Julia Warnings

Just like Python, Julia lets you do the following:

```
In [2]:
         1 let
                 x = [1,2,3]
          3
                 @show x
                 y = x # NEVER DO THIS, EDITING ONE WILL NOW EDIT BOTH
          4
                 y[3] = 100 # this will now modify both y and x
          6
                 x[1] = 300 \#  this will now modify both y and x
          7
          8
          9
         10
                 @show x
         11 end
        x = [1, 2, 3]
        y = [300, 2, 100]
        x = [300, 2, 100]
Out[2]: 3-element Vector{Int64}:
         300
           2
         100
In [3]:
          1
            # to avoid this, here are two alternatives
          2
            let
          3
                x = [1,2,3]
          4
                 @show x
          5
          6
                 y1 = 1*x
                                   # this is fine
          7
                 y2 = deepcopy(x) # this is also fine
          8
          9
                 x[2] = 200 \# only edits x
                 y1[1] = 400 # only edits y1
y2[3] = 100 # only edits y2
         10
         11
         12
         13
                 @show x
                 @show y1
         14
         15
                 @show y2
         16 end
        x = [1, 2, 3]
        x = [1, 200, 3]
        y1 = [400, 2, 3]
        y2 = [1, 2, 100]
Out[3]: 3-element Vector{Int64}:
           1
           2
```

Optional function arguments

100

We can have optional keyword arguments for functions in Julia, like the following:

```
In [4]:
          1 ## optional arguments in functions
           3 # we can have functions with optional arguments after a ; that have default values
           4 let
           5
                  function f1(a, b; c=4, d=5)
           6
                       @show a,b,c,d
           7
           8
           9
                                           # this means c and d will take on default value
          10
                  f1(1,2;c = 100,d = 2) # specify c and d
                  f1(1,2;d = -30)
                                          # or we can only specify one of them
          11
          12 end
         (a, b, c, d) = (1, 2, 4, 5)
(a, b, c, d) = (1, 2, 100, 2)
(a, b, c, d) = (1, 2, 4, -30)
Out[4]: (1, 2, 4, -30)
```

Q1: Integration (20 pts)

In this question we are going to integrate the equations of motion for a double pendulum using multiple explicit and implicit integrators. We will write a generic simulation function for each of the two categories (explicit and implicit), and compare 6 different integrators.

The continuous time dynamics of the cartpole are written as a function:

```
\dot{x} = f(x)
```

In the code you will see xdot = dynamics(params, x).

Part A (10 pts): Explicit Integration

Here we are going to implement the following explicit integrators:

- Forward Euler (explicit)
- · Midpoint (explicit)
- RK4 (explicit)

```
In [5]:
           1 # these two functions are given, no TODO's here
              function double_pendulum_dynamics(params::NamedTuple, x::Vector)
                    \# continuous time dynamics for a double pendulum given state x,
            3
                    # also known as the "equations of motion".
            4
            5
                    # returns the time derivative of the state, \dot{x} (dx/dt)
            6
            7
                    # the state is the following:
            8
                    \theta1,\theta1,\theta2,\theta2 = x
            9
          10
                    # system parameters
                    m1, m2, L1, L2, g = params.m1, params.m2, params.L1, params.L2, params.g
          11
          12
          13
                    # dynamics
                   c = cos(\theta 1 - \theta 2)
          14
          15
                    s = \sin(\theta 1 - \theta 2)
          16
          17
                    \dot{x} = [
                         Θĺ;
          18
                          (\ m2*g*\sin(\theta 2)*c - m2*s*(L1*c*\theta 1^2 + L2*\theta 2^2) - (m1+m2)*g*\sin(\theta 1) \ ) \ / \ (\ L1 *(m1+m2*s^2) \ ); 
          19
          20
                         θŻ:
                         ((m1+m2)*(L1*\theta\dot{1}^2*s - g*sin(\theta 2) + g*sin(\theta 1)*c) + m2*L2*\theta\dot{2}^2*s*c) / (L2 * (m1 + m2*s^2));
          21
          22
          23
          24
                    return x
          25
               end
               function double_pendulum_energy(params::NamedTuple, x::Vector)::Real
          26
          27
                    # calculate the total energy (kinetic + potential) of a double pendulum given a state x
          28
          29
          30
                    # the state is the following:
          31
                    \theta1,\theta1,\theta2,\theta2 = x
          32
          33
                    # system parameters
          34
                    m1, m2, L1, L2, g = params.m1, params.m2, params.L1, params.L2, params.g
          35
          36
                    # cartesian positions/velocities of the masses
          37
                    r1 = [L1*sin(\theta 1), 0, -params.L1*cos(\theta 1) + 2]
                   r2 = r1 + [params.L2*sin(\theta2), 0, -params.L2*cos(\theta2)]

v1 = [L1*\thetai*cos(\theta1), 0, L1*\thetai*sin(\theta1)]
          38
          39
                    v2 = v1 + [L2*\theta\dot{2}*cos(\theta 2), 0, L2*\theta\dot{2}*sin(\theta 2)]
          40
          41
          42
                    # energy calculation
          43
                    kinetic = 0.5*(m1*v1'*v1 + m2*v2'*v2)
                    potential = m1*g*r1[3] + m2*g*r2[3]
          44
          45
                    return kinetic + potential
          46 end
```

Out[5]: double_pendulum_energy (generic function with 1 method)

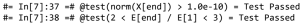
Now we are going to simulate this double pendulum by integrating the equations of motion with the simplest explicit integrator, the Forward Euler method:

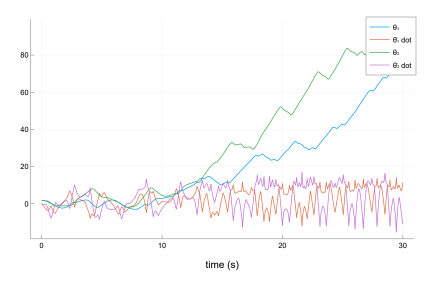
```
x_{k+1} = x_k + \Delta t \cdot f(x_k) Forward Euler (explicit)
```

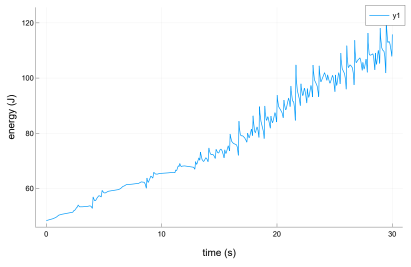
```
1 """
In [6]:
                 x_{k+1} = forward_euler(params, dynamics, x_k, dt)
          4 Given \dot{x} = dynamics(params, x), take in the current state \dot{x} and integrate it forward \dot{d}
          5 using Forward Euler method.
          6
          7
             function forward_euler(params::NamedTuple, dynamics::Function, x::Vector, dt::Real)::Vector
          8
                 \dot{x} = dynamics(params, x)
                 # TODO: implement forward euler
                 # error("forward euler not implemented")
         10
         11
                 return x + \dot{x} .* dt
         12 end
```

Out[6]: forward_euler

```
In [7]:
           include(joinpath(@__DIR__, "animation.jl"))
           3
              let
           4
           5
                   # parameters for the simulation
           6
                   params = (
           7
                        m1 = 1.0.
           8
                        m2 = 1.0,
           9
                        L1 = 1.0,
          10
                        L2 = 1.0
                        g = 9.8
          11
          12
                   )
          13
          14
                   # initial condition
          15
                   x0 = [pi/1.6; 0; pi/1.8; 0]
          16
          17
                   # time step size (s)
          18
                   dt = 0.01
          19
                   tf = 30.0
                   t_vec = 0:dt:tf
          20
          21
                   N = length(t_vec)
          22
          23
                   # store the trajectory in a vector of vectors
          24
                   X = [zeros(4) for i = 1:N]
          25
                   X[1] = 1*x0
          26
                   # TODO: simulate the double pendulum with `forward_euler`
          27
          28
                   \# X[k] = `x_k`, so X[k+1] = forward\_euler(params, double\_pendulum\_dynamics, <math>X[k], dt)
          29
          30
                   for k in 1:N-1
          31
                        X[k+1] = forward_euler(params, double_pendulum_dynamics, X[k], dt)
          32
          33
          34
                   # calculate energy
          35
                   E = [double_pendulum_energy(params,x) for x in X]
          36
          37
                   @show @test norm(X[end]) > 1e-10 # make sure all X's were updated
          38
                   @show @test 2 < (E[end]/E[1]) < 3 # energy should be increasing
          39
                   # plot state history, energy history, and animate it
          40
                   \label{eq:display} \begin{split} & \text{display(plot(t_vec, hcat(X...)',xlabel = "time (s)", label = ["$\theta_1$" "$\theta_1$" dot" "$\theta_2$" "$\theta_2$" dot"]))} \\ & \text{display(plot(t_vec, E, xlabel = "time (s)", ylabel = "energy (J)"))} \end{split}
          41
          42
          43
                   meshcat_animate(params,X,dt,N)
          44
          45
          46
              end
          47
          48
```

