Kepler Problem

Yingbo Ma, Chris Rackauckas

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The Hamiltonian \mathcal{H} and the angular momentum L for the Kepler problem are

$$\mathcal{H} = \frac{1}{2}(\dot{q}_1^2 + \dot{q}_2^2) - \frac{1}{\sqrt{q_1^2 + q_2^2}}, \quad L = q_1\dot{q}_2 - \dot{q}_1q_2$$

Also, we know that

$$\frac{\mathrm{d}\boldsymbol{p}}{\mathrm{d}t} = -\frac{\partial \mathcal{H}}{\partial \boldsymbol{q}} \quad , \quad \frac{\mathrm{d}\boldsymbol{q}}{\mathrm{d}t} = +\frac{\partial \mathcal{H}}{\partial \boldsymbol{p}}$$

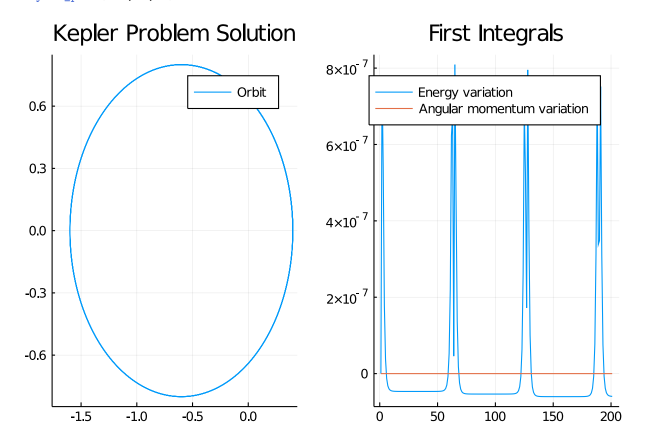
```
using OrdinaryDiffEq, LinearAlgebra, ForwardDiff, Plots; gr()
H(q,p) = norm(p)^2/2 - inv(norm(q))
L(q,p) = q[1]*p[2] - p[1]*q[2]
pdot(dp,p,q,params,t) = ForwardDiff.gradient!(dp, q->-H(q, p), q)
qdot(dq,p,q,params,t) = ForwardDiff.gradient!(dq, p-> H(q, p), p)
initial_position = [.4, 0]
initial_velocity = [0., 2.]
initial_cond = (initial_position, initial_velocity)
initial_first_integrals = (H(initial_cond...), L(initial_cond...))
tspan = (0,20.)
prob = DynamicalODEProblem(pdot, qdot, initial_velocity, initial_position, tspan)
sol = solve(prob, KahanLi6(), dt=1//10);
retcode: Success
Interpolation: 3rd order Hermite
t: 201-element Array{Float64,1}:
  0.0
  0.1
  0.2
  0.3000000000000004
  0.4
  0.5
  0.6
  0.7
  0.799999999999999
  0.899999999999999
 19.200000000000003
 19.300000000000004
 19.400000000000006
 19.500000000000007
```

```
19.60000000000001
  19.70000000000001
  19 8000000000001
  19.900000000000013
 20.0
u: 201-element Array{RecursiveArrayTools.ArrayPartition{Float64,Tuple{Array
{Float64,1},Array{Float64,1}},1}:
  [0.0, 2.0][0.4, 0.0]
  [-0.5830949354540153, 1.8556656829703986] [0.36982713146498514, 0.195035965
14776078]
  [-0.9788105843777312, 1.5274462532150213] [0.28987830863610903, 0.364959747
35762693]
  [-1.17547762665905, 1.1751394486895783] [0.18078065407309682, 0.49984577206
18293]
  [-1.2440239387295458, 0.8720450804540057] [0.05902925334751511, 0.601695680]
2132387]
  [-1.2441259417439434, 0.6289994697149073] [-0.06577256855272472, 0.67627471
022914821
  [-1.210142434136089, 0.4368770315976506] [-0.188677607179601, 0.72919425685]
  [-1.159918613868923, 0.28408169071815415] [-0.30726896099260204, 0.76495839
909355681
  [-1.1025329550493486, 0.16100716005909121] [-0.42042727561865095, 0.7869985
1798978897
  [-1.0426125487031446, 0.06047044972523817][-0.5276934467175253, 0.79790892]
70264804]
  [-1.2216434770974676, 1.0146166139270498] [0.12021680827053512, 0.555011377
  [-1.2499499900381417, 0.7423750265723883] [-0.003918416420213356, 0.6423528
468283568]
  [-1.2298310873691611, 0.5265058660314975] [-0.12818281922639643, 0.70537248
  [-1.1861148292768293, 0.3555788492114466] [-0.24911096207713992, 0.74915056
05432051]
  [-1.1314960903670108, 0.2188164842264573] [-0.36504892367796954, 0.77762418
23022721]
  [-1.0724336821492226, 0.10787192092148691][-0.47526538003987784, 0.7937719]
633967374]
  [-1.0122234000273465, 0.016617787590286977] [-0.579499110310746, 0.79985306
90972269]
  [-0.9525349461454056, -0.05939856743324051] [-0.6777283550436937, 0.7976021]
3488854071
   \hbox{ $[-0.894185739649566, $-0.12343221924182135] $[-0.7700512224747644, $0.78837185] $[-0.894185739649566, $-0.12343221924182135] $[-0.894185739649566, $-0.894185739649566] $[-0.894185739649566, $-0.894185739649566] $[-0.894185739649566, $-0.894185739649566] $[-0.894185739649566] $[-0.894185739649566] $[-0.894185739649566] $[-0.894185739649566] $[-0.894185739649566] $[-0.894185739649566] $[-0.894185739649566] $[-0.894185739649566] $[-0.89418573964956] $[-0.89418573964] $[-0.89418573964] $[-0.89418573964] $[-0.8941857396] $[-0.8941857396] $[-0.8941857396] $[-0.8941857396] $[-0.8941857396] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.894185739] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.8941857] $[-0.894185
320841]
Let's plot the orbit and check the energy and angular momentum variation. We know that
plot_orbit(sol) = plot(sol,vars=(3,4), lab="Orbit", title="Kepler Problem Solution")
```

