

Annealing code minAone User Guide

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1 About minAone

The annealing code `minAone` described in this document is used for calculating action levels of dynamical systems. The code is developed as an extension of `minAzero` written by Bryan Toth and Chris Knowlton. That is where the name `minAone` comes from. In another aspect, following the lowest action level A_0 , A_1 represents the second lowest one, which is an interesting quantity we care about and has significant application in statistical data assimilation.

2 Problem Statement

Given a dynamical system modeled by D -dimensional discrete map

$$x_a(n+1) = f_a(\mathbf{x}(n)), \quad a = 1, \dots, D$$

the probability distribution of its states can be expressed as $P(X|Y) = \exp(-A_0)$, when L -dimensional observations Y are present. If one assumes both measurement noises and model error are independent and gaussian, the action A_0 has the format of

$$A_0(X) = \sum_{n=0}^m \frac{R_m(n)}{2} \sum_{l=1}^L [x_l(n) - y_l(n)]^2 + \frac{R_f}{2} \sum_{n=0}^{m-1} \sum_{a=1}^D [x_a(n+1) - f_a(\mathbf{x}(n))]^2. \quad (1)$$

where R_m and R_f are the inverse of variances.

The annealing method is based on the observation that the minima solution X^q of A_0 at $R_f = 0$ is $x_l(n) = y_l(n)$, the other $D - L$ components of the model state vector are undetermined, and the solution is degenerate. As we increase R_f , the action levels split, and depending on R_m , R_f , L and the precise form of the dynamical vector field $\mathbf{f}(\mathbf{x})$, there will be 1, 2, ... minima of A_0 .

3 Annealing Procedure

The annealing process proceeds as follows: with very small initial R_f , we call it R_{f0} , solve the $(m+1)D$ -dimensional search problem with an optimization algorithm that seeks minima of $A_0(X)$. Start the search with a set of trial paths whose components are selected from a uniform distribution within limits suggested by examining the times series generated by the

model $\mathbf{x} \rightarrow \mathbf{f}(\mathbf{x})$ (or any other selection process for the initial guess). This will generate a collection of approximate paths X^q . Increase R_f by a small increment (we choose $R_f = \{R_{f0}\alpha^\beta\}$, where $\alpha = 2, \beta = 0, 1, \dots$ in our examples), and using the paths found for the smaller R_f as initial guesses, find a new set of approximate X^q . Continue this process until the lowest action level path X^0 produces a $A_0(X^0)$ near expected value, which can be identified from our knowledge of measurement noises. In our example, as the values $[y_l(n) - x_l(n)] \sim \mathcal{N}(0, \sigma^2)$ by our choice, the measurement error term $\sum_{n=0}^m \sum_{l=1}^L [(x_l(n) - y_l(n))/\sigma]^2/2$ has a χ^2 distribution with $L(m+1)$ degrees of freedom. The mean and uncertainty of this distribution over different choices of noise waveforms are $(m+1)L/2$ and $\sqrt{(m+1)L/2}$, respectively.

After identifying the global minima and other local minima of A_0 , we can employ laplace method to approximate the expected value $\langle G(X) \rangle$ of a function $G(X)$ is

$$\langle G(X) \rangle = \frac{\int dX G(X) \exp[-A_0(X)]}{\int dX \exp[-A_0(X)]} \approx G(X^0). \quad (2)$$

plus exponentially small corrections. If the action level $A_0(X^0)$ is substantially less than the action level on the next path $A_0(X^0) \ll A_0(X^1)$, all statistical data assimilation expected values $\langle G(X) \rangle$ are given by X^0 and fluctuations about that path with exponential accuracy of order $\exp[-(A_0(X^1) - A_0(X^0))]$.

4 Installing Required Programs and Packages

This document will assume that the user is using a Linux distribution and has basic compilers installed including gcc, gfortran and python.

4.1 Python Packages

These python scripts link to the sympy library. To install these, use `apt-get/yum install sympy` or download directly from sympy.org.