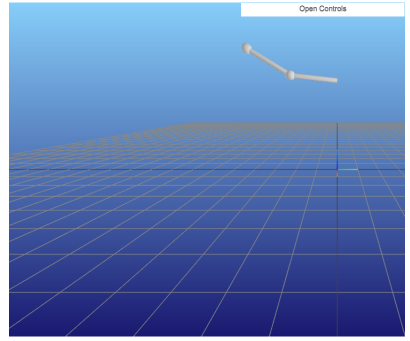






Info: MeshCat server started. You can open the visualizer by visiting the following URL in your browser:
http://127.0.0.1:8713





Now let's implement the next two integrators:

Midpoint:

$$x_m = x_k + \frac{\Delta t}{2} \cdot f(x_k)$$

$$x_{k+1} = x_k + \Delta t \cdot f(x_m)$$

RK4:

$$\begin{aligned} k_1 &= \Delta t \cdot f(x_k) \\ k_2 &= \Delta t \cdot f(x_k + k_1/2) \\ k_3 &= \Delta t \cdot f(x_k + k_2/2) \\ k_4 &= \Delta t \cdot f(x_k + k_3) \\ x_{k+1} &= x_k + (1/6) \cdot (k_1 + 2k_2 + 2k_3 + k_4) \end{aligned}$$

1

```
In [8]:
         1 function midpoint(params::NamedTuple, dynamics::Function, x::Vector, dt::Real)::Vector
                # TODO: implement explicit midpoint
          3 #
                  error("midpoint not implemented")
          4
                 \dot{x}_k = dynamics(params, x)
          5
                 x_m = x + dt * \dot{x}_k / 2
                 return x + dynamics(params, x<sub>m</sub>) .* dt
          7
            end
          8
             function rk4(params::NamedTuple, dynamics::Function, x::Vector, dt::Real)::Vector
          9
                # TODO: implement RK4
                  error("rk4 not implemented")
         10 #
                 k_1 = dynamics(params, x) .* dt
         11
                 k_2 = dynamics(params, x + k_1 / 2) .* dt
         12
         13
                 k_3 = dynamics(params, x + k_2 / 2) .* dt
                 k_4 = dynamics(params, x + k_3) .* dt
         15
                 return x + (k_1 + 2*k_2 + 2*k_3 + k_4) / 6
```

Out[8]: rk4 (generic function with 1 method)

```
In [9]:
        1 function simulate_explicit(params::NamedTuple,dynamics::Function,integrator::Function,x0::Vector,dt::Real,tf::Real)
                # TOOD: update this function to simulate dynamics forward
          3
                # with the given explicit integrator
          4
          5
          6
                # take in
          7
                t_vec = 0:dt:tf
          8
                N = length(t_vec)
                X = [zeros(length(x0)) for i = 1:N]
          9
         10
                X[1] = x0
         11
                # TODO: simulate X forward
         12
         13
                for k in 1:N-1
                    X[k+1] = integrator(params, dynamics, X[k], dt)
         14
         15
         16
                # return state history X and energy E
         17
                E = [double_pendulum_energy(params,x) for x in X]
         18
         19
                return X, E
```

Out[9]: simulate_explicit (generic function with 1 method)

Out[10]: (m1 = 1.0, m2 = 1.0, L1 = 1.0, L2 = 1.0, g = 9.8)

Part B (10 pts): Implicit Integrators

Explicit integrators work by calling a function with x_k and Δt as arguments, and returning x_{k+1} like this:

$$x_{k+1} = f_{explicit}(x_k, \Delta t)$$

Implicit integrators on the other hand have the following relationship between the state at x_k and x_{k+1} :

$$f_{implicit}(x_k, x_{k+1}, \Delta t) = 0$$

This means that if we want to get x_{k+1} from x_k , we have to solve for a x_{k+1} that satisfies the above equation. This is a rootfinding problem in x_{k+1} (our unknown), so we juse have to use Newton's method.

Here are the three implicit integrators we are looking at, the first being Backward Euler (1st order):

$$f(x_k, x_{k+1}, \Delta t) = x_k + \Delta t \cdot \dot{x}_{k+1} - x_{k+1} = 0$$
 Backward Euler

Implicit Midpoint (2nd order)

$$x_{k+1/2} = \frac{1}{2}(x_k + x_{k+1})$$

$$f(x_k, x_{k+1}, \Delta t) = x_k + \Delta t \cdot \dot{x}_{k+1/2} - x_{k+1} = 0$$
 Implicit Midpoint

Hermite Simpson (3rd order)

$$\begin{split} x_{k+1/2} &= \frac{1}{2}(x_k + x_{k+1}) + \frac{\Delta t}{8}(\dot{x}_k - \dot{x}_{k+1}) \\ f(x_k, x_{k+1}, \Delta t) &= x_k + \frac{\Delta t}{6} \cdot (\dot{x}_k + 4\dot{x}_{k+1/2} + \dot{x}_{k+1}) - x_{k+1} = 0 \end{split}$$
 Hermite-Simpson

When you implement these integrators, you will update the functions such that they take in a dynamics function, x_k and x_{k+1} , and return the residuals described above. We are NOT solving these yet, we are simply returning the residuals for each implicit integrator that we want to be 0.

```
In [11]:
          1 # since these are explicit integrators, these function will return the residuals described above
           2 # NOTE: we are NOT solving anything here, simply return the residuals
           3 function backward_euler(params::NamedTuple, dynamics::Function, x1::Vector, x2::Vector, dt::Real)::Vector
                   error("backward euler not implemented")
           5
                  return x1 + dt .* dynamics(params, x2) - x2
           6 end
              function implicit_midpoint(params::NamedTuple, dynamics::Function, x1::Vector, x2::Vector, dt::Real)::Vector
           8
                    error("implicit midpoint not implemented")
                  x_{mid} = (x1 + x2) / 2
           9
          10
                  return x1 + dt .* dynamics(params, x_mid) - x2
          11 end
          12 function hermite_simpson(params::NamedTuple, dynamics::Function, x1::Vector, x2::Vector, dt::Real)::Vector
          13 #
                    error("hermite simpson not implemented")
                  xk_dot = dynamics(params, x1)
          14
          15
                  xk1_dot = dynamics(params, x2)
          16
          17
                  xk_half = 0.5 * (x1 + x2) + dt .* (xk_dot - xk1_dot) / 8
                  xk_half_dot = dynamics(params, xk_half)
return x1 + dt .* (xk_dot + 4 * xk_half_dot + xk1_dot) / 6 - x2
          18
          19
          20 end
```

Out[11]: hermite simpson (generic function with 1 method)

```
In [12]:
          1 # TODO
           2 \mid # this function takes in a dynamics function, implicit integrator function, and x1
           3 # and uses Newton's method to solve for an x2 that satsifies the implicit integration equations
           4 # that we wrote about in the functions above
           5 function implicit_integrator_solve(params::NamedTuple, dynamics::Function, implicit_integrator::Function, x1::Vector, dt::Re
           7
                  # initialize auess
           8
                  x2 = 1*x1
           9
          10
                  # TODO: use Newton's method to solve for x2 such that residual for the integrator is 0
                  # DO NOT USE A WHILE LOOP
          11
          12
          13
                  for i = 1:max_iters
          14
                      # TODO: return x2 when the norm of the residual is below tol
          15
          16
                      xn = implicit_integrator(params, dynamics, x1, x2, dt)
          17
                      \Delta x = - FD.jacobian(x2 -> implicit_integrator(params, dynamics, x1, x2, dt),x2)\xn
          18
                      x2 = x2 + \Delta x
          19
          20
                      if norm(xn) < tol</pre>
          21
                          return x2
          22
                      end
          23
          24
          25
                  error("implicit integrator solve failed")
          26 end
```

Out[12]: implicit_integrator_solve (generic function with 1 method)

```
In [13]:
           1 @testset "implicit integrator check" begin
           3
                  dt = 1e-1
                  x1 = [.1, .2, .3, .4]
           4
           5
                  for integrator in [backward_euler, implicit_midpoint, hermite_simpson]
           6
           7
                      println("----testing $integrator -----")
           8
                      x2 = implicit_integrator_solve(params, double_pendulum_dynamics, integrator, x1, dt)
           9
                      @test norm(integrator(params, double_pendulum_dynamics, x1, x2, dt)) < 1e-10</pre>
          10
                  end
          11
          12 end
         ----testing backward_euler -----
```

```
----testing implicit_midpoint -----
----testing hermite_simpson -----

Test Summary: | Pass Total
implicit integrator check | 3 3
```

