Ordinary Differential Equations





Ordinary Differential Equations

An Ordinary Differential Equation is any equation in the form:

$$\dot{y} = f(y, t)$$

- \blacksquare Where y is the state variable
- lacktriangleright ...And f is a function, providing the gradient of the state variable

The peculiarities:

- y is actually a function or the t variable
- The t variable typically (but not always) represents time
- ...Hence y(t) is the state at time t
- ullet The gradient f depends on both the current state and current time

Ordinary = does not feature partial derivatives





Initial Value Problem

An Initial Value Problem consists of an ODE and a initial condition

$$\dot{y} = f(y, t)$$
$$y(0) = y_0$$

- This can be interpreted as running a simulation
- Given that the initial state y(0) is y_0 , how will the state unfold?

Initial values problem can be solved (a.k.a. integrated):

Exactly, using symbolic approaches, e.g.

$$\dot{y} = a, y(0) = b \Rightarrow y(t) = ay + b$$

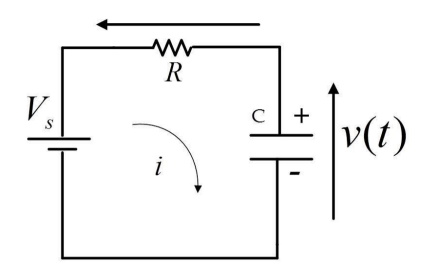
- This is the method considered in typical calculus courses
- ...Or approximately, via numerical approach





An Example

As an example, let's consider a simple RC circuit



It's dynamic behavior is described by the ODE:

$$\dot{V} = \frac{1}{\tau}(V_s - V)$$

• Where $\tau = RC$





Euler Method

The simplest numerical approach for ODEs is called Euler Method

This is obtained by:

- Considering a fixed sequence of evaluation points $\{t_k\}_{k=0}^n$
- Using a linear approximation for y(t) within each interval $[t_k, t_{k+1}]$
- lacksquare Approximating the slope with the gradient at time t_k

The pseudo code of the method consists of a single loop

• for k = 1..n:

$$y_k = y_{k-1} + (t_k - t_{k-1}) f(y_{k-1}, t_{k-1})$$

The output is a sequence $\{y_k\}_{i=0}^n$

- y_k is the state at time t_k
- y_0 is also an input for the algorithm





Euler Method for the RC Circuit

A typical Initial Value Problem solver API requires to define

The function characterizing the equation, i.e. f(y, t):

```
In [2]: tau, Vs = 8, 12
f = lambda y, t: 1./tau * (Vs - y)
```

The initial state y_0 and the evaluation points $\{t_i\}_{i=0}^n$

```
In [3]: y0 = (0,) # We start from an empty capacitor t = np.linspace(0, 40, 12)
```

Then we can call the solver itself (the code is in the util module)

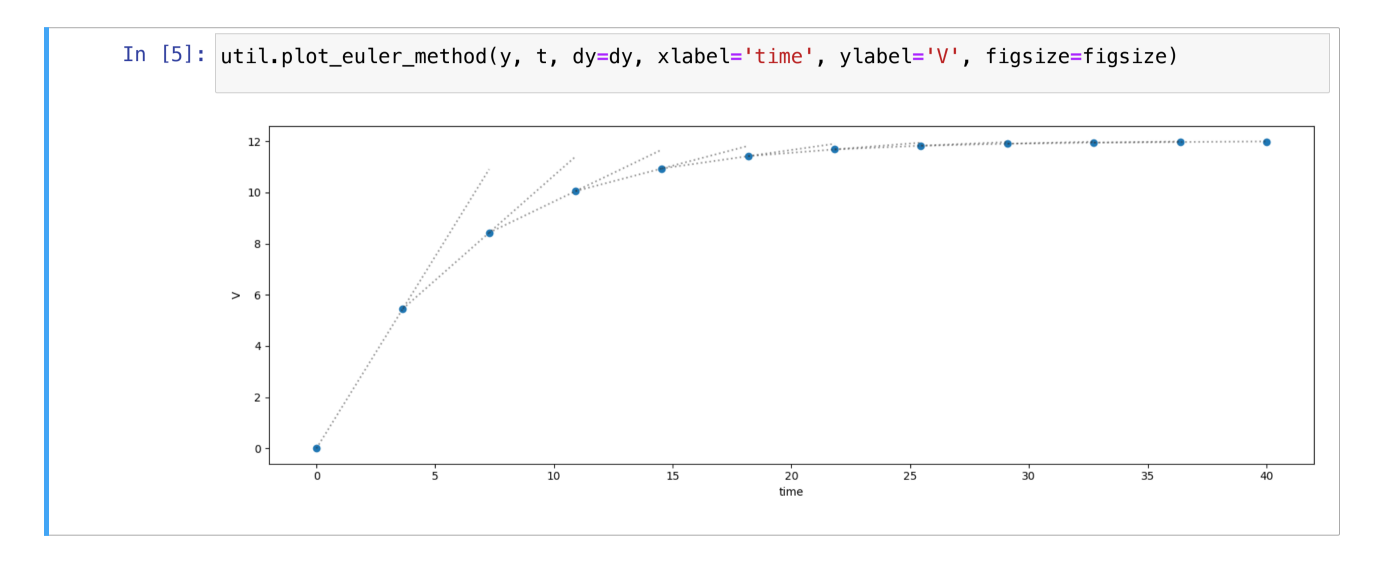
```
In [4]:
y, dy = util.euler_method(f, y0, t, return_gradients=True)
```