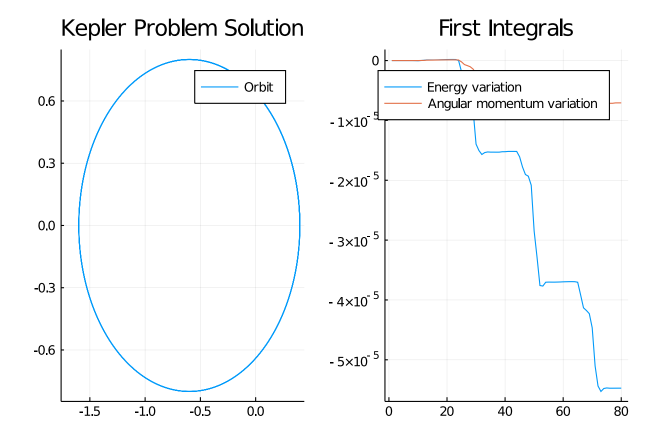
energy and angular momentum should be constant, and they are also called first integrals.

```
function plot_first_integrals(sol, H, L)
   plot(initial_first_integrals[1].-map(u->H(u[2,:], u[1,:]), sol.u), lab="Energy
variation", title="First Integrals")
   plot!(initial_first_integrals[2].-map(u->L(u[2,:], u[1,:]), sol.u), lab="Angular
momentum variation")
analysis_plot(sol, H, L) = plot(plot_orbit(sol), plot_first_integrals(sol, H, L))
```

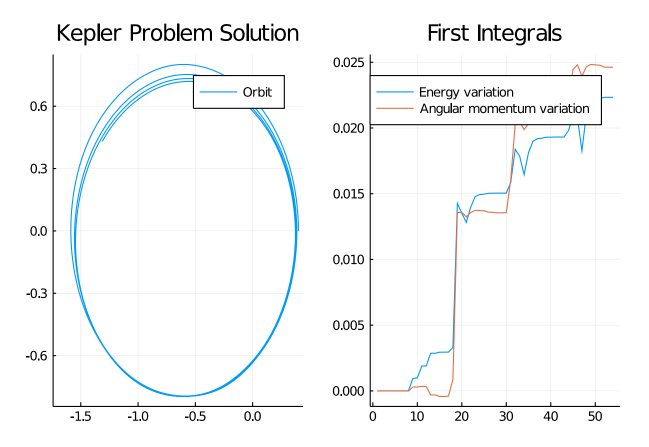
analysis_plot(sol, H, L)



Let's try to use a Runge-Kutta-Nyström solver to solve this problem and check the first integrals' variation.