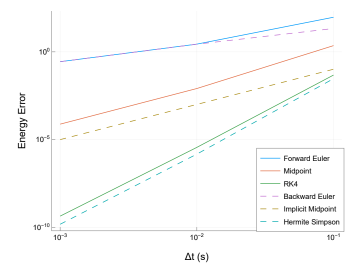
Out[13]: Test.DefaultTestSet("implicit integrator check", Any[], 3, false, false)

```
In [14]:
              function simulate_implicit(params::NamedTuple,dynamics::Function,implicit_integrator::Function,x0::Vector,dt::Real,tf::Real);
           1
                  t vec = 0:dt:tf
           2
           3
                  N = length(t_vec)
           4
                  X = [zeros(length(x0)) for i = 1:N]
           5
                  X[1] = x0
           6
           7
                  # TODO: do a forward simulation with the selected implicit integrator
           8
                  # hint: use your `implicit_integrator_solve` function
           9
          10
                  for k in 1:N-1
                      X[k+1] = implicit_integrator_solve(params, dynamics, implicit_integrator, X[k], dt, tol=tol)
          11
          12
          13
                  E = [double_pendulum_energy(params,x) for x in X]
          14
          15
                  @assert length(X)==N
                  @assert length(E)==N
          16
          17
                  return X, E
          18
             end
              ∢ I
```

Out[14]: simulate_implicit (generic function with 1 method)

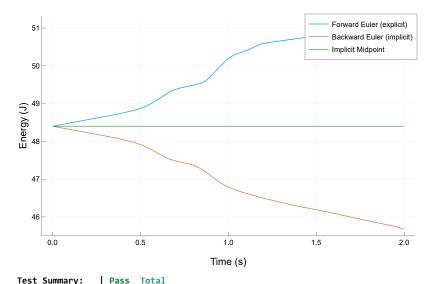
```
In [15]:
           1 function max_err_E(E)
           2
                  E0 = E[1]
           3
                  err = abs.(E - E0)
           4
                  return maximum(err)
           5
              end
           6
              function get_explicit_energy_error(integrator::Function, dts::Vector)
           7
                  [max_err_E(simulate_explicit(params,double_pendulum_dynamics,integrator,x0,dt,tf)[2]) for dt in dts]
           8
              function get_implicit_energy_error(integrator::Function, dts::Vector)
          10
                  [\max\_err\_E(simulate\_implicit(params,double\_pendulum\_dynamics,integrator,x0,dt,tf)[2]) \ \ for \ \ dt \ \ in \ \ dts]
          11
              end
          12
          13
              const tf = 2.0
          14
          15
              let
          16
                  # here we compare everything
          17
                  dts = [1e-3, 1e-2, 1e-1]
          18
                  explicit_integrators = [forward_euler, midpoint, rk4]
                  implicit_integrators = [backward_euler, implicit_midpoint, hermite_simpson]
          19
          20
          21
                  explicit_data = [get_explicit_energy_error(integrator, dts) for integrator in explicit_integrators]
          22
                  implicit_data = [get_implicit_energy_error(integrator, dts) for integrator in implicit_integrators]
          23
                  plot(dts, hcat(explicit_data...),label = ["Forward Euler" "Midpoint" "RK4"],xaxis=:log10,yaxis=:log10, xlabel = "Δt (s)"
          24
                  plot!(dts, hcat(implicit_data...),ls = :dash, label = ["Backward Euler" "Implicit Midpoint" "Hermite Simpson"])
          25
          26
                  plot!(legend=:bottomright)
          27
              end
          28
```

Out[15]:



What we can see above is the maximum energy error for each of the integration methods. In general, the implicit methods of the same order are slightly better than the explicit ones.

```
In [16]:
              @testset "energy behavior" begin
           1
           3
                  # simulate with all integrators
           4
                  dt = 0.01
           5
                  t_vec = 0:dt:tf
           6
                  E1 = simulate_explicit(params,double_pendulum_dynamics,forward_euler,x0,dt,tf)[2]
                  E2 = simulate_implicit(params,double_pendulum_dynamics,backward_euler,x0,dt,tf)[2]
           7
           8
                  E3 = simulate_implicit(params,double_pendulum_dynamics,implicit_midpoint,x0,dt,tf)[2]
           9
                  E4 = simulate_implicit(params,double_pendulum_dynamics,hermite_simpson,x0,dt,tf)[2]
          10
                  E5 = simulate_explicit(params,double_pendulum_dynamics,midpoint,x0,dt,tf)[2]
                  E6 = simulate_explicit(params,double_pendulum_dynamics,rk4,x0,dt,tf)[2]
          11
          12
          13
                  # plot forward/backward euler and implicit midpoint
          14
                  plot(t_vec,E1, label = "Forward Euler (explicit)")
          15
                  plot!(t_vec,E2, label = "Backward Euler (implicit)")
                  display(plot!(t_vec,E3, label = "Implicit Midpoint",xlabel = "Time (s)", ylabel="Energy (J)"))
          16
          17
          18
                  # test energy behavior
          19
                  E0 = E1[1]
          20
                  @test 2.5 < (E1[end] - E0) < 3.0
          21
          22
                  @test -3.0 < (E2[end] - E0) < -2.5
          23
                  @test abs(E3[end] - E0) < 1e-2</pre>
                  @test abs(E0 - E4[end]) < 1e-4</pre>
          24
          25
                  @test abs(E0 - E5[end]) < 1e-1</pre>
                  @test abs(E0 - E6[end]) < 1e-4</pre>
          26
          27
              end
          28
```



Out[16]: Test.DefaultTestSet("energy behavior", Any[], 6, false, false)

6

energy behavior |

Another important takeaway from these integrators is that explicit Euler results in unstable behavior (as shown here by the growing energy), and implicit Euler results in artificial damping (losing energy). Implicit midpoint however maintains the correct energy. Even though the solution from implicit midpoint will vary from the initial energy, it does not move secularly one way or the other.

```
In [13]: import Pkg
    Pkg.activate(@__DIR__)
    Pkg.instantiate()
    using LinearAlgebra, Plots
    import ForwardDiff as FD
    using MeshCat
    using Test
    using Plots;
```

Activating environment at `~/ocrl ws/16-745/HW1 S23/Project.toml`

Q2: Equality Constrained Optimization (20 pts)

In this problem, we are going to use Newton's method to solve some constrained optimization problems. We will start with a smaller problem where we can experiment with Full Newton vs Gauss-Newton, then we will use these methods to solve for the motor torques that make a quadruped balance on one leg.

Part A (10 pts)

Here we are going to solve some equality-constrained optimization problems with Newton's method. We are given a problem

$$\min_{x} \quad f(x) \tag{1}$$

$$st \quad c(x) = 0 \tag{2}$$

Which has the following Lagrangian:

$$\mathcal{L}(x,\lambda) = f(x) + \lambda^T c(x),$$

and the following KKT conditions for optimality:

$$\nabla_x \mathcal{L} = \nabla_x f(x) + \left[\frac{\partial c}{\partial x} \right]^T \lambda = 0$$
 (3)

$$c(x) = 0 (4)$$

Which is just a root-finding problem. To solve this, we are going to solve for a $z=[x^T,\lambda]^T$ that satisfies these KKT conditions.

Newton's Method with a Linesearch

We use Newton's method to solve for when r(z)=0. To do this, we specify res_fx(z) as r(z), and res_jac_fx(z) as $\partial r/\partial z$. To calculate a Newton step, we do the following:

12/02/2023, 15:59

$$\Delta z = -iggl[rac{\partial r}{\partial z}iggr]^{-1} r(z_k)$$

We then decide the step length with a linesearch that finds the largest $\alpha \leq 1$ such that the following is true:

$$\phi(z_k + \alpha \Delta z) < \phi(z_k)$$

Where ϕ is a "merit function", or <code>merit_fx(z)</code> in the code. In this assignment you will use a backtracking linesearch where α is initialized as $\alpha=1.0$, and is divided by 2 until the above condition is satisfied.

NOTE: YOU DO NOT NEED TO (AND SHOULD NOT) USE A WHILE LOOP ANYWHERE IN THIS ASSIGNMENT.

```
In [14]: function linesearch(z::Vector, Δz::Vector, merit fx::Function;
                               max_ls_iters = 10)::Float64 # optional argument with a def
              # TODO: return maximum \alpha \le 1 such that merit f_X(z + \alpha * \Delta z) < merit <math>f_X(z)
              # with a backtracking linesearch (\alpha = \alpha/2 after each iteration)
              # NOTE: DO NOT USE A WHILE LOOP
              \alpha = 1.0
              for i = 1:max_ls_iters
                  # TODO: return \alpha when merit fx(z + \alpha * \Delta z) < merit <math>fx(z)
                  if merit_fx(z + \alpha*\Delta z) < merit_fx(z)
                       return \alpha
                  end
                  \alpha = \alpha/2
              error("linesearch failed")
          end
          function newtons_method(z0::Vector, res_fx::Function, res_jac_fx::Function, me
                                    tol = 1e-10, max iters = 50, verbose = false)::Vector{
              # TODO: implement Newton's method given the following inputs:
              # - z0, initial guess
              # - res fx, residual function
              # - res jac fx, Jacobian of residual function wrt z
              # - merit fx, merit function for use in linesearch
              # optional arguments
              # - tol, tolerance for convergence. Return when norm(residual)<tol
              # - max iter, max # of iterations
              # - verbose, bool telling the function to output information at each itera
              # return a vector of vectors containing the iterates
              # the last vector in this vector of vectors should be the approx. solution
              # NOTE: DO NOT USE A WHILE LOOP ANYWHERE
              # return the history of guesses as a vector
              Z = [zeros(length(z0)) for i = 1:max iters]
```

```
Z[1] = z0
    for i = 1:(max_iters - 1)
        # NOTE: everything here is a suggestion, do whatever you want to
        # TODO: evaluate current residual
        curr_r = res_fx(Z[i])
        norm_r = norm(curr_r) # TODO: update this
        if verbose
             print("iter: $i |r|: $norm r
        end
        # TODO: check convergence with norm of residual < tol
        # if converged, return Z[1:i]
        if norm r < tol</pre>
             return Z[1:i]
        end
        # TODO: caculate Newton step (don't forget the negative sign)
        \Delta Z = -res jac fx(Z[i]) \setminus curr r
        # TODO: linesearch and update z
        \alpha = linesearch(Z[i], \Delta Z, merit_fx)
        Z[i+1] = Z[i] + \alpha .* \Delta Z
        if verbose
             print("\alpha: $\alpha \n")
        end
    error("Newton's method did not converge")
end
```

