An Intro to Differential Equations.jl

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0.1 Basic Introduction Via Ordinary Differential Equations

This notebook will get you started with Differential Equations.jl by introducing you to the functionality for solving ordinary differential equations (ODEs). The corresponding documentation page is the ODE tutorial. While some of the syntax may be different for other types of equations, the same general principles hold in each case. Our goal is to give a gentle and thorough introduction that highlights these principles in a way that will help you generalize what you have learned.

0.1.1 Background

If you are new to the study of differential equations, it can be helpful to do a quick background read on the definition of ordinary differential equations. We define an ordinary differential equation as an equation which describes the way that a variable u changes, that is

$$u' = f(u, p, t) \tag{1}$$

where p are the parameters of the model, t is the time variable, and f is the nonlinear model of how u changes. The initial value problem also includes the information about the starting value:

$$u(t_0) = u_0 \tag{2}$$

Together, if you know the starting value and you know how the value will change with time, then you know what the value will be at any time point in the future. This is the intuitive definition of a differential equation.

0.1.2 First Model: Exponential Growth

Our first model will be the canonical exponential growth model. This model says that the rate of change is proportional to the current value, and is this:

$$u' = au \tag{3}$$

where we have a starting value $u(0) = u_0$. Let's say we put 1 dollar into Bitcoin which is increasing at a rate of 98% per year. Then calling now t = 0 and measuring time in years, our model is:

$$u' = 0.98u \tag{4}$$

and u(0) = 1.0. We encode this into Julia by noticing that, in this setup, we match the general form when

```
f(u,p,t) = 0.98u
```

with $u_0 = 1.0$ \$. If we want to solve this model on a time span from t=0.0 to t=1.0, then we define an ODEProblem by specifying this function f, this initial condition u0, and this time span as follows:

```
using DifferentialEquations
f(u,p,t) = 0.98u
u0 = 1.0
tspan = (0.0,1.0)
prob = ODEProblem(f,u0,tspan)
```

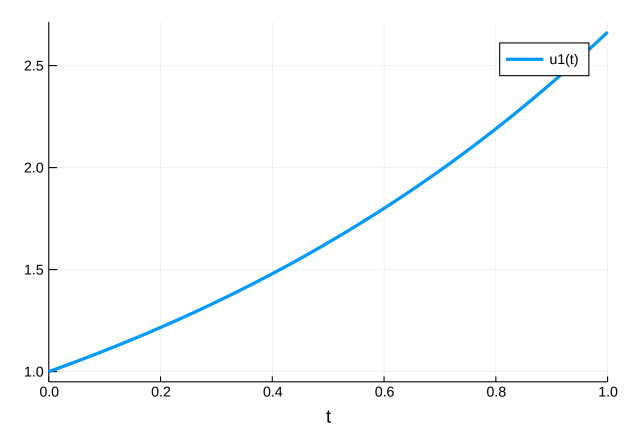
To solve our ODEProblem we use the command solve.

```
sol = solve(prob)
```

and that's it: we have succesfully solved our first ODE!

Analyzing the Solution Of course, the solution type is not interesting in and of itself. We want to understand the solution! The documentation page which explains in detail the functions for analyzing the solution is the Solution Handling page. Here we will describe some of the basics. You can plot the solution using the plot recipe provided by Plots.jl:

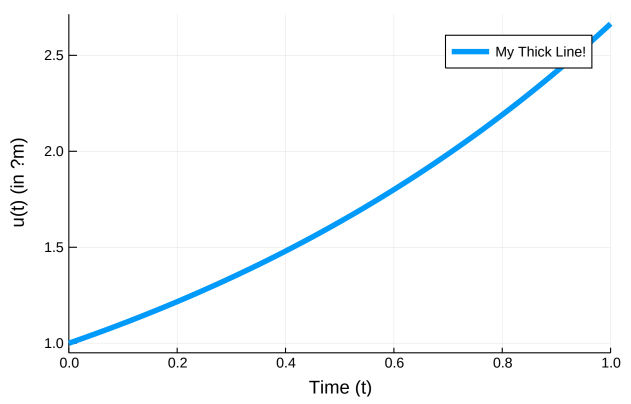
```
using Plots; gr()
plot(sol)
```



From the picture we see that the solution is an exponential curve, which matches our intuition. As a plot recipe, we can annotate the result using any of the Plots.jl attributes. For example:

```
plot(sol,linewidth=5,title="Solution to the linear ODE with a thick line", xaxis="Time (t)",yaxis="u(t) (in \mum)",label="My Thick Line!") # legend=false
```