Euler Method for the RC Circuit

Visually, the method works as follows:



- The dots represent evaluated states
- The slope of the lines corresponds to the gradient at each step

ODE Integration Methods

Euler method is the simplest ODE integration approach

- ...But also one of the worst in terms of accuracy
- This is due to errors in the local approximation
- ...And forces to use very small steps to obtain high-quality results

There are many alternative integration methods

Some examples include:

- Backward Euler method
 - Like Euler method, but we use the gradient at the next state
 - In practice it requires to solve a (typically non-linear) equation
- Runge-Kutta methods
 - It's a family of method (Euler method is the simplest version)
 - They combine multiple gradients to obtain a local slope





"Learning" ODEs





"Learning" ODEs

The parameters of an ODE can be estimated from data

Formally, training amounts to solving:

$$\operatorname{argmin}_{\omega} \left\{ L(\hat{y}(t), y) \mid \dot{\hat{y}} = f(\hat{y}, t; \theta), \dot{y}(0) = y_0 \right\}$$

Where:

- $\{t_k\}_{k=0}^n$ is a sequence of points for which measurements are available
- $\{y_k\}_{k=0}^n$ are the corresponding state measurements
- f is a parameterized gradient function
- lacksquare L is a loss function (e.g. the classical MSE)

Intuitively, we require the integrated ODE to be close to the real one

- The goal is to choose the parameters (e.g. au, V_s) so as to achieve this





"Learning" ODEs

A viable approach is to "discretize, then optimize"

...Which can be be done by relying on an automatic differentiation engine

- First we solve the initial value problem using a numerical method
 - ...Making sure we evaluate every point in $\{t_k\}$
- lacksquare Then, we compute the loss $oldsymbol{L}$
- ...And view the whole process as a single compute graph

Then optimization over heta can be performed (e.g.) via gradient descent

This is possible since every integration step is differentiable

This is true for Euler method, but also for other (better) integration methods

- In particular, it's doable for the whole Runge-Kutta family
- ...But a bit more complicated in implicit methods (e.g. backward Euler)





Building Our Ground Truth

We'll see an example using our simple RC circuit

Let's start by building a high-quality ground truth sequence

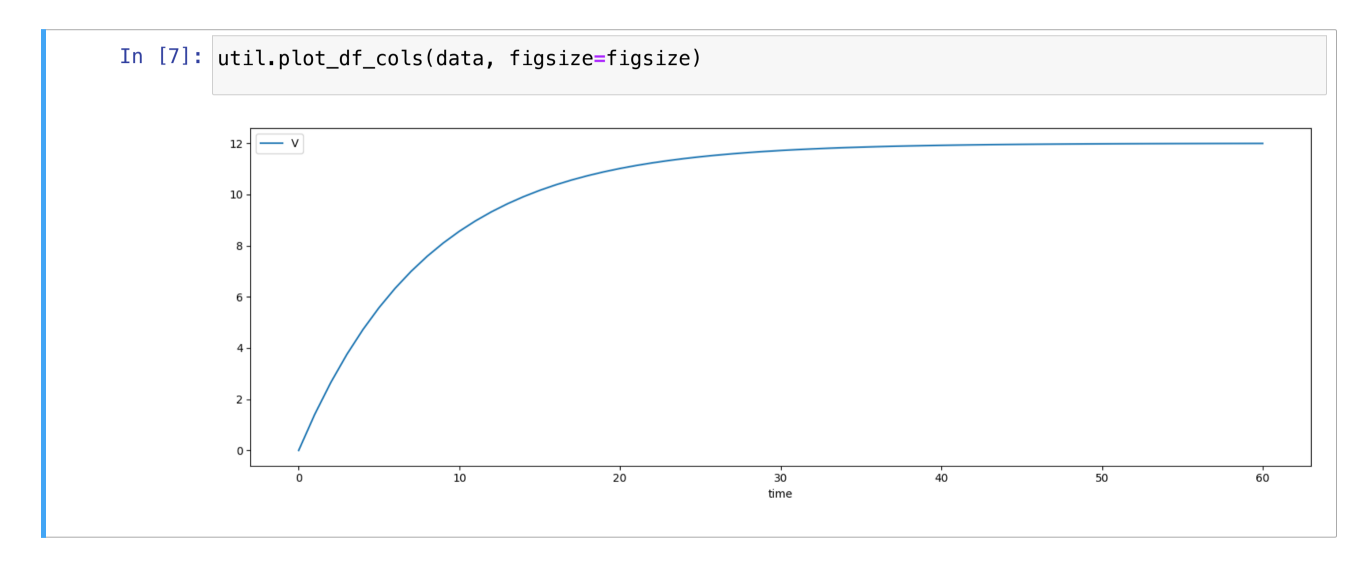
- We will use the odeint solver from scikit learn for this
- The code can be found in the **simulate_RC** function



