *

In [3]: help(e7_A1_eigenvalues)
   # This brings up online help for this function

In [4]: e7_A1_eigenvalues(vacua_v70 "#3")
Out [4]:
   (array([ 1.33758921 -0.09352799 j, 1.33758921 -0.0935279 9 j ,
   1.09478134+0.00705399 j , 1.33758921 -0.09352799 j, 
   1.33758921 -0.09352799 j , 1.33758921 -0.09352799 j, 
   1.33758921 -0.09352799 j , 1.33758921 -0.09352799 j]),
   [ 1.499987205717096 ,
   1.499987205717116 ,
   1.0000000000469873 ,
   1.499987205717116 ,
   1.499987205717089 ,
   1.499987205717109 ,
   1.499987205717116 ,
   1.499987205717105])

In [5]: e7_test_stationarity_Q(vacua_v70 "SU(4)")
Out [5]: 3.0086481280021352 e -15
   # This shows that the Q-tensor stationarity condition is
   # satisfied to high numerical accuracy for the known
   # stationary point with SU(4) symmetry.

Figure 1: An interactive ipython session with the validation code.

Python documentation of this module. It provides the functions tensor_print(),
tensor_read(), and tensor_write().

2.3.2 The e7_combinators.py module

This component provides a few combinatorics-related definitions that are of use to
other components. In particular, it defines two-index and four-index ranges running
from (0,0) to (7,7) and (0,0,0,0) to (7,7,7,7), respectively. Also, it introduces
iteration over 3-permutations and 4-permutations. Here, it has to be kept in mind
that index counting starts at 0, not 1. Matching published data to numerical results
hence requires shifting all index ranges found in the literature by one.

2.3.3 The e7_definitions.py module

This component provides definitions related to the spin(8) algebra that are then
used to define explicit forms for the 133 complex $56 \times 56 E_7^{(1)}$ generator matrices
of the fundamental representation. The most important definitions are these non-
sparse numpy higher-rank arrays:

- $T_{e7}$ – the $133 \times 56 \times 56$ tensor $T^{(E_7)}_{A\hat{A}B\hat{B}}$ from formula (A.12) in [13].
- $so8_{\gamma\alpha\beta\dot{\alpha}\dot{\beta}}$ – the $8 \times 8 \times 8 \times 8$ tensor $\gamma_{a\dot{a}}$ containing spin(8) ‘gamma matrices’. The order of indices is vector, spinor, co-spinor, as indicated by
the name.
- $so8_{\gamma\alpha\beta\dot{a}\alpha\dot{a}\beta}$, $so8_{\gamma\alpha\beta\dot{a}\alpha\dot{a}\dot{a}}$, $so8_{\gamma\alpha\beta\dot{a}\alpha\dot{a}\dot{a}}$ – likewise, these (non-sparse)
  $8 \times 8 \times 8 \times 8 \times 8 \times 8 \times 8 \times 8$ arrays give the tensors $\gamma^{ab}_{a\beta}$, $\gamma^{ab}_{\alpha\beta}$,
  $\gamma^{abcd}_{a\beta}$, and $\gamma^{abcd}_{\alpha\beta}$.
from e7_vacua import *
from e7_analysis import *

for n in range(21):
    key='#'+str(n)
    v70=vacua_v70[key]
    (gens, svd)=e7_residual_gauge_group_generators(v70)

    print( "Solution:", key, \\
    "P=", e7_potential_from_v70(v70), \\
    "Q=", e7_test_stationarity_Q(v70), \\
    "dim(res. GG)="", len(gens)
    )

Figure 2: Example code demonstrating scripting use of the validation code and its output.
Furthermore, there are functions to map numerical vectors of length 70 that contain $35 + 35$ coefficients of the ‘boost’ generators of $E_7(+7)$ to the language of self-dual and anti-self-dual four-forms and back. Internally, four-form coefficients are represented as Python dictionaries (hash tables). The `format_abcd_to_string()` and `format_abcd_to_latex()` functions can be used to bring these into human-readable form.

The function `abcd_from_v70()` maps a 70-vector to a four-form Python dictionary. Its inverse is given by `v70_from_abcd()`. Normalization conventions are such that if a 70-vector $v$ has a single non-zero entry, then the non-zero entries (192 of them) in the corresponding four-form dictionary are all $\pm 1$.

### 2.3.4 The `e7_potential.py` Module

This component contains the definitions needed to calculate the supergravity scalar potential. The most important function is `e7_potential_from_v70()`, which computes and returns the potential to numerical accuracy. As this function can also return the most important intermediate quantities used in the calculation, such as the tensors $A_1$ and $A_2$, when called with an additional dictionary parameter `info` to store data in, the other auxiliary functions defined here – such as `e7_A12()` – are usually not needed.

Based on the potential function, the function `e7_test_gradient()` uses finite differences to numerically calculate an approximation to the potential’s gradient as well as its length at a given point. This provides a direct test for stationarity and hence allows the numerical validation of solutions.

### 2.3.5 The `e7_vacua.py` Module

This module provides just a single constant, the dictionary `vacua_v70` that contains 70-vectors for all the known stationary points in the scalar potential. Some solutions are associated with named keys, such as ‘G2’, and all of them can be addressed with keys of the form ‘#17’, corresponding to the list given in [13].

This module also provides a function `e7_verify_vacua()` that re-does the automatic gradient check for all known stationary points. This function uses the function `e7_test_gradient()` from the `e7_potential.py` module.