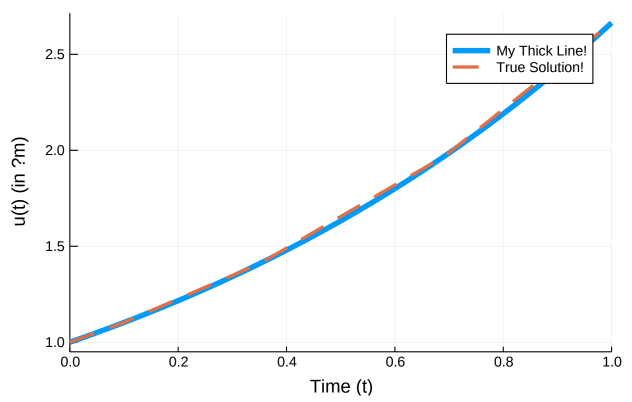
Solution to the linear ODE with a thick line



Using the mutating plot! command we can add other pieces to our plot. For this ODE we know that the true solution is $u(t) = u_0 exp(at)$, so let's add some of the true solution to our plot:

plot!(sol.t, t->1.0*exp(0.98t), lw=3, ls=:dash, label="True Solution!")





In the previous command I demonstrated sol.t, which grabs the array of time points that the solution was saved at:

sol.t

5-element Array{Float64,1}:

- 0.0
- 0.10042494449239292
- 0.35218555997054785
- 0.6934428593452983
- 1.0

We can get the array of solution values using sol.u:

sol.u

5-element Array{Float64,1}:

- 1.0
- 1.1034222047865465
- 1.4121902211481592
- 1.9730369899955797
- 2.664456142481388

sol.u[i] is the value of the solution at time sol.t[i]. We can compute arrays of functions of the solution values using standard comprehensions, like:

```
[t+u for (u,t) in tuples(sol)]

5-element Array{Float64,1}:
1.0
1.2038471492789395
1.764375781118707
2.666479849340878
3.664456142481388
```

However, one interesting feature is that, by default, the solution is a continuous function. If we check the print out again:

sol

```
retcode: Success
Interpolation: Automatic order switching interpolation
t: 5-element Array{Float64,1}:
0.0
0.10042494449239292
0.35218555997054785
0.6934428593452983
1.0
u: 5-element Array{Float64,1}:
1.0
1.1034222047865465
1.4121902211481592
1.9730369899955797
2.664456142481388
```

you see that it says that the solution has a order changing interpolation. The default algorithm automatically switches between methods in order to handle all types of problems. For non-stiff equations (like the one we are solving), it is a continuous function of 4th order accuracy. We can call the solution as a function of time sol(t). For example, to get the value at t=0.45, we can use the command:

```
sol(0.45)
```

1.554261048052598

Controlling the Solver Differential Equations.jl has a common set of solver controls among its algorithms which can be found at the Common Solver Options page. We will detail some of the most widely used options.

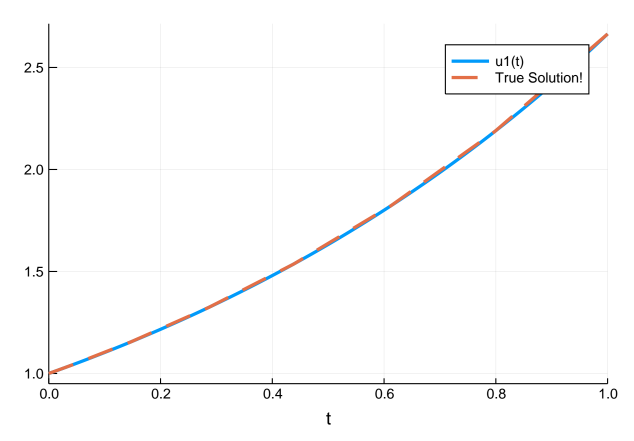
The most useful options are the tolerances abstol and reltol. These tell the internal adaptive time stepping engine how precise of a solution you want. Generally, reltol is the relative accuracy while abstol is the accuracy when u is near zero. These tolerances are local tolerances and thus are not global guarantees. However, a good rule of thumb is that the total solution accuracy is 1-2 digits less than the relative tolerances. Thus for the defaults

abstol=1e-6 and reltol=1e-3, you can expect a global accuracy of about 1-2 digits. If we want to get around 6 digits of accuracy, we can use the commands:

```
sol = solve(prob,abstol=1e-8,reltol=1e-8)
```