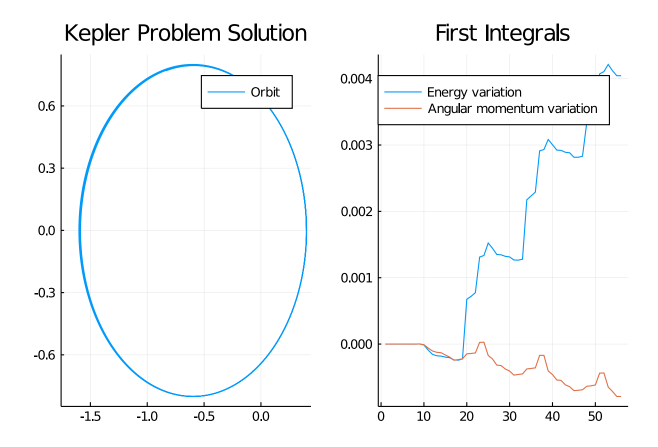
Let's then try to solve the same problem by the ${\tt ERKN4}$ solver, which is specialized for sinusoid-like periodic function



We can see that ERKN4 does a bad job for this problem, because this problem is not sinusoid-like.

One advantage of using DynamicalODEProblem is that it can implicitly convert the second order ODE problem to a *normal* system of first order ODEs, which is solvable for other ODE solvers. Let's use the Tsit5 solver for the next example.

```
sol4 = solve(prob, Tsit5())
@show sol4.u |> length
sol4.u |> length = 56
analysis_plot(sol4, H, L)
```



Note There is drifting for all the solutions, and high order methods are drifting less because they are more accurate.

0.0.1 Conclusion

Symplectic integrator does not conserve the energy completely at all time, but the energy can come back. In order to make sure that the energy fluctuation comes back eventually, symplectic integrator has to have a fixed time step. Despite the energy variation, symplectic integrator conserves the angular momentum perfectly.

Both Runge-Kutta-Nyström and Runge-Kutta integrator do not conserve energy nor the angular momentum, and the first integrals do not tend to come back. An advantage Runge-Kutta-Nyström integrator over symplectic integrator is that RKN integrator can have adaptivity. An advantage Runge-Kutta-Nyström integrator over Runge-Kutta integrator is that RKN integrator has less function evaluation per step. The ERKN4 solver works best for sinusoid-like solutions.

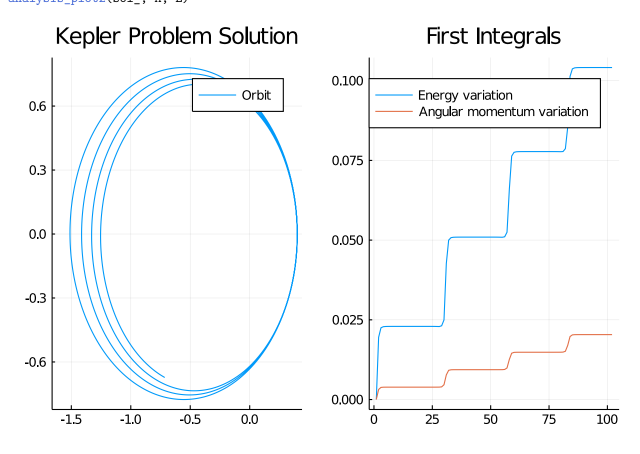
0.1 Manifold Projection

In this example, we know that energy and angular momentum should be conserved. We can achieve this through mainfold projection. As the name implies, it is a procedure to project the ODE solution to a manifold. Let's start with a base case, where mainfold projection isn't being used.

using DiffEqCallbacks

```
- Run `import Pkg; Pkg.add("DiffEqCallbacks")` to install the DiffEqCallbac
ks package.
plot_orbit2(sol) = plot(sol,vars=(1,2), lab="Orbit", title="Kepler Problem Solution")
function plot_first_integrals2(sol, H, L)
    plot(initial_first_integrals[1].-map(u->H(u[1:2],u[3:4]), sol.u), lab="Energy
variation", title="First Integrals")
    plot!(initial_first_integrals[2].-map(u->L(u[1:2],u[3:4]), sol.u), lab="Angular
momentum variation")
end
analysis_plot2(sol, H, L) = plot(plot_orbit2(sol), plot_first_integrals2(sol, H, L))
function hamiltonian(du,u,params,t)
    q, p = u[1:2], u[3:4]
    qdot(@view(du[1:2]), p, q, params, t)
    pdot(@view(du[3:4]), p, q, params, t)
end
prob2 = ODEProblem(hamiltonian, [initial_position; initial_velocity], tspan)
sol_ = solve(prob2, RK4(), dt=1//5, adaptive=false)
analysis_plot2(sol_, H, L)
```

Error: ArgumentError: Package DiffEqCallbacks not found in current path:



There is a significant fluctuation in the first integrals, when there is no mainfold projection.

```
function first_integrals_manifold(residual,u)
    residual[1:2] .= initial_first_integrals[1] - H(u[1:2], u[3:4])
    residual[3:4] .= initial_first_integrals[2] - L(u[1:2], u[3:4])
end

cb = ManifoldProjection(first_integrals_manifold)
```

```
Error: UndefVarError: ManifoldProjection not defined
sol5 = solve(prob2, RK4(), dt=1//5, adaptive=false, callback=cb)
Error: UndefVarError: cb not defined
analysis_plot2(sol5, H, L)
Error: UndefVarError: sol5 not defined
We can see that thanks to the manifold projection, the first integrals' variation is very small,
although we are using RK4 which is not symplectic. But wait, what if we only project to the
energy conservation manifold?
function energy_manifold(residual,u)
    residual[1:2] .= initial_first_integrals[1] - H(u[1:2], u[3:4])
    residual[3:4] = 0
energy_cb = ManifoldProjection(energy_manifold)
Error: UndefVarError: ManifoldProjection not defined
sol6 = solve(prob2, RK4(), dt=1//5, adaptive=false, callback=energy_cb)
Error: UndefVarError: energy_cb not defined
analysis_plot2(sol6, H, L)
Error: UndefVarError: sol6 not defined
There is almost no energy variation but angular momentum varies quite bit. How about
only project to the angular momentum conservation manifold?
function angular_manifold(residual,u)
    residual[1:2] .= initial_first_integrals[2] - L(u[1:2], u[3:4])
    residual[3:4] .= 0
end
angular_cb = ManifoldProjection(angular_manifold)
Error: UndefVarError: ManifoldProjection not defined
sol7 = solve(prob2, RK4(), dt=1//5, adaptive=false, callback=angular_cb)
Error: UndefVarError: angular_cb not defined
analysis plot2(sol7, H, L)
Error: UndefVarError: sol7 not defined
```

0.2 Appendix

Again, we see what we expect.

This tutorial is part of the DiffEqTutorials.jl repository, found at: https://github.com/JuliaDiffEq/DiffEqTutorials.jl repository, found at: https://github.com/JuliaDiffEq/DiffEqTutorials.jl repository, found at: https://github.com/JuliaDiffEq/DiffEq

```
using DiffEqTutorials
DiffEqTutorials.weave_file("models","05-kepler_problem.jmd")
Computer Information:
Julia Version 1.4.2
Commit 44fa15b150* (2020-05-23 18:35 UTC)
Platform Info:
 OS: Linux (x86_64-pc-linux-gnu)
 CPU: Intel(R) Core(TM) i7-9700K CPU @ 3.60GHz
 WORD_SIZE: 64
 LIBM: libopenlibm
 LLVM: libLLVM-8.0.1 (ORCJIT, skylake)
Environment:
 JULIA_DEPOT_PATH = /builds/JuliaGPU/DiffEqTutorials.jl/.julia
 JULIA_CUDA_MEMORY_LIMIT = 536870912
 JULIA PROJECT = @.
 JULIA NUM THREADS = 4
Package Information:
Status `/builds/JuliaGPU/DiffEqTutorials.jl/tutorials/models/Project.toml`
[eb300fae-53e8-50a0-950c-e21f52c2b7e0] DiffEqBiological 4.3.0
[f3b72e0c-5b89-59e1-b016-84e28bfd966d] DiffEqDevTools 2.22.0
[055956cb-9e8b-5191-98cc-73ae4a59e68a] DiffEqPhysics 3.2.0
[Oc46a032-eb83-5123-abaf-570d42b7fbaa] DifferentialEquations 6.14.0
[31c24e10-a181-5473-b8eb-7969acd0382f] Distributions 0.23.4
[587475ba-b771-5e3f-ad9e-33799f191a9c] Flux 0.10.4
[f6369f11-7733-5829-9624-2563aa707210] ForwardDiff 0.10.10
[23fbe1c1-3f47-55db-b15f-69d7ec21a316] Latexify 0.13.5
[961ee093-0014-501f-94e3-6117800e7a78] ModelingToolkit 3.10.2
```

[2774e3e8-f4cf-5e23-947b-6d7e65073b56] NLsolve 4.4.0

[429524aa-4258-5aef-a3af-852621145aeb] Optim 0.21.0

[91a5bcdd-55d7-5caf-9e0b-520d859cae80] Plots 1.4.3

[37e2e46d-f89d-539d-b4ee-838fcccc9c8e] LinearAlgebra [2f01184e-e22b-5df5-ae63-d93ebab69eaf] SparseArrays

[8faf48c0-8b73-11e9-0e63-2155955bfa4d] NeuralNetDiffEq 1.6.0

[1dea7af3-3e70-54e6-95c3-0bf5283fa5ed] OrdinaryDiffEq 5.41.0

[731186ca-8d62-57ce-b412-fbd966d074cd] RecursiveArrayTools 2.5.0 [789caeaf-c7a9-5a7d-9973-96adeb23e2a0] StochasticDiffEq 6.23.1