4.2 IPOPT

Download

Get it here: <https://projects.coin-or.org/Ipopt>

- Download and unzip latest version of IPOPT

- As of right now this is 3.11.7 - Efficacy of installation instructions may degrade over time as packages are updated.
- Go into ThirdParty folder in the IPOPT directory then do the following commands.

```
$ cd Blas
$ ./get.Blas
$ cd ../Lapack
$ ./get.Lapack
$ cd ../ASL
$ ./get.ASL
$ cd ../Metis
$ ./get.Metis
```

- Get the HSL subroutines from <http://hsl.rl.ac.uk/ipopt>
- Note that there are two releases for HSL - you will want the more complete one that contains ma57, ma77, and ma97.
- While the freely available ma27 will work for many problems, the newer packages are faster, work on larger problems, and can use multi-core architecture.
- This will require filling out a form stating essentially that you are in academia and waiting a couple hours for a link to download.
- Unpack the resulting library into the ThirdParty folder such that the path is (IPOPT Path)/ThirdParty/HSL/coinhsl

Install

- Go to the IPOPT directory

```
$ mkdir build
$ cd build
$ ../configure
```

- Note that if you have lapack or blas installed previously you can use `-with-lapack` and `-with-blas` to link to those packages
- If something goes wrong refer here <http://www.coin-or.org/Ipopt/documentation/node19.html#ExpertInstall>

- Assuming everything worked:

```
$ make
$ make test
$ make install
```

5 minAone.py Description

minAone is a python script used to write C++ code and compiler instructions using the IPOPT (Interior Point OPTimization) libraries to estimate unmeasured states and parameters in dynamical systems with limited measurements. The scripts take a set of differential equations and state and parameter names provided by a text file "equations.txt" and returns a set of C++ files consisting of a set of constraints based on a discretized version of those differential equations. A second text file 'specs.txt' allows for changes in run specific quantities state and parameter bounds, as well as input files without the need to recompile.

List of Files

- discAone.py
-Discretizes equations and creates strings for Jacobian and Hessian Elements.
- makecppAone.py
-Writes C++ file linking to IPOPT libraries using strings from discAone.py
- makehppAone.py
-Writes header file for above
- makemakeAone.py
-Writes makefile for problem. Will need to be changed based on install location of IPOPT
- makeoptAone.py
-Writes settings file for IPOPT

These files can be put in /usr/local/sbin for ease of use

Modify makemakeAone.py

The Makefile compiles C ++ object files and links them with the installed IPOPT libraries, in order to create an executable. Since the location of the IPOPT libraries, as well as the flags used to compile them, differ between installations, this file will be unique to a given machine. Modification of the makemake.py script to give correct Makefiles for a given machine consists of:

- Ensure that the IPOPT installation proceeded correctly, as evidenced by zero errors for the make install step.
- In the IPOPT build directory, try to compile (make) one of the examples, for instance at /build/Ipopt/examples/hs071.cpp.
- If this compiles and runs correctly, open the Makefile in this directory.
- Make note of the entries in the following fields of this Makefile: CXX, CXXFLAGS, CXXLINK-FLAGS, INCL, LIBS.
- In makemake.py, replace the default entries for these fields with those given in the example Makefile.
 - makemakeAone.py is formatted differently than a Makefile, since it is a python code generation script.
 - Lines that begin with the # sign will be comments in the Makefile - leave these alone.
 - All lines must end with \n\ in order for the Makefile to be generated correctly.
 - The best way to ensure that all the compile flags are correct is to copy and paste from the example Makefile, ensuring that the end line characters are in place.
- The modification of makemake.py must only be done once for a given machine, unless IPOPT is reinstalled for whatever reason.

6 Running the Code

minAone uses two text documents (along with any needed data files) as input, `equations.txt` and `specs.txt`. Once these are filled

`equations.txt` contains information on the model and is used once for generating the needed `cpp` and `hpp` files for the run. The file should be written as described below in this order.