Now we can see no visible difference against the true solution:

```
plot(sol)
plot!(sol.t, t->1.0*exp(0.98t),lw=3,ls=:dash,label="True Solution!")
```



Notice that by decreasing the tolerance, the number of steps the solver had to take was 9 instead of the previous 5. There is a trade off between accuracy and speed, and it is up to you to determine what is the right balance for your problem.

Another common option is to use saveat to make the solver save at specific time points. For example, if we want the solution at an even grid of t=0.1k for integers k, we would use the command:

```
sol = solve(prob,saveat=0.1)
```

Notice that when saveat is used the continuous output variables are no longer saved and thus sol(t), the interpolation, is only first order. We can save at an uneven grid of points by passing a collection of values to saveat. For example:

```
sol = solve(prob, saveat=[0.2, 0.7, 0.9])
```

If we need to reduce the amount of saving, we can also turn off the continuous output directly via dense=false:

```
sol = solve(prob,dense=false)
```

and to turn off all intermediate saving we can use save everystep=false:

```
sol = solve(prob, save_everystep=false)
```

If we want to solve and only save the final value, we can even set save_start=false.

```
sol = solve(prob, saveat=[0.2,0.7,0.9], save_start = false)
```

Note that similarly on the other side there is save end=false.

More advanced saving behaviors, such as saving functionals of the solution, are handled via the SavingCallback in the Callback Library which will be addressed later in the tutorial.

Choosing Solver Algorithms There is no best algorithm for numerically solving a differential equation. When you call <code>solve(prob)</code>, DifferentialEquations.jl makes a guess at a good algorithm for your problem, given the properties that you ask for (the tolerances, the saving information, etc.). However, in many cases you may want more direct control. A later notebook will help introduce the various algorithms in DifferentialEquations.jl, but for now let's introduce the <code>syntax</code>.

The most crucial determining factor in choosing a numerical method is the stiffness of the model. Stiffness is roughly characterized by a Jacobian f with large eigenvalues. That's quite mathematical, and we can think of it more intuitively: if you have big numbers in f (like parameters of order 1e5), then it's probably stiff. Or, as the creator of the MATLAB ODE Suite, Lawrence Shampine, likes to define it, if the standard algorithms are slow, then it's stiff. We will go into more depth about diagnosing stiffness in a later tutorial, but for now note that if you believe your model may be stiff, you can hint this to the algorithm chooser via alg_hints = [:stiff].

```
sol = solve(prob,alg_hints=[:stiff])
```

Stiff algorithms have to solve implicit equations and linear systems at each step so they should only be used when required.

If we want to choose an algorithm directly, you can pass the algorithm type after the problem as solve(prob,alg). For example, let's solve this problem using the Tsit5() algorithm, and just for show let's change the relative tolerance to 1e-6 at the same time:

```
sol = solve(prob,Tsit5(),reltol=1e-6)
```

0.1.3Systems of ODEs: The Lorenz Equation

Now let's move to a system of ODEs. The Lorenz equation is the famous "butterfly attractor" that spawned chaos theory. It is defined by the system of ODEs:

$$\frac{dx}{dt} = \sigma(y - x) \tag{5}$$

$$\frac{dy}{dt} = x(\rho - z) - y \tag{6}$$

$$\frac{dx}{dt} = \sigma(y - x) \tag{5}$$

$$\frac{dy}{dt} = x(\rho - z) - y \tag{6}$$

$$\frac{dz}{dt} = xy - \beta z \tag{7}$$

To define a system of differential equations in Differential Equations.jl, we define our f as a vector function with a vector initial condition. Thus, for the vector $\mathbf{u} = [x,y,z]$, we have the derivative function:

```
function lorenz! (du,u,p,t)
    \sigma, \rho, \beta = p
    du[1] = \sigma*(u[2]-u[1])
    du[2] = u[1]*(\rho-u[3]) - u[2]
    du[3] = u[1]*u[2] - \beta*u[3]
end
```

lorenz! (generic function with 1 method)

