Implementation of an Iterative Linear Quadratic Regulator (iLQR)

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Problem statement

The iLQR algorithm

Our implementation

Demonstration time

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General formulation

Dynamics function:

$$x_{t+1} = f(x_t, u_t)$$

- Goal: minimize a quadratic cost function
- Cost function:

$$J(u) = \sum_{t=0}^{T-1} \left(x_t^{\top} Q x_t + u_t^{\top} R u_t \right) + \frac{1}{2} (x_T - x^*)^{\top} Q_f(x_T - x^*)$$

- Q: state cost matrix
- Q_f : final state cost matrix
- R: control cost matrix

Example: Simple Pendulum

- State: $x = [\theta \ \dot{\theta}]$
- Control: u, torque applied to the pendulum
- Dynamics: physical laws (simulator)
- Target: $x = [0 \ 0]$
- Cost function:

$$J(u) = \frac{1}{2} \left(\theta_f^2 + \dot{\theta}_f^2 \right) + \frac{1}{2} \int_0^T r u^2(t) dt$$

corresponding to $Q_f = I_2$, $Q = 0_2$, $R = rI_1$

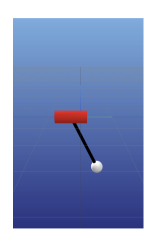


Example: Cartpole

- State: $x = [y \ \theta \ \dot{y} \ \dot{\theta}]$
- Control: u, force applied to the cart
- Dynamics: physical laws (simulator)
- Target: $x = [0 \ 0 \ 0 \ 0]$
- Cost function:

$$J(u) = \frac{1}{2} \left(\theta_f^2 + \dot{\theta}_f^2 + y_f^2 + \dot{y}_f^2 \right) + \frac{1}{2} \int_0^T r u^2(t) dt$$

corresponding to $Q_f = I_4$, $Q = 0_4$, $R = rI_1$



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General idea

- iLQR is an iterative algorithm
- Start with an initial trajectory
- Iteratively improve it using a local linear approximation
- Stop when the trajectory converges

Linearizing the dynamics

The equation $x_{t+1} = f(x_t, u_t)$ is linearized (at each step) as:

$$\delta x_{t+1} = A_t \delta x_t + B_t \delta u_t$$

with:

- A_t : Jacobian of f with respect to x evaluated at (x_t, u_t)
- B_t : Jacobian of f with respect to u evaluated at (x_t, u_t)

We are in LQR (Linear Quadratic Regulator, cf. TP5) setup!

Trajectory refinement using LQR

- 1. Forward pass: compute the successive states (x_t) for the current controls (u_t) , and the corresponding cost J
- 2. **Backward pass**: compute the gains, i.e. how much we should change the controls in each direction to minimize the cost
- 3. Forward rollout: apply the gains to the controls to obtain a new trajectory
- 4. Repeat until convergence

Computing the Jacobians

Finite differences method

We want to compute:

- $A_t=rac{\partial f}{\partial x}(x_t,u_t)$, i.e. how much the state at time t+1 changes when we slightly change the state at time t
- $B_t=rac{\partial f}{\partial u}(x_t,u_t)$, i.e. how much the state at time t+1 changes when we slightly change the control at time t

In a black box setting, we can use finite differences:

$$[A_t]_i \approx \frac{f(x_t + \varepsilon e_i, u_t) - f(x_t - \varepsilon e_i, u_t)}{2\varepsilon}$$
$$[B_t]_i \approx \frac{f(x_t, u_t + \varepsilon e_i) - f(x_t, u_t - \varepsilon e_i)}{2\varepsilon}$$

for some small ε and the canonical basis (e_i)

Tricks for practical convergence

• Gradient clipping: limit the size of the control updates norm to α to avoid divergence

$$\delta u_t = \frac{\delta u_i}{\max\left(1, \frac{\|\delta u_i\|}{\alpha}\right)}$$

 Gaussian initialization: start with a small random control sequence instead of a zero sequence

$$u_t \sim \mathcal{N}(0, \Sigma)$$

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What language to use?

Python
 Easy to use
 Fast
 Support for many libraries
 Not very funny
 Embarrassingly slow

Therefore, we chose to have a Rust core with Python bindings



From Rust to Python, and the other way around

- Instantiate the solver in Python
- Use Python libraries to define the dynamics
- The Rust solver does the computations, and calls the Python dynamics function

API Basic usage

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```
def dynamics(x, u):
    return ... # simulator
Q = np.zeros((state_dim, state_dim)) # state cost
Qf = np.eye(state dim) # final state cost
R = 1e-5 * np.eve(control_dim) # control cost (minimize the energy)
s = ilgr.ILQRSolver(state dim, control dim, Q, Qf, R)
target = np.zeros(state dim) # upright pendulum with no velocity
output = s.solve(np.concatenate((q0, v0)), target, dynamics, time steps=N,
                 gradient_clip=10.0, # max norm of the gradient
                 initialization=0.5) # std of the Gaussian initialization
```

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