### 2.3.6 The `e7_analysis.py` Module

This module provides additional functions to analyze the properties of stationary points. The most important ones are:

- `e7_test_stationarity_Q()` – to verify whether a point satisfies the $Q_{abcd} + \frac{1}{24} g_{abcdefgh} Q_{efgh} = 0$ stationarity condition. This provides an independent test in addition to the direct numerical evaluation of the gradient provided by `e7_test_gradient()`.

- `e7_residual_gauge_group_generators()` – to automatically determine a basis of the subgroup of spin(8) that is left unbroken by the point in question.

- `e7_susy_A2()` – to numerically determine the number of unbroken supersymmetries via the $\epsilon^i A_2^{ijkl} = 0$ condition.

- `e7_A1_eigenvalues()` – to numerically determine the eigenvalues of the $A_1$ tensor. In addition to these eigenvalues $E$, the values $-6 |E|^2 / P$ are returned as well. This gives a second test for unbroken supersymmetry.

- `e7_scalar_massmatrix()` – to numerically calculate the re-scaled scalar mass matrix given by formula (2.25) in [8]:

$$96e^{-1} g^{-2} L_M = -\left(\frac{2}{3} P(V) + \frac{13}{72} A_2^{mnpq} A_2^{mnpq}\right) \sum_{ijkl} \Sigma^{ijkl}$$
\[- \left( 6A_{2k}^{mn}A_{2j}^{mn\ell} - \frac{3}{2}A_{2n}^{mij}A_{2n}^{mk\ell} \right) \Sigma_{ijpq}\Sigma_{k\ell pq} \\
+ \frac{2}{3}A_{2j}^{mn\epsilon}A_{2q}^{j\epsilon\ell} \Sigma_{mn\epsilon q}\Sigma_{j\epsilon\ell}. \]

One normally will want to re-scale this matrix by multiplying with $-6/P$ ($P$ being the value of the potential) in order to obtain masses in AdS-units. After such re-scaling, the Breitenlohner-Freedman stability bound [24] is $-(d-1)^2/4 = -9/4 = -2.25$.

- $e7\mathcal{A12.masses2()}$ – to calculate the fermion masses-squared according to formula (2.14) in [8].

### 3 Outlook

Deeper numerical investigations than those presented in [13] suggest that the number of yet unpublished stationary points in the scalar potential studied here as well as related ones may be fairly high. A number of these solutions may easily turn out to be sufficiently interesting to warrant a deeper investigation. One might think that the idea of using numerical techniques to find approximate solutions should be of very limited use in particular for those solutions that break the gauge group (almost) completely: Studying a stationary point numerically in order to get ideas that may guide a fully analytic approach seems to be a fairly unrewarding exercise unless it has a sufficient degree of residual symmetry. However, this is not necessarily the case. First of all, it is possible to eliminate almost all of the arbitrariness in the raw numbers that is associated with the freedom to apply a spin(8)-rotation to a given solution, using the techniques that were employed in [15, 13] to minimize the number of non-zero coefficient vector entries. There is no fundamental obstacle to modifying readily existing techniques to use very high precision arithmetics (with hundreds of digits) rather than machine-precision 16-digit arithmetics in order to determine parameters very precisely. This then allows one to utilize powerful integer relation algorithms such as in particular PSLQ [17] to systematically ‘guess’ even very complicated analytic expressions. To demonstrate this, let us specifically consider a number such as the coordinate $z(2)$ from table A.1 in [6]: this is given as:

\[
z(2) = \frac{1}{4} \left( 1 - \frac{i}{3^{3/4}} \sqrt{2 + \sqrt{3}} \right) (3 + \sqrt{3} - 3^{3/4} \sqrt{10}) \tag{1} \]

To 300 digits, the imaginary part of $z(2)$ is:

\[
\text{Im } z(2) \approx -0.209269477042954393209112597384263059035631424 \\
839824728046049646351961493949022694957040180268 \\
46085412921545142577924797777536023519697290138 \\
356018555716825868238074107476584372107488197263 \tag{2} \\
11336237435710197764338927716172598148291052464 \\
018329383941096143123970705213689657726314686822 \\
409336284550127068
\]

If we truncate this number to 200 digits and use the PSLQ algorithm to then find a vector of integer coefficients for integer powers of this number so that the sum is closer to zero than $10^{-170}$, the PSLQ algorithm produces the following polynomial coefficients:

\[
[-64, 0, 704, 0, -240, 0, 32, 0, -1] \tag{3}
\]
One could say that this analytic expression allows us to make a testable 'prediction' for the next 100 digits. In fact, using Newton's algorithm to find the zero of interest of this polynomial reproduces Im $\zeta(2)$ to 100 digits beyond the previous 200. This is demonstrated in detail the example code file e7\_example\_mpmath.py. In order to run this piece of code, the 'mpmath' Python extension \cite{23} has to be installed.

It seems hence very likely that even for new stationary points that can be expected to require even more complicated analytic expressions, obtaining – and most likely even rigorously proving – analytic results can be fully mechanized.

The codebase that has been presented in this work (a) correctly reproduces the known properties of the ‘classical’ non-trivial stationary points, (b) should be simple enough to be readily validated against the formulae given in \cite{13}, and (c) should in the future help to considerably reduce the effort required to check claims about the properties of new stationary points.

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