Out[14]: newtons_method (generic function with 1 method)

```
α: 1.0
iter: 1
        |r|: 0.9995239729818045
iter: 2
         |r|: 0.9421342427117169
                               \alpha: 0.5
iter: 3
         |r|: 0.1753172908866053
                               \alpha: 1.0
iter: 4
         |r|: 0.0018472215879181287
                                  \alpha: 1.0
iter: 5
         |r|: 2.1010529101114843e-9
                                  \alpha: 1.0
iter: 6
         check Newton |
                2
```

Out[15]: Test.DefaultTestSet("check Newton", Any[], 2, false, false)

We will now use Newton's method to solve the following constrained optimization problem. We will write functions for the full Newton Jacobian, as well as the Gauss-Newton Jacobian.

```
Out[16]:
                                                            Cost Function
                                                                                                              constraint
                   1.0
                                                                                                       12
                                                                                                       10
                   0.5
                                                                                                       8
              \times
                   0.0
                                                                                                       6
                                                                                                       4
                  -0.5
                  -1.0
                                        -0.5
                                                         0.0
                                                                           0.5
                       -1.0
                                                                                            1.0
                                                         X_1
4
```

```
In [25]: # we will use Newton's method to solve the constrained optimization problem sho
function cost(x::Vector)
    Q = [1.65539     2.89376; 2.89376    6.51521]
    q = [2;-3]
    return 0.5*x'*Q*x + q'*x + exp(-1.3*x[1] + 0.3*x[2]^2)
end
function constraint(x::Vector)
    norm(x) - 0.5
end
# HINT: use this if you want to, but you don't have to
```

```
function constraint jacobian(x::Vector)::Matrix
    # since `constraint` returns a scalar value, ForwardDiff
    # will only allow us to compute a gradient of this function
    # (instead of a Jacobian). This means we have two options for
    # computing the Jacobian: Option 1 is to just reshape the gradient
    # into a row vector
    \# J = reshape(FD.gradient(constraint, x), 1, 2)
    # or we can just make the output of constraint an array,
    # where is this '_x' defined?
    constraint_array(_x) = [constraint(_x)]
    J = FD.jacobian(constraint_array, x)
    # assert the jacobian has # rows = # outputs
    # and # columns = # inputs
    @assert size(J) == (length(constraint(x)), length(x))
    return J
end
function kkt conditions(z::Vector)::Vector
    # TODO: return the KKT conditions
    x = z[1:2]
    \lambda = z[3:3]
    # TODO: return the stationarity condition for the cost function
    # and the primal feasibility
      error("kkt not implemented")
    primal feasibility = [constraint(x)]
    J = FD.gradient(cost, x)
    \delta c = constraint jacobian(x)
    stationarity = J + \delta c' * \lambda
    return [stationarity ;primal_feasibility]
end
function fn kkt jac(z::Vector)::Matrix
    # TODO: return full Newton Jacobian of kkt conditions wrt z
    x = z[1:2]
    \lambda = z[3:3]
    \beta = 1e-3
    # TODO: return full Newton jacobian with a 1e-3 regularizer
     error("fn kkt jac not implemented")
    primal feasibility = constraint(x)
    H = FD.hessian(x -> cost(x), x)
    δc = constraint_jacobian(x)
```

```
double derivative of L = H + FD.jacobian(x -> (constraint jacobian(x)' * \lambda
              kkt jacobian = [double derivative of L \delta c'; \delta c 0]
                    e = eigvals(kkt jacobian)
                    while !(sum(e .> 0) == length(x) \&\& sum(e .< 0) == length(\lambda))
#
                                 kkt \ jacobian = kkt \ jacobian + Diagonal([\beta*ones(length(x)); -\beta*ones(length(x)); -
#
                                 e = eigvals(kkt jacobian)
#
                    end
              kkt jacobian = kkt jacobian + Diagonal([\beta*ones(length(x)); -\beta*ones(length(x))]
              return kkt_jacobian
end
function gn_kkt_jac(z::Vector)::Matrix
             # TODO: return Gauss-Newton Jacobian of kkt conditions wrt z
             x = z[1:2]
             \lambda = z[3]
             \beta = 1e-3
            # TODO: return Gauss-Newton jacobian with a 1e-3 regularizer
                    error("gn kkt jac not implemented")
             primal feasibility = constraint(x)
            H = FD.hessian(x \rightarrow cost(x), x)
             \delta c = constraint jacobian(x)
             double_derivative_of_L = H
             gn_kkt_jacobian = [double_derivative_of_L δc'; δc 0]
                    e = eigvals(gn kkt jacobian)
                    while !(sum(e .> 0) == length(x) \&\& sum(e .< 0) == length(\lambda))
                                 gn kkt jacobian = gn kkt jacobian + Diagonal([\beta*ones(length(x)); -\beta*(length(x))]
                                 e = eigvals(gn kkt jacobian)
                    end
             gn_kkt_jacobian = gn_kkt_jacobian + Diagonal([\beta*ones(length(x)); -\beta*ones(length(x)); -\beta*ones(length(x)); -\beta*ones(length(x));
              return gn_kkt_jacobian
end
```

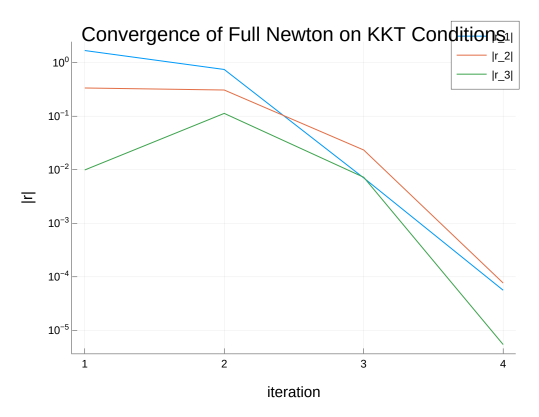
Out[25]: gn_kkt_jac (generic function with 1 method)