- The first line is the problem name, this name will be used to name the resulting executable.
- The second line tells minAzero how many dynamical variables, parameters, coupling terms, stimuli, functions, and measurements there are, in that order as a comma delimited list. It is essential that these numbers are accurate as minAzero uses this to know how many lines to read for each component of the code.
- A list of every differential equation.
- The measurement term of the cost function. A penalty term for coupling terms is suggested as any coupling to measurements is not present in physical systems.
- The names of all the variables. These must be the same as used in the differential equations and should be multiple letters/and or numbers such that variable name is contained in any other name or common function.
- The names of parameters, names of couplings, names of data, and names of stimuli, in that order. Again use fully unique names.
- Function names and number of arguments of that function separated by a comma. Use a function if there is some component of the dynamics with a removable singularity or other difficult numerical object that requires an alternative local definition.
- Functions will require an additional file `'myfunctions.cpp'` containing the function definition along with its jacobian and hessian (an example of this is included)

specs.txt contains run specific information such as file names, variable bounds, and problem length. This file can be edited without recompiling the code.

- First line is the number of full steps the code will use. Because the code is compiled using a midpoint method, the actual problem length will double this plus one.
- Second line is the number of lines in each input file to skip. This allows for the code to start at any point in a long data set.
- Third line is double the time step of the data. Again since a midpoint method is used, the time step is for a whole step - which includes two points.
- If you wish to start at a non constant guess, you can put a 1 followed by a line with an initial condition file. This file should have one column for each state. If you do not want to include an initial condition file, use 0
- One line for each of the measured data file names. Each file should be a single column.
- One line for each of the stimulus data file names. Each file should be a single column.
- For each variable, the lower bound, upper bound, and RF0 value separated by commas.
- For each parameter a lower bound, upper bound
- One line for annealing settings, alpha, incresement of beta and maximum beta separated by commas

Once everything is filled out and all data files are present, you can run the python scripts:

```
$ minAone.py
$ make
$ ./(problem_name)_cpp
```

If data files are missing or too short, the code will segfault. The output file contains annealing result for one path named like **D5_M1_PATH0.dat**. Each line of **D5_M1_PATH0.dat** contains the optimal path at different values of

beta. The first three numbers are beta exitflag and action value, respectively. Exitflag can be 0 or 1. 1 means IPopt routines find the optimal path and 0 means it fails. The rest numbers represent the optimal path.

```
beta exitflag action_value
optimal_path[x1(0) x2(0) x3(0) x4(0) x5(0) x1(1) x2(1) x3(1)
x4(1) x5(1) ... x1(NT) x2(NT) x3(NT) x4(NT) x5(NT)
p(1) p(2) ... p(NP)]
```

7 Run in Parallel

One excute `(problem_name)_cpp` can obtain the result for only one random initial path. To explore the landscape of action A_0 , we need to start from different random paths and each of them will converge to different local minima. Since all those paths are independent from each other, it is easy to implement the calculation in parallel using array job.

Here we give a example submission scripts on ccom-boom cluster

```
#!/bin/bash
#$ -t 1-100
#$ -N job_name
#$ -cwd
#$ -j y
#$ -M your@email.com
#$ -S /bin/bash
#$ -m beas
#$ -o ./output
#$ -e ./error
#$ -q batch.q
./problem_name_cpp $SGE_TASK_ID
```

Each path will be stored in individual file with the name like `D5_M1_PATH0.dat`, `D5_M1_PATH2.dat`, ..., `D5_M1_PATH100.dat`.

8 Examples

Two examples are provided: the first one is Lorenz96 D=5 to show the basic settings of `equations.txt` and `specs.txt`. And the other example is NaKL to show how to include external stimuli in `equations.txt` and `specs.txt`.

8.1 Lorenz96 D=5

Lorenz96 D=5 Vector Field

$$\begin{aligned}\frac{dx_1}{dt} &= x_5(x_2 - x_4) - x_1 + f \\ \frac{dx_2}{dt} &= x_1(x_3 - x_5) - x_2 + f \\ \frac{dx_3}{dt} &= x_2(x_4 - x_1) - x_3 + f \\ \frac{dx_4}{dt} &= x_3(x_5 - x_2) - x_4 + f \\ \frac{dx_5}{dt} &= x_4(x_1 - x_3) - x_5 + f\end{aligned}$$