Notice here we used the in-place format which writes the output to the preallocated vector du. For systems of equations the in-place format is faster. We use the initial condition $u_0 = [1.0, 0.0, 0.0]$ as follows:

```
u0 = [1.0, 0.0, 0.0]
```

Lastly, for this model we made use of the parameters p. We need to set this value in the ODEProblem as well. For our model we want to solve using the parameters $\sigma = 10$, $\rho = 28$, and $\beta = 8/3$, and thus we build the parameter collection:

```
p = (10,28,8/3) # we could also make this an array, or any other type!
```

Now we generate the ODEProblem type. In this case, since we have parameters, we add the parameter values to the end of the constructor call. Let's solve this on a time span of t=0 to t=100:

```
tspan = (0.0, 100.0)
prob = ODEProblem(lorenz!,u0,tspan,p)
```

Now, just as before, we solve the problem:

```
sol = solve(prob)
```

The same solution handling features apply to this case. Thus sol.t stores the time points and sol.u is an array storing the solution at the corresponding time points.

However, there are a few extra features which are good to know when dealing with systems of equations. First of all, sol also acts like an array. sol[i] returns the solution at the ith time point.

```
sol.t[10],sol[10]
(0.0836852441654334, [1.08886, 2.05232, 0.0740254])
```

Additionally, the solution acts like a matrix where sol[j,i] is the value of the jth variable at time i:

```
sol[2,10]
```

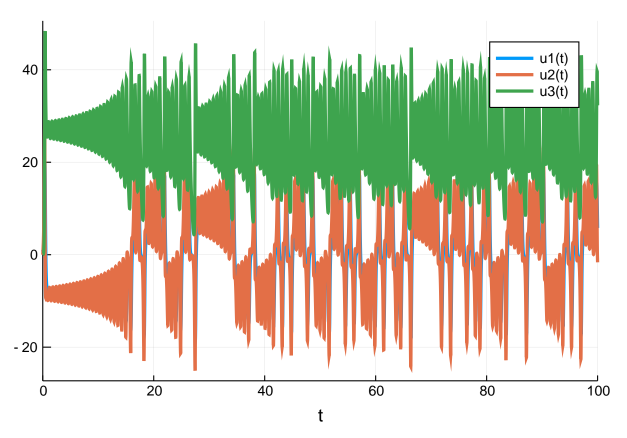
2.052321820923891

We can get a real matrix by performing a conversion:

```
A = Array(sol)
```

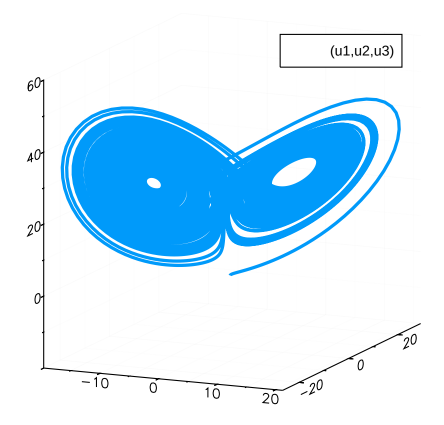
This is the same as sol, i.e. sol[i,j] = A[i,j], but now it's a true matrix. Plotting will by default show the time series for each variable:

```
plot(sol)
```



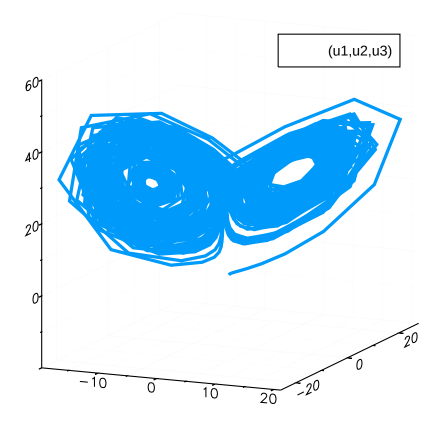
If we instead want to plot values against each other, we can use the vars command. Let's plot variable 1 against variable 2 against variable 3:

plot(sol, vars=(1,2,3))