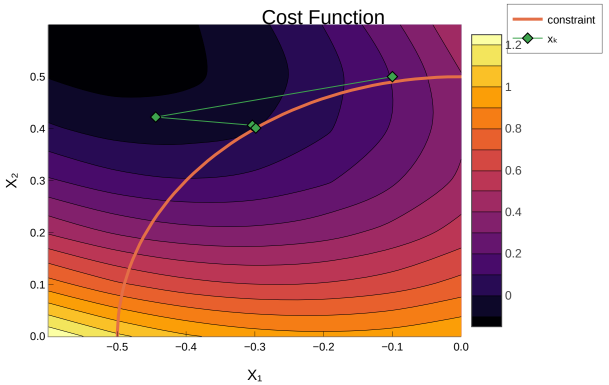
```
In [26]: @testset "Test Jacobians" begin

# first we check the regularizer
z = randn(3)
J_fn = fn_kkt_jac(z)
J_gn = gn_kkt_jac(z)

# check what should/shouldn't be the same between
@test norm(J_fn[1:2,1:2] - J_gn[1:2,1:2]) > 1e-10
@test abs(J_fn[3,3] + 1e-3) < 1e-10
@test abs(J_gn[3,3] + 1e-3) < 1e-10
@test norm(J_fn[1:2,3] - J_gn[1:2,3]) < 1e-10
@test norm(J_fn[3,1:2] - J_gn[3,1:2]) < 1e-10
end</pre>
```

```
Test Summary: | Pass Total
         Test Jacobians | 5
Out[26]: Test.DefaultTestSet("Test Jacobians", Any[], 5, false, false)
In [27]: @testset "Full Newton" begin
              z0 = [-.1, .5, 0] # initial guess
              merit fx(z) = norm(kkt conditions(z)) # simple merit function
              Z = newtons_method(z0, kkt_conditions, fn_kkt_jac, merit_fx; tol = 1e-4, max
              R = kkt conditions.(Z)
              # make sure we converged on a solution to the KKT conditions
             @test norm(kkt conditions(Z[end])) < 1e-4</pre>
             @test length(R) < 6</pre>
              # ------plotting stuff-----
             Rp = [[abs(R[i][ii]) + 1e-15 \text{ for } i = 1:length(R)] \text{ for } ii = 1:length(R[1])]
              plot(Rp[1],yaxis=:log,ylabel = "|r|",xlabel = "iteration",
                   yticks= [1.0*10.0^{(-x)} \text{ for } x = float(15:-1:-2)],
                   title = "Convergence of Full Newton on KKT Conditions", label = "|r 1|
              plot!(Rp[2], label = "|r 2|")
              display(plot!(Rp[3],label = "|r 3|"))
              contour(-.6:.1:0,0:.1:.6, (x1,x2)-> cost([x1;x2]),title = "Cost Function",
                      xlabel = "X_1", ylabel = "X_2", fill = true)
              xcirc = [.5*\cos(\theta) \text{ for } \theta \text{ in } range(0, 2*pi, length = 200)]
              ycirc = [.5*\sin(\theta) for \theta in range(0, 2*pi, length = 200)]
              plot!(xcirc, ycirc, lw = 3.0, xlim = (-.6, 0), ylim = (0, .6), label = "constant"
              z1 \text{ hist} = [z[1] \text{ for } z \text{ in } Z]
              z2 \text{ hist} = [z[2] \text{ for } z \text{ in } Z]
              display(plot!(z1 hist, z2 hist, marker = :d, label = "xk"))
                   -----plotting stuff-----
         end
         iter: 1
                     |r|: 1.7188450769812715
                                                α: 1.0
         iter: 2
                     |r|: 0.8150495962203247
                                                α: 1.0
         iter: 3
                     |r|: 0.025448943695826287
                                                 α: 1.0
         iter: 4
                     |r|: 9.501514353500914e-5
```





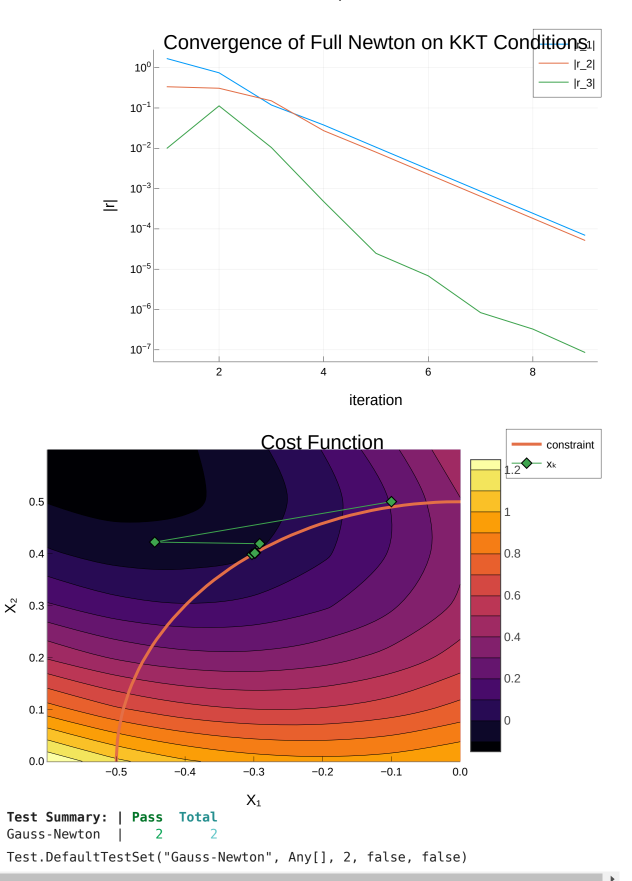
Test Summary: | Pass Total
Full Newton | 2 2

Out[27]: Test.DefaultTestSet("Full Newton", Any[], 2, false, false)

```
In [28]: @testset "Gauss-Newton" begin

z0 = [-.1, .5, 0] # initial guess
merit_fx(_z) = norm(kkt_conditions(_z)) # simple merit function
```

```
# the only difference in this block vs the previous is `gn kkt jac` instea
    Z = newtons_method(z0, kkt_conditions, gn_kkt_jac, merit_fx; tol = 1e-4, max
    R = kkt conditions.(Z)
    # make sure we converged on a solution to the KKT conditions
    @test norm(kkt conditions(Z[end])) < 1e-4</pre>
    @test length(R) < 10
    # -----plotting stuff-----
    Rp = [[abs(R[i][ii]) + 1e-15 \text{ for } i = 1:length(R)] \text{ for } ii = 1:length(R[1])]
    plot(Rp[1],yaxis=:log,ylabel = "|r|",xlabel = "iteration",
         yticks= [1.0*10.0^{(-x)} \text{ for } x = float(15:-1:-2)],
         title = "Convergence of Full Newton on KKT Conditions", label = "|r 1|
    plot!(Rp[2], label = "|r 2|")
    display(plot!(Rp[3], label = "|r_3|"))
    contour(-.6:.1:0,0:.1:.6, (x1,x2)-> cost([x1;x2]),title = "Cost Function",
            xlabel = "X_1", ylabel = "X_2", fill = true)
    xcirc = [.5*\cos(\theta) \text{ for } \theta \text{ in } range(0, 2*pi, length = 200)]
    ycirc = [.5*\sin(\theta) for \theta in range(0, 2*pi, length = 200)]
    plot!(xcirc,ycirc, lw = 3.0, xlim = (-.6, 0), ylim = (0, .6), label = "constant"
    z1_{hist} = [z[1] \text{ for } z \text{ in } Z]
    z2 \text{ hist} = [z[2] \text{ for } z \text{ in } Z]
    display(plot!(z1 hist, z2 hist, marker = :d, label = "xk"))
    # -----plotting stuff-----
end
iter: 1
           |r|: 1.7188450769812715
                                       \alpha: 1.0
iter: 2
           |r|: 0.8150495962203247
                                       α: 1.0
iter: 3
           |r|: 0.19186516708148574
                                        α: 1.0
iter: 4
           |r|: 0.04663490553083029
                                        α: 1.0
iter: 5
           |r|: 0.01332977842954523
                                        α: 1.0
iter: 6
          |r|: 0.0037714013578573355 \alpha: 1.0
iter: 7
                                         α: 1.0
           |r|: 0.001071165054782875
iter: 8
           |r|: 0.00030392210707413806
                                           \alpha: 1.0
iter: 9
           |r|: 8.625764141582568e-5
```



Part B (10 pts): Balance a quadruped

Now we are going to solve for the control input $u \in \mathbb{R}^{12}$, and state $x \in \mathbb{R}^{30}$, such that the quadruped is balancing up on one leg. First, let's load in a model and display the rough "guess" configuration that we are going for:

Open Controls

Now, we are going to solve for the state and control that get us a statically stable stance on just one leg. We are going to do this by solving the following optimization problem:

12/02/2023, 15:59

$$\min_{x,u} \quad \frac{1}{2} (x - x_{guess})^T (x - x_{guess}) + \frac{1}{2} 10^{-3} u^T u \tag{5}$$

$$st \quad f(x,u) = 0 \tag{6}$$

Where our primal variables are $x\in\mathbb{R}^{30}$ and $u\in\mathbb{R}^{12}$, that we can stack up in a new variable $y=[x^T,u^T]^T\in\mathbb{R}^{42}$. We have a constraint $f(x,u)=\dot{x}=0$, which will ensure the resulting configuration is stable. This constraint is enforced with a dual variable $\lambda\in\mathbb{R}^{30}$. We are now ready to use Newton's method to solve this equality constrained optimization problem, where we will solve for a variable $z=[y^T,\lambda^T]^T\in\mathbb{R}^{72}$.