Lorenz96 D=5 equations.txt

```
# Problem Name
lorenz96
# nY,nP,nU,nI,nF,nM
5,1,0,0,0,1
# equations
yy5*(yy2-yy4)-yy1+FF1
yy1*(yy3-yy5)-yy2+FF1
yy2*(yy4-yy1)-yy3+FF1
yy3*(yy5-yy2)-yy4+FF1
yy4*(yy1-yy3)-yy5+FF1
# Objective/Cost function
4*(data1-yy1)*(data1-yy1)
# variable names
yy1
yy2
yy3
yy4
yy5
# parameter names
FF1
# data names
data1
# stimuli names
```

Lorenz96 D=5 specs.txt

```

# Includes the problem length
80
# How much data to skip
# In case you do not want to start at the beginning of the data file
100
# Time step - this is twice the time step of the data,
# since the data includes time and midpoints.
0.02
# Data File names - input
x1.dat
# Data File name - stimuli
# No stimuli for this problem
# Boundary & initial conditions
# 0 for no initial data file, 1 for data file
# A data file must include values for all state variables
# at each time point.
0
# If above is 1, list name of data file next. If 0, no entry needed.
# State Variables:
# These are in the formats: lower bound, upper bound, Rf0
# y1
-15, 15, 0.01
# y2
-15, 15, 0.01
# y3
-15, 15, 0.01
# y4
-15, 15, 0.01
# y5
-15, 15, 0.01
# Parameters:
0, 20, 8.17
#annealing setting:  $R_f = R_{f0} \cdot \alpha^{\beta}$ 
#There are in the formats: alpha, incresement of beta, maximum beta
# here we have alpha=2, beta = 0, 1, 2, 3, ..., 29
2,1,30

```

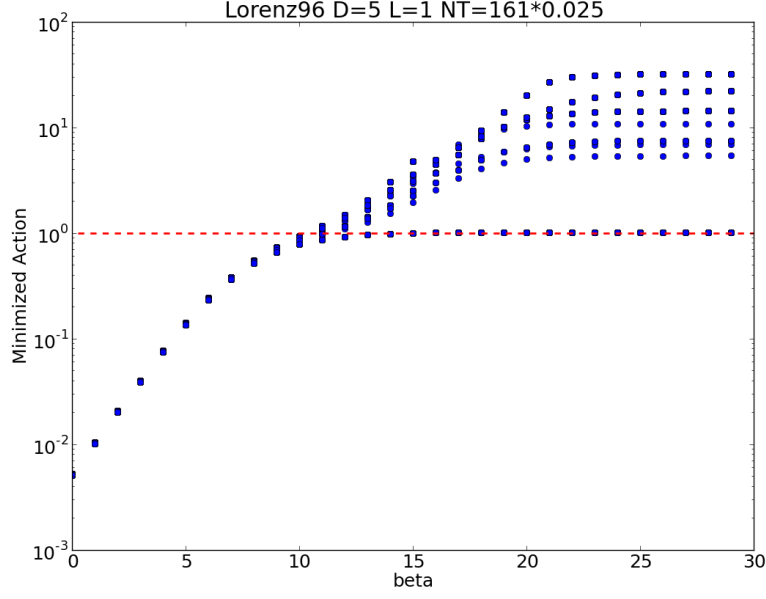


Figure 1: Lorenz 96 D=5 L=1 Action Level

8.2 NaKL

NaKL Vector Field

$$\frac{dV}{dt} = C I_{inj}(t) + g_{Na} m^3 h (E_{Na} - V) + g_K n^4 (E_K - V) + g_L (E_L - V)$$

$$\frac{da}{dt} = \frac{a_{\infty} - a}{\tau_a}, \quad a = \{m, h, n\}$$

$$a_{\infty} = \frac{1}{2} + \frac{1}{2} \tanh\left(\frac{V - V_a}{\Delta V_a}\right)$$

$$\tau_a = \tau_{a0} + \tau_{a0} \left(1 - \tanh^2\left(\frac{V - V_a}{\Delta V_a}\right)\right)$$