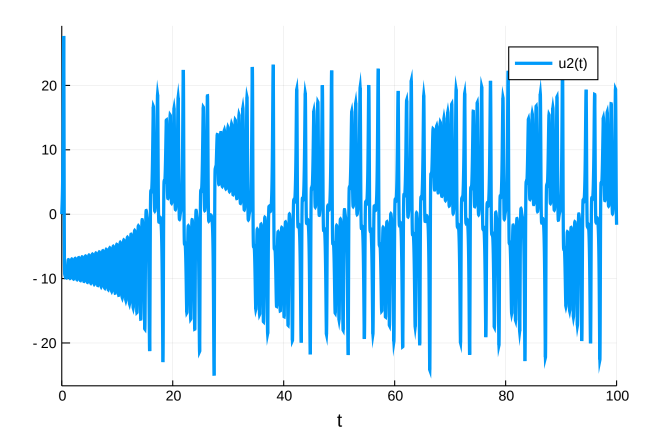
This is the classic Lorenz attractor plot, where the x axis is u[1], the y axis is u[2], and the z axis is u[3]. Note that the plot recipe by default uses the interpolation, but we can turn this off:

plot(sol, vars=(1,2,3), denseplot=false)



Yikes! This shows how calculating the continuous solution has saved a lot of computational effort by computing only a sparse solution and filling in the values! Note that in vars, O=time, and thus we can plot the time series of a single component like:

```
plot(sol,vars=(0,2))
```



0.1.4A DSL for Parameterized Functions

In many cases you may be defining a lot of functions with parameters. There exists the domain-specific language (DSL) defined by the **@ode_def** macro for helping with this common problem. For example, we can define the Lotka-Volterra equation:

$$\frac{dx}{dt} = ax - bxy \tag{8}$$

$$\frac{dx}{dt} = ax - bxy$$

$$\frac{dy}{dt} = -cy + dxy$$
(8)

as follows:

```
function lotka_volterra!(du,u,p,t)
 du[1] = p[1]*u[1] - p[2]*u[1]*u[2]
 du[2] = -p[3]*u[2] + p[4]*u[1]*u[2]
end
```

lotka_volterra! (generic function with 1 method)

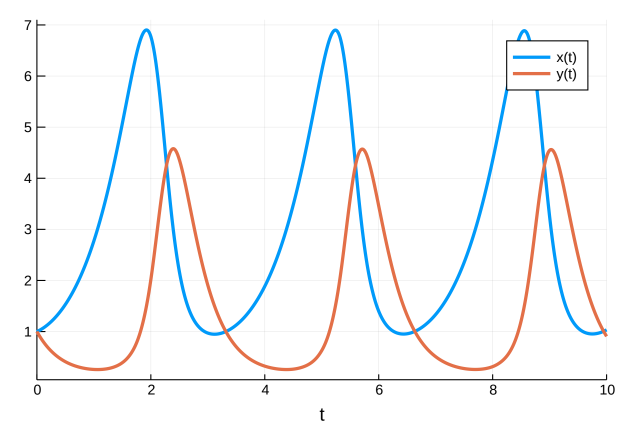
However, that can be hard to follow since there's a lot of "programming" getting in the way. Instead, you can use the **@ode_def** macro from ParameterizedFunctions.jl:

using ParameterizedFunctions

```
lv! = @ode_def LotkaVolterra begin
dx = a*x - b*x*y
dy = -c*y + d*x*y
end a b c d
```

We can then use the result just like an ODE function from before:

```
u0 = [1.0,1.0]
p = (1.5,1.0,3.0,1.0)
tspan = (0.0,10.0)
prob = ODEProblem(lv!,u0,tspan,p)
sol = solve(prob)
plot(sol)
```



Not only is the DSL convenient syntax, but it does some magic behind the scenes. For example, further parts of the tutorial will describe how solvers for stiff differential equations have to make use of the Jacobian in calculations. Here, the DSL uses symbolic differentiation to automatically derive that function:

```
lv!.Jex
```

```
quote
    internal_var___J[1, 1] = internal_var___p[1] - internal_var___p[2] * in
ternal_var___u[2]
    internal_var___J[1, 2] = -(internal_var___p[2]) * internal_var___u[1]
    internal_var___J[2, 1] = internal_var___p[4] * internal_var___u[2]
    internal_var___J[2, 2] = -(internal_var___p[3]) + internal_var___p[4] *
internal_var___u[1]
```

```
nothing end
```

The DSL can derive many other functions; this ability is used to speed up the solvers. An extension to Differential Equations.jl, Latexify.jl, allows you to extract these pieces as LaTeX expressions.