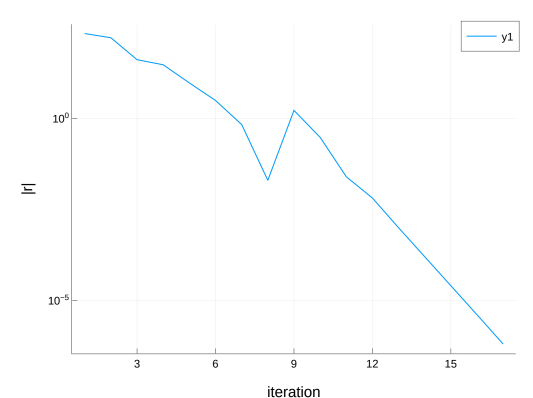
In this next section, you should fill out $quadruped_kkt(z)$ with the KKT conditions for this optimization problem, given the constraint is that dynamics(model, x, u) = zeros(30). When forming the Jacobian of the KKT conditions, use the Gauss-Newton approximation for the hessian of the Lagrangian (see example above if you're having trouble with this).

```
In [30]: # initial guess
          const x guess = initial state(model)
          # indexing stuff
          const idx x = 1:30
          const idx_u = 31:42
          const idx c = 43:72
          # I like stacking up all the primal variables in y, where y = [x;u]
          # Newton's method will solve for z = [x; u; \lambda], or z = [y; \lambda]
          function quadruped cost(y::Vector)
              # cost function
              @assert length(y) == 42
              x = y[idx x]
              u = y[idx u]
              return 0.5 * (x - x \text{ guess})' * (x - x \text{ guess}) + 0.5 * (10^-3) * u' * u
          end
          function quadruped constraint(y::Vector)::Vector
              # constraint function
              @assert length(y) == 42
              x = y[idx x]
              u = y[idx_u]
              return dynamics(model, x, u)
          end
          function quadruped kkt(z::Vector)::Vector
              @assert length(z) == 72
              x = z[idx_x]
              u = z[idx u]
              \lambda = z[idx c]
              y = [x;u]
              primal feasibility = quadruped constraint(y)
```

```
J = FD.gradient(quadruped cost, y)
             δc = FD.jacobian(y -> quadruped constraint(y), y)
             stationarity = J + \delta c' * \lambda
             return [stationarity ;primal_feasibility]
         end
         function quadruped kkt jac(z::Vector)::Matrix
             @assert length(z) == 72
             x = z[idx x]
             u = z[idx u]
             \lambda = z[idx c]
             \beta = 1e-5
             y = [x;u]
             H = FD.hessian(quadruped cost, y)
             δc = FD.jacobian( y -> quadruped constraint( y), y)
             double derivative of L = H
             return kkt jacobian
         end
         WARNING: redefinition of constant x guess. This may fail, cause incorrect answ
         ers, or produce other errors.
Out[30]: quadruped kkt jac (generic function with 1 method)
In [31]: function quadruped merit(z)
             # merit function for the quadruped problem
             @assert length(z) == 72
             r = quadruped kkt(z)
             return norm(r[1:42]) + 1e4*norm(r[43:end])
         end
         @testset "quadruped standing" begin
             z0 = [x_guess; zeros(12); zeros(30)]
             Z = newtons method(z0, quadruped kkt, quadruped kkt jac, quadruped merit;
             set configuration!(mvis, Z[end][1:state dim(model)÷2])
             R = norm.(quadruped kkt.(Z))
             display(plot(1:length(R), R, yaxis=:log,xlabel = "iteration", ylabel = "|r
             @test R[end] < 1e-6
             @test length(Z) < 25
             x,u = Z[end][idx_x], Z[end][idx_u]
             @test norm(dynamics(model, x, u)) < 1e-6</pre>
         end
```

02

```
α: 1.0
iter: 1
            |r|: 217.37236872332227
iter: 2
            |r|: 166.30172438182936
                                         \alpha: 1.0
iter: 3
            |r|: 41.47732635798011
                                        \alpha: 0.25
iter: 4
            |r|: 30.165366152039404
                                         \alpha: 1.0
iter: 5
            |r|: 9.547200505196036
                                        \alpha: 1.0
iter: 6
            |r|: 3.1522081784314824
                                         \alpha: 1.0
iter: 7
            |r|: 0.6883639538094173
                                         α: 1.0
iter: 8
            |r|: 0.020229959592048152
                                            \alpha: 1.0
iter: 9
            |r|: 1.6831359405438966
                                         \alpha: 1.0
iter: 10
             |r|: 0.30646762831705066
                                            \alpha: 1.0
iter: 11
             |r|: 0.02538589021613
                                        \alpha: 1.0
iter: 12
             |r|: 0.006601332797307665
                                             α: 1.0
iter: 13
             |r|: 0.0010077080791744822
                                              \alpha: 1.0
iter: 14
              |r|: 0.00016605797758123474
                                               \alpha: 1.0
iter: 15
             |r|: 2.5750782144045896e-5
                                              \alpha: 1.0
iter: 16
             |r|: 4.103870588678761e-6
                                             α: 1.0
             |r|: 6.439334108144757e-7
iter: 17
```



Out[31]: Test.DefaultTestSet("quadruped standing", Any[], 3, false, false)

```
In [32]: let

# let's visualize the balancing position we found

z0 = [x_guess; zeros(12); zeros(30)]
Z = newtons_method(z0, quadruped_kkt, quadruped_kkt_jac, quadruped_merit;
# visualizer
mvis = initialize_visualizer(model)
set_configuration!(mvis, Z[end][1:state_dim(model)÷2])
render(mvis)
```

end

 $_{\mbox{\sc \Gamma}}$ Info: MeshCat server started. You can open the visualizer by visiting the following URL in your browser:

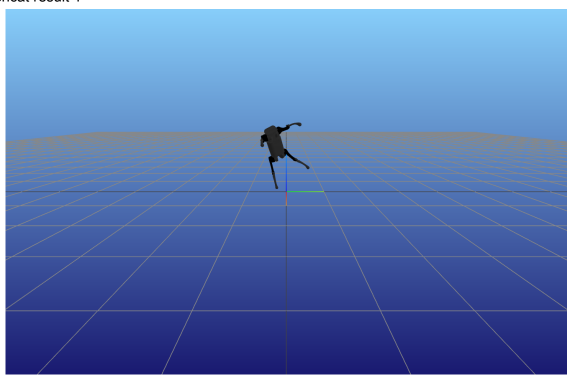
http://127.0.0.1:8708

Out[32]:

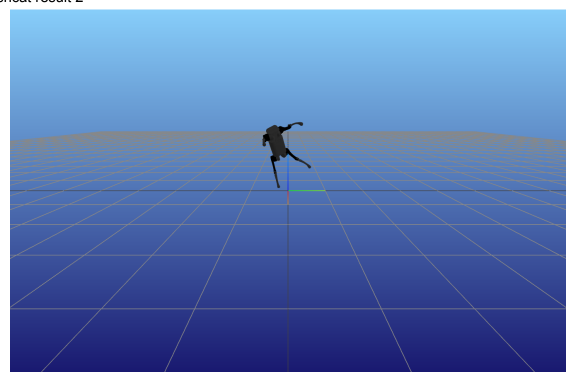
Open Controls

Meshcat plots were no rendering Q2 Part B hence attaching it here:

Meshcat result 1



Meshcat result 2



```
In [6]: import Pkg
   Pkg.activate(@__DIR__)
   Pkg.instantiate()
   using LinearAlgebra, Plots
   import ForwardDiff as FD
   using Printf
   using JLD2
```

Activating environment at `~/ocrl_ws/16-745/HW1_S23/Project.toml`

Q2 (20 pts): Augmented Lagrangian Quadratic Program Solver

Here we are going to use the augmented lagrangian method described here in a video, with the corresponding pdf here to solve the following problem:

$$\min_{x} \quad \frac{1}{2} x^T Q x + q^T x$$
 (1)

$$s.t. \quad Ax - b = 0 \tag{2}$$

$$Gx - h \le 0 \tag{3}$$

where the cost function is described by $Q\in\mathbb{R}^{n\times n}$, $q\in\mathbb{R}^n$, an equality constraint is described by $A\in\mathbb{R}^{m\times n}$ and $b\in\mathbb{R}^m$, and an inequality constraint is described by $G\in\mathbb{R}^{p\times n}$ and $h\in\mathbb{R}^p$.

By introducing a dual variable $\lambda \in \mathbb{R}^m$ for the equality constraint, and $\mu \in \mathbb{R}^p$ for the inequality constraint, we have the following KKT conditions for optimality:

$$Qx + q + A^T\lambda + G^T\mu = 0$$
 stationarity (4)

$$Ax - b = 0$$
 primal feasibility (5)

$$Gx - h \le 0$$
 primal feasibility (6)

$$\mu \ge 0$$
 dual feasibility (7)

$$\mu \circ (Gx - h) = 0$$
 complementarity (8)