NaKL equations.txt

```
simple_nakl
```

```
# nY,nP,nU,nI,nF,nM
```

```
4,19,0,1,0,1
```

```
#vector field
```

```
gNa*(m0*m0*m0*h0)*(ENa-V0)+gK*n0*n0*n0*n0*(EK-V0)+gL*(EL-V0)+Area*Iinj
```

```
(0.5*(1+tanh((V0-Vmo)*dVm)) - m0)/(Cm1+Cm2*(1.0-tanh((V0-Vmo)*dVm)*tanh((V0-Vmo)*dVm))
```

```

(0.5*(1+tanh((V0-Vho)*dVh)) - h0)/(Ch1+Ch2*(1.0-tanh((V0-Vho)*dVh)*tanh((V0-Vho)*dVh))
(0.5*(1+tanh((V0-Vno)*dVn)) - n0)/(Cn1+Cn2*(1.0-tanh((V0-Vno)*dVn)*tanh((V0-Vno)*dVn))
#obj function
(VDATA0 - V0)*(VDATA0 - V0)
#states
V0
m0
h0
n0
#parameters
gNa
ENa
gK
EK
gL
EL
Area
Vmo
dVm
Cm1
Cm2
Vho
dVh
Ch1
Ch2
Vno
dVn
Cn1
Cn2
#data names
VDATA0
#stimuli
Iinj

NaKL specs.txt

3000
0
0.04
#data

```

```

./noise_measured.dat
#stimuli
./current.dat
0
#./allstates.dat
# state bounds and Rf0
-150,70,1e-3
0, 1,1e1
0, 1,1e1
0, 1,1e1
# parameter bounds
#gna
50,200,100,120
#Ena
0,100,50,50
#gki
5,40,30,20
#Ek
-100,-50,-70,-77
#gl
0.1,1,.2,.3
#El
-60,-50,-52,-54
#Area
0.5,1.5,1,0.8
#mv1
-60,-30,-45,-40
#mv2
.01,0.1,.075,0.06667
#cm1
0.05,.25,.15,.1
#cm2
.1,1,.4,.4
#hv1
-70,-40,-50,-60
#hv2
-0.1,-.01,-.05,-.06667
#ch1
.1,5,1.2,1
#ch2

```

```

1,15,6,7
#nv1
-70,-40,-52,-55
#nv2
.01,0.1,.03,.03333
#cn1
.1,5,.8,1
#cn2
2,12,5,5
#anneal settings
2,1,30

```

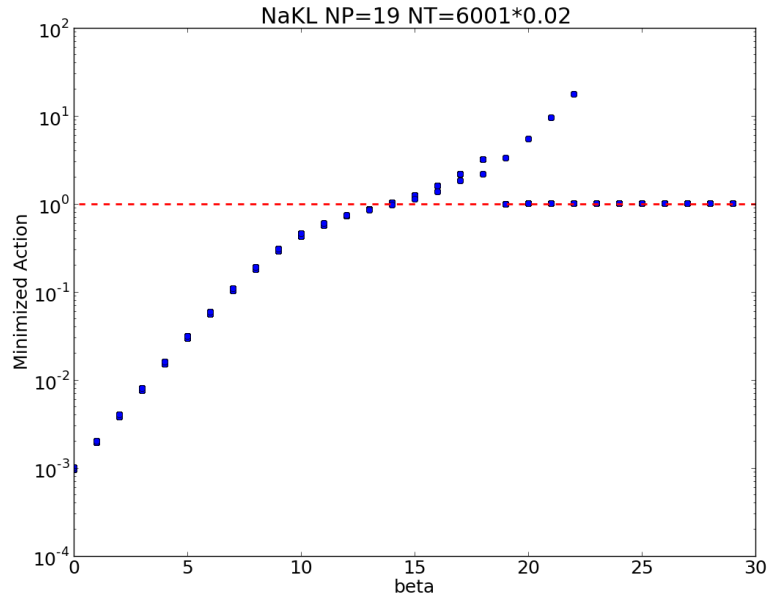


Figure 2: NaKL Action Level

9 Troubleshooting

I have tested these scripts over a wide range of problems, so I believe that the algorithms are correct. However, there are a few common errors that may crop up.

- Variable and parameter naming is very important. A few common problems can crop up. Never use a variable name that includes the

name of another variable. For instance p1 and p11 would be bad, since p11 includes p1. In this case, p01 and p11 would be adequate. Along this vein, all variable names should be at least 2 characters long, just in case.