0.2 Internal Types

The last basic user-interface feature to explore is the choice of types. DifferentialEquations.jl respects your input types to determine the internal types that are used. Thus since in the previous cases, when we used Float64 values for the initial condition, this meant that the internal values would be solved using Float64. We made sure that time was specified via Float64 values, meaning that time steps would utilize 64-bit floats as well. But, by simply changing these types we can change what is used internally.

As a quick example, let's say we want to solve an ODE defined by a matrix. To do this, we can simply use a matrix as input.

```
A = [1. 0 0 -5
    4 -2 4 -3
    -4 0 0 1
    5 -2 2 3]
u0 = rand(4,2)
tspan = (0.0,1.0)
f(u,p,t) = A*u
prob = ODEProblem(f,u0,tspan)
sol = solve(prob)
```

There is no real difference from what we did before, but now in this case u0 is a 4x2 matrix. Because of that, the solution at each time point is matrix:

```
sol[3]
```

```
4×2 Array{Float64,2}:
0.325714 -0.0275025
0.548522 0.445243
0.50286 0.878366
1.38471 0.238715
```

In DifferentialEquations.jl, you can use any type that defines +, -, *, /, and has an appropriate norm. For example, if we want arbitrary precision floating point numbers, we can change the input to be a matrix of BigFloat:

```
big_u0 = big_u0)
```

and we can solve the ODEProblem with arbitrary precision numbers by using that initial condition:

```
prob = ODEProblem(f,big_u0,tspan)
sol = solve(prob)
sol[1,3]
```

-1.24462245861786783235698848817863208166206930525840630025592058823375357226417

To really make use of this, we would want to change abstol and reltol to be small! Notice that the type for "time" is different than the type for the dependent variables, and this can be used to optimize the algorithm via keeping multiple precisions. We can convert time to be arbitrary precision as well by defining our time span with BigFloat variables:

```
prob = ODEProblem(f,big_u0,big.(tspan))
sol = solve(prob)
```

Let's end by showing a more complicated use of types. For small arrays, it's usually faster to do operations on static arrays via the package StaticArrays.jl. The syntax is similar to that of normal arrays, but for these special arrays we utilize the @SMatrix macro to indicate we want to create a static array.

```
using StaticArrays
A = @SMatrix [ 1.0 0.0 0.0 -5.0
               4.0 -2.0 4.0 -3.0
               -4.0 0.0 0.0 1.0
               5.0 -2.0 2.0 3.0]
u0 = @SMatrix rand(4,2)
tspan = (0.0, 1.0)
f(u,p,t) = A*u
prob = ODEProblem(f,u0,tspan)
sol = solve(prob)
sol[3]
4×2 StaticArrays.SArray{Tuple{4,2},Float64,2,8}:
  0.678593 0.503025
  0.482739 0.690649
 -0.118648 0.632936
  0.993407 0.860104
```

0.3 Conclusion

These are the basic controls in DifferentialEquations.jl. All equations are defined via a problem type, and the solve command is used with an algorithm choice (or the default) to get a solution. Every solution acts the same, like an array sol[i] with sol.t[i], and also like a continuous function sol(t) with a nice plot command plot(sol). The Common

Solver Options can be used to control the solver for any equation type. Lastly, the types used in the numerical solving are determined by the input types, and this can be used to solve with arbitrary precision and add additional optimizations (this can be used to solve via GPUs for example!). While this was shown on ODEs, these techniques generalize to other types of equations as well.