where o is element-wise multiplication.

```
In [22]: # TODO: read below
# NOTE: DO NOT USE A WHILE LOOP ANYWHERE
"""
The data for the QP is stored in `qp` the following way:
     @load joinpath(@__DIR__, "qp_data.jld2") qp

which is a NamedTuple, where
     Q, q, A, b, G, h = qp.Q, qp.q, qp.A, qp.b, qp.G, qp.h

contains all of the problem data you will need for the QP.

Your job is to make the following function
```

```
x, \lambda, \mu = solve qp(qp; verbose = true, max iters = 100, tol = 1e-8)
You can use (or not use) any of the additional functions:
You can use (or not use) any of the additional functions:
You can use (or not use) any of the additional functions:
You can use (or not use) any of the additional functions:
as long as solve qp works.
function cost(qp::NamedTuple, x::Vector)::Real
    0.5*x'*qp.Q*x + dot(qp.q,x)
end
function c_eq(qp::NamedTuple, x::Vector)::Vector
    qp.A*x - qp.b
end
function h ineq(qp::NamedTuple, x::Vector)::Vector
    qp.G*x - qp.h
end
function mask matrix(qp::NamedTuple, x::Vector, μ::Vector, ρ::Real)::Matrix
      error("not implemented")
       possible point of failure maybe think of some diff method to calculate
    h = h_{ineq}(qp, x)
    Ip = zeros(length(h),length(h))
    for i = 1:length(h)
        if h[i] < 0 \&\& \mu[i] == 0
            I\rho[i,i] = 0
        else
            I\rho[i,i] = \rho
        end
    end
    return Ip
end
function lagrangian(qp::NamedTuple, x::Vector, λ::Vector, μ::Vector, ρ::Real):
    cx = c eq(qp, x)
    hx = h ineq(qp, x)
    return cost(qp ,x)+ \lambda' * cx + \mu' * hx
end
function augmented lagrangian(qp::NamedTuple, x::Vector, λ::Vector, μ::Vector,
#
#
      error("not implemented")
    cx = c_eq(qp, x)
    hx = h_{ineq}(qp, x)
    Iρ = mask matrix(qp , x , μ, ρ)
    lag = lagrangian(qp ,x , λ ,μ, ρ)
    return lag + 0.5 * \rho * cx' * cx + 0.5 * hx' * I\rho * hx
```

```
end
function augemented_lag_grad(qp::NamedTuple, x::Vector, λ::Vector, μ::Vector,
    Iρ = mask matrix(qp , x , μ, ρ)
    return qp.Q * x + qp.q + qp.A' * (\lambda + \rho * c eq(qp,x)) + qp.G' * (\mu + Ip)
end
function augemented lag hessian(qp::NamedTuple, x::Vector, λ::Vector, μ::Vecto
    Iρ = mask_matrix(qp , x , μ, ρ)
    return qp.Q + \rho * qp.A' * qp.A + qp.G' * Ip * qp.G
end
function stationarity(qp::NamedTuple, x::Vector, \lambda::Vector, \mu::Vector, \rho::Real
    return FD.gradient(x -> augmented_lagrangian(qp ,x , \lambda ,\mu, \rho), x)
#
           \delta cx = FD. jacobian(x-> c eq(qp, x), x)
           \delta hx = FD.jacobian(x-> h ineq(qp, x),x)
           return FD.gradient(x-> cost(qp ,x),x) + \delta cx'*\lambda + \delta hx'\mu
end
function logging(qp::NamedTuple, main iter::Int, AL gradient::Vector, x::Vecto
    # TODO: stationarity norm
    stationarity norm = norm(stationarity(qp, x, \lambda, \mu, \rho)) # fill this in
    @printf("%3d % 7.2e % 7.2e % 7.2e % 7.2e % 7.2e %5.0e\n",
           main_iter, stationarity_norm, norm(AL_gradient), maximum(h_ineq(qp,x)
           norm(c eq(qp,x),Inf), abs(dot(\mu,h ineq(qp,x))), \rho)
end
function kkt cond(qp::NamedTuple, x::Vector, \lambda::Vector, \mu::Vector, \rho::Real)
    cx = c eq(qp, x)
    hx = h_ineq(qp, x)
      return [stationarity(qp, x, \lambda, \mu, \rho) ; cx; hx; \mu; abs(dot(\mu, h_ineq(qp,x))
    return stationarity(qp, x, \lambda, \mu, \rho)
end
function newton_step(qp::NamedTuple, x::Vector, λ::Vector, μ::Vector, ρ::Real)
    kkt jacobian = FD.hessian(x -> augmented lagrangian(qp, x, \lambda, \mu, \rho), x)
    return -kkt jacobian \ kkt cond(qp,x,\lambda,\mu,\rho)
end
function solve qp(qp; verbose = true, max iters = 100, tol = 1e-8)
    x = zeros(length(qp.q))
    \lambda = zeros(length(qp.b))
    \mu = zeros(length(qp.h))
    \phi = 10
    \alpha = 1.0
    \rho = 1.0
    if verbose
        @printf "iter |\nabla L_{\times}| |\nabla AL_{\times}| max(h) |c|
                                                                           compl
        @printf "-----
    end
    # TOD0:
    for main_iter = 1:max_iters
```

03

```
if verbose
              logging(qp, main_iter, augemented_lag_grad(qp, x, \lambda, \mu, \rho), x, \lambda,
            return x, \lambda, \mu
         Use Newton method to calculate the change in x
         \Delta x = newton_step(qp, x, \lambda, \mu, \rho)
            skipping line search as it is specified that use \alpha = 1
            update x
         x = x + \alpha .* \Delta x
            update λ & μ
         \lambda = \lambda + \rho * c_eq(qp,x)
         \mu = \max.(0, (\mu + \rho * h\_ineq(qp ,x)))
         \rho = \rho * \phi
         # TODO: convergence criteria based on tol
         if norm(c eq(qp,x), Inf) < tol && max(0,maximum(h ineq(qp,x))) < tol
              return x, λ, μ
         end
    end
     error("qp solver did not converge")
end
let
    # example solving qp
    @load joinpath(@__DIR__, "qp_data.jld2") qp
     x, \lambda, \mu = solve qp(qp; verbose = true, tol = 1e-8)
end
                     |∇AL<sub>×</sub>|
iter
        |\nabla L_{\times}|
                                  max(h)
                                                | c |
                                                             compl
                                                                         ρ
       5.60e+01
                    5.60e+01
                               4.38e+00
                                               6.49e+00
                                                            0.00e+00
                                                                        1e+00
                                                                        1e+01
```

```
2
   7.50e+01
              7.50e+01
                         1.55e+00
                                     1.31e+00
                                                2.64e+00
3
   9.63e+01
              9.63e+01
                         2.96e-02
                                    3.04e-01
                                                4.74e-02
                                                         1e+02
   4.21e+01
              4.21e+01
                          6.37e-03
                                     1.35e-02
                                                7.39e-03
                                                         1e+03
5
   2.34e+03
              2.34e+03
                          6.84e-02
                                     1.55e-04
                                                4.67e+00
                                                          1e+04
   2.12e+03
              2.12e+03
                          2.12e-06
                                     3.74e-06
                                                2.71e-04
6
                                                         1e+05
   1.30e-01
               1.30e-01
                        -1.94e-08
                                     3.42e-08
                                                2.18e-08
                                                         1e+06
```

QP Solver test (10 pts)

```
In [23]: # 10 points
using Test
@testset "qp solver" begin
    @load joinpath(@__DIR__, "qp_data.jld2") qp
    x, \(\lambda\), \(\mu = \solve_qp(qp; \text{ verbose} = \text{ true}, \text{ max_iters} = 100, \text{ tol} = 1e-6)

@load joinpath(@__DIR__, "qp_solutions.jld2") qp_solutions
@test norm(x - qp_solutions.x,Inf)<1e-3;</pre>
```