0.4 Appendix

```
using DiffEqTutorials
DiffEqTutorials.tutorial_footer(WEAVE_ARGS[:folder],WEAVE_ARGS[:file])
These benchmarks are part of the DiffEqTutorials.jl repository, found at:
https://github.com/JuliaDiffEq/DiffEqTutorials.jl
To locally run this tutorial, do the following commands:
using DiffEqTutorials
DiffEqTutorials.weave_file("introduction", "ode_introduction.jmd")
Computer Information:
Julia Version 1.1.0
Commit 80516ca202 (2019-01-21 21:24 UTC)
Platform Info:
  OS: Windows (x86_64-w64-mingw32)
  CPU: Intel(R) Core(TM) i7-8700 CPU @ 3.20GHz
  WORD_SIZE: 64
  LIBM: libopenlibm
  LLVM: libLLVM-6.0.1 (ORCJIT, skylake)
Environment:
  JULIA_EDITOR = "C:\Users\accou\AppData\Local\atom\app-1.34.0\atom.exe" -a
  JULIA_NUM_THREADS = 6
Package Information:
    Status `C:\Users\accou\.julia\environments\v1.1\Project.toml`
  [7e558dbc] ArbNumerics v0.3.6
  [c52e3926] Atom v0.7.14
  [6e4b80f9] BenchmarkTools v0.4.2
  [336ed68f] CSV v0.4.3
  [3895d2a7] CUDAapi v0.5.4
  [be33ccc6] CUDAnative v1.0.1
  [3a865a2d] CuArrays v0.9.1
  [a93c6f00] DataFrames v0.17.1
  [55939f99] DecFP v0.4.8
  [abce61dc] Decimals v0.4.0
  [39dd38d3] Dierckx v0.4.1
  [bb2cbb15] DiffEqBenchmarks v0.0.0 [`C:\Users\accou\.julia\external\DiffE
qBenchmarks.jl`]
  [459566f4] DiffEqCallbacks v2.5.2
  [f3b72e0c] DiffEqDevTools v2.6.1
  [aae7a2af] DiffEqFlux v0.2.0
  [c894b116] DiffEqJump v6.1.0+ [`C:\Users\accou\.julia\dev\DiffEqJump`]
  [1130ab10] DiffEqParamEstim v1.6.0+ [`C:\Users\accou\.julia\dev\DiffEqPar
```

```
amEstim`]
  [055956cb] DiffEqPhysics v3.1.0
  [a077e3f3] DiffEqProblemLibrary v4.1.0
  [225cb15b] DiffEqTutorials v0.0.0 [`C:\Users\accou\.julia\external\DiffEq
Tutorials.jl`]
  [0c46a032] DifferentialEquations v6.3.0
  [497a8b3b] DoubleFloats v0.7.5
  [587475ba] Flux v0.7.3
  [f6369f11] ForwardDiff v0.10.3+ [`C:\Users\accou\.julia\dev\ForwardDiff`]
  [28b8d3ca] GR v0.38.1
  [7073ff75] IJulia v1.17.0
  [c601a237] Interact v0.9.1
  [b6b21f68] Ipopt v0.5.4
  [4076af6c] JuMP v0.19.0
  [e5e0dc1b] Juno v0.5.4
  [7f56f5a3] LSODA v0.4.0
  [eff96d63] Measurements v2.0.0
  [76087f3c] NLopt v0.5.1
  [c030b06c] ODE v2.4.0
  [54ca160b] ODEInterface v0.4.5+ [`C:\Users\accou\.julia\dev\ODEInterface`
  [09606e27] ODEInterfaceDiffEq v3.0.0
  [429524aa] Optim v0.17.2
  [1dea7af3] OrdinaryDiffEq v5.2.1+ [`C:\Users\accou\.julia\dev\OrdinaryDif
fEq`]
  [65888b18] ParameterizedFunctions v4.1.1
  [91a5bcdd] Plots v0.23.0
  [71ad9d73] PuMaS v0.0.0 [`C:\Users\accou\.julia\dev\PuMaS`]
  [d330b81b] PyPlot v2.7.0
  [731186ca] RecursiveArrayTools v0.20.0
  [90137ffa] StaticArrays v0.10.2
  [789caeaf] StochasticDiffEq v6.1.1+ [`C:\Users\accou\.julia\dev\Stochasti
cDiffEq ]
  [c3572dad] Sundials v3.0.0
  [1986cc42] Unitful v0.14.0
  [2a06ce6d] UnitfulPlots v0.0.0 #master (https://github.com/ajkeller34/Uni
tfulPlots.jl)
  [44d3d7a6] Weave v0.7.1 [`C:\Users\accou\.julia\dev\Weave`]
```