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```
@test norm(λ - qp_solutions.λ,Inf)<1e-3;
@test norm(μ - qp_solutions.μ,Inf)<1e-3;
end</pre>
```

iter	\(\nabla \) L \(\times \)	\(\nabla \) L \(\)	max(h)	c	compl	ρ
1	5.60e+01 7.50e+01	5.60e+01 7.50e+01	4.38e+00 1.55e+00	6.49e+00 1.31e+00	0.00e+00 2.64e+00	1e+00 1e+01
3	9.63e+01	9.63e+01	2.96e-02	3.04e-01	4.74e-02	1e+02
4 5	4.21e+01 2.34e+03	4.21e+01 2.34e+03	6.37e-03 6.84e-02	1.35e-02 1.55e-04	7.39e-03 4.67e+00	1e+03 1e+04
6 Test	2.12e+03 Summary:	2.12e+03 Pass Total	2.12e-06	3.74e-06	2.71e-04	1e+05
qp solver 3 3						

Out[23]: Test.DefaultTestSet("qp solver", Any[], 3, false, false)

Simulating a Falling Brick with QPs

In this question we'll be simulating a brick falling and sliding on ice in 2D. You will show that this problem can be formulated as a QP, which you will solve using an Augmented Lagrangian method.

The Dynamics

The dynamics of the brick can be written in continuous time as

$$M\dot{v}+Mg=J^T\lambda$$
 where $M=mI_{2 imes2},\;g=\left[egin{array}{c}0\9.81\end{array}
ight],\;J=\left[egin{array}{c}0&1\end{array}
ight]$

and $\lambda \in \mathbb{R}$ is the normal force. The velocity $v \in \mathbb{R}^2$ and position $q \in \mathbb{R}^2$ are composed of the horizontal and vertical components.

We can discretize the dynamics with backward Euler: \$\$

$$\left[\begin{smallmatrix} v_{k+1} \\ q_{k+1} \end{smallmatrix} \right]$$

=

$$\left[egin{array}{c} v_k \ q_k \end{array}
ight]$$

• \Delta t \cdot

$$\left[egin{array}{c} rac{1}{m}J^T\lambda_{k+1}-g\ v_{k+1} \end{array}
ight]$$

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We also have the following contact constraints:

$$Jq_{k+1} \ge 0$$
 (don't fall through the ice) (9)

$$\lambda_{k+1} \ge 0$$
 (normal forces only push, not pull) (10)

$$\lambda_{k+1} J q_{k+1} = 0$$
 (no force at a distance) (11)

Q3

Part (a): QP formulation (5 pts)

Show that these discrete-time dynamics are equivalent to the following QP by writing down the KKT conditions.

minimize_{$$v_{k+1}$$} $\frac{1}{2}v_{k+1}^T M v_{k+1} + [M(\Delta t \cdot g - v_k)]^T v_{k+1}$ (12)

subject to
$$-J(q_k + \Delta t \cdot v_{k+1}) \le 0 \tag{13}$$

TASK: Write down the KKT conditions for the optimization problem above, and show that it's equivalent to the dynamics problem stated previously. Use LaTeX markdown.

PUT ANSWER HERE:

Comparing the above given cost function and constraints to the form

$$\min_{x} \quad \frac{1}{2}x^{T}Qx + q^{T}x \tag{14}$$

$$s.t. \quad Ax - b = 0 \tag{15}$$

$$Gx - h \le 0 \tag{16}$$

we have.

$$Q=M=mI_{2 imes2} \ x=v_{k+1} \ q_k=M(\Delta t\cdot g-v_k) \ G=-J\Delta t \ h=Jq_k$$

Since, there i-s not equality constraint Ax - b does not exist

The KKT conditions for the above are:

$$\frac{\delta L}{\delta x} = Qx + q + G^T \mu = 0$$
 stationarity (17)

(18)

$$Gx - h \le 0$$
 primal feasibility (19)
 $\mu \ge 0$ dual feasibility (20)

$$\mu \ge 0$$
 dual feasibility (20)

$$\mu \circ (Gx - h) = 0$$
 complementarity (21)

Now we will expand these conditions to retrive the discrete-time dynamics:

1. Expanding the stationarity:

$$Qx + q + G^T \mu = 0$$
 $Mv_{k+1} + M(\Delta t \cdot g - v_k) + (-J\Delta t)^T \mu = 0$

Q3

- a. since Δt is a scalar $(-J\Delta t)^T=-J^T\Delta t$
- b. setting $\mu=\lambda_{k+1}$ where $\lambda_{k+1}\geq 0$ (normal force only push, not pull) eq 11

$$egin{aligned} Mv_{k+1} + M(\Delta t \cdot g - v_k) + -J^T \Delta t \lambda_{k+1} &= 0 \ M(v_{k+1} + (\Delta t \cdot g - v_k)) + -J^T \Delta t \lambda_{k+1} &= 0 \ M(v_{k+1} + (\Delta t \cdot g - v_k)) &= J^T \Delta t \lambda_{k+1} \ (v_{k+1} + (\Delta t \cdot g - v_k)) &= M^{-1} J^T \Delta t \lambda_{k+1} \ v_{k+1} &= -(\Delta t \cdot g - v_k)) + M^{-1} J^T \Delta t \lambda_{k+1} \ v_{k+1} &= v_k + \Delta t \cdot (-g + M^{-1} J^T \lambda_{k+1}) \end{aligned}$$

Which is the dynamics equation 1

2. Expanding $Gx-h\leq 0$ - primal feasibility

$$egin{aligned} Gx-h &\leq 0 \ -J\Delta t v_{k+1} - Jq_k &\leq 0 \ J\Delta t v_{k+1} + Jq_k &\geq 0 \ J(\Delta t v_{k+1} + q_k) &\geq 0 \end{aligned}$$

setting $q_{k+1} = (\Delta t v_{k+1} + q_k)$ from the dynamics eq2

$$Jq_{k+1} \geq 0$$

Which gives us the constraint - don't fall through the ice eq 9

3. Expanding $\mu \circ (Gx - h) = 0$ - complementarity

$$\lambda_{k+1}\circ Jq_{k+1}=0$$
 - no force at a distance eq 11

Thus we can see that the KKT conditions are equivalent to the dynamics problem stated proviously.

Brick Simulation (5 pts)

In [12]: function brick_simulation_qp(q, v; mass = 1.0, Δt = 0.01)

```
Q3
    # TODO: fill in the QP problem data for a simulation step
    # fill in Q, q, G, h, but leave A, b the same
    # this is because there are no equality constraints in this qp
    g = [0; 9.81]
    J = [0 \ 1]
    M = mass * [1.0 0.0 ; 0.0 1.0]
    qp = (
        Q = M
        q = M * (\Delta t .* g - v),
        A = zeros(0,2), # don't edit this
        b = zeros(0), # don't edit this
        G = -J * \Delta t
        h = J * q
    )
    return qp
end
```

Out[12]: brick simulation qp (generic function with 1 method)

```
In [13]: @testset "brick qp" begin
           q = [1,3.0]
           v = [2, -3.0]
           qp = brick simulation qp(q,v)
           # check all the types to make sure they're right
           qp.0::Matrix{Float64}
           qp.q::Vector{Float64}
           qp.A::Matrix{Float64}
           qp.b::Vector{Float64}
           qp.G::Matrix{Float64}
           qp.h::Vector{Float64}
           (qp.Q) = (2,2)
           (q_1, q_2, q_3) = (2, q_4)
           (0,2)
           (0, 0) = (0, 0)
           (qp.G) = (1,2)
           @test size(qp.h) == (1,)
           (qp.Q) - 2 < 1e-10
           (qe.q - [-2.0, 3.0981]) < 1e-10
           (qp.G - [0 -.01]) < 1e-10
           (qp.h[1] -3) < 1e-10
        end
```

```
Test Summary: | Pass Total
brick qp
                  10
                        10
```

Out[13]: Test.DefaultTestSet("brick qp", Any[], 10, false, false)

```
In [46]: include(joinpath(@_DIR__, "animate_brick.jl"))
         function kkt brick(qp::NamedTuple, x::Vector, μ::Real)
```

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```
return qp.Q * x + qp.q + qp.G' * \mu
end
function brick newton step(qp::NamedTuple, x::Vector, μ::Real)
    kkt func(x) = [kkt brick(qp,x,\mu)]
    @show kkt_brick(qp,x,μ)
    @show qp
    @show x
    @show µ
    kkt jacobian = FD.jacobian(dx \rightarrow kkt func(dx), x)
    @show kkt jacobian(qp,x,\mu)
    return -kkt_jacobian \ kkt_brick(qp,x,μ)
end
let
    dt = 0.01
    T = 3.0
    t \text{ vec} = 0:dt:T
    N = length(t vec)
    qs = [zeros(2) for i = 1:N]
    vs = [zeros(2) for i = 1:N]
    qs[1] = [0, 1.0]
    vs[1] = [1, 4.5]
    mass = 1.0
    tol = 1e-5
    g = [0; 9.81]
    J = [0 \ 1]
    \lambda = 9.81
    # TODO: simulate the brick by forming and solving a qp
    # at each timestep. Your QP should solve for vs[k+1], and
    # you should use this to update qs[k+1]
    for k = 1:N-1
        Use Newton method to calculate the change in x
        brick qp = brick simulation qp(qs[k], vs[k])
        vs[k+1], _, \lambda = solve_qp(brick_qp; verbose = false, tol = 1e-8)
        qs[k+1] = qs[k] + vs[k+1]*dt
    end
    xs = [q[1] \text{ for } q \text{ in } qs]
    ys = [q[2] for q in qs]
    @show @test abs(maximum(ys)-2)<1e-1</pre>
    @show @test minimum(ys) > -1e-2
    @show @test abs(xs[end] - 3) < 1e-2
    xdot = diff(xs)/dt
    @show @test maximum(xdot) < 1.0001</pre>
    @show @test minimum(xdot) > 0.9999
    @show @test ys[110] > 1e-2
    @show @test abs(ys[111]) < 1e-2
```

#= In[46]:59 =# @test(ys[110] > 0.01) = Test Passed #= In[46]:60 =# @test(abs(ys[111]) < 0.01) = Test Passed #= In[46]:61 =# @test(abs(ys[112]) < 0.01) = Test Passed

```
@show @test abs(ys[112]) < le-2

display(plot(xs, ys, ylabel = "y (m)", xlabel = "x (m)"))

animate_brick(qs)

end

#= In[46]:52 =# @test(abs(maximum(ys) - 2) < 0.1) = Test Passed
#= In[46]:53 =# @test(minimum(ys) > -0.01) = Test Passed
#= In[46]:54 =# @test(abs(xs[end] - 3) < 0.01) = Test Passed
#= In[46]:57 =# @test(maximum(xdot) < 1.0001) = Test Passed
#= In[46]:58 =# @test(minimum(xdot) > 0.9999) = Test Passed
```

b' \n'

```
Info: MeshCat server started. You can open the visualizer by visiting the fo
llowing URL in your browser:
http://127.0.0.1:8702
```