steps_per_unit defines how many evaluations to perform per unit of time

Building Our Ground Truth

Let' check (visually) that the result is smooth enough



We need this step since we treat this as the ground truth



Outline of the Approach

We will go for a simple, but relatively general approach

- We will view the (parameterized) gradient function $f(y, t; \theta)$ as a layer type
- ...And we will use a keras. Model to encode Euler method, i.e.

$$\hat{y}(t_k) = \hat{y}(t_{k-1}) + (t_k - t_{k-1})f(\hat{y}(t_{k-1}), t_{k-1}; \theta)$$

- Each step of the method can be viewed as layer instance
- ...And all instances share the same weights

In terms of input/output:

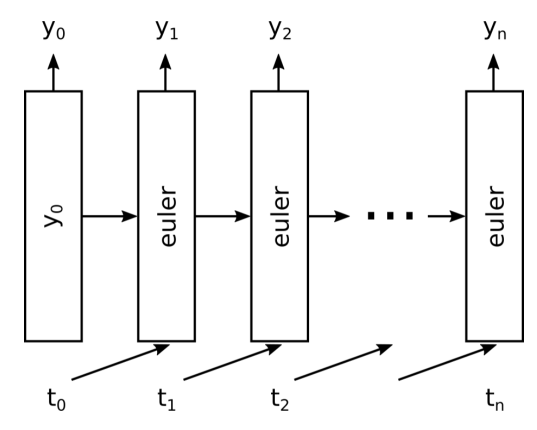
- The initial state corresponds to the input
- ...And a secondary input is given by the sequence $\{t_k\}_{k=0}^n$
- The output is the state for each evaluation step





Outline of the Approach

Overall, our "architecture" looks like this:



- lacktriangle The input includes the initial state y_0 and the evaluation points $\{t_k\}_{k=0}^n$
- The output consists of the sequence of state evaluations $\{\hat{y}_k\}_{k=0}^n$

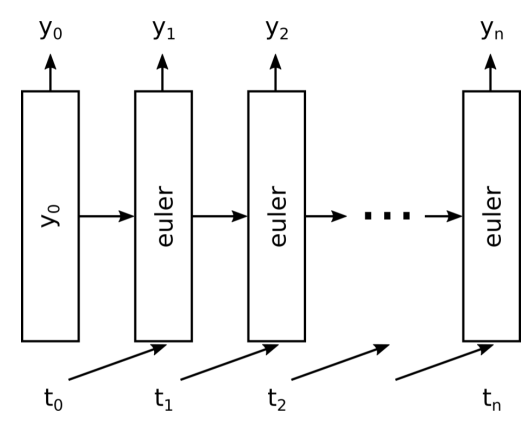
Overall, the signature is analogous to that of an ODE solver





Outline of the Approach

Overall, our "architecture" looks like this:



- Each "example" corresponds to a distinct integration of the same system
- ...And the architecture is very similar to a recurrent NN
- In particular, the "depth" grows with the number of evaluation points





Details Matter

In our RC circuit case, we have:

argmin_{$$\omega$$} $L(\hat{y}(t), y)$
subject to $\dot{\hat{y}} = \frac{1}{\tau}(V_s - \hat{y})$
 $\hat{y}(0) = y_0$

lacktriangle Where the parameters to be learned are au and V_s

There are a few details we need to account for

- For both parameters, negative values make no sense
- Moreover, since we plan to use gradient descent for training
- ...We need to make sure that our initial guesses are reasonable





Details Matter

We can meet both conditions by adopting the reformulation:

$$au = \sigma_{ au} e^{ heta_{ au}} \ V_{ ext{ iny S}} = \sigma_{V_{ ext{ iny S}}} e^{ heta_{V_{ ext{ iny S}}}}$$

Where the parameters to be learned are now $heta_{ au}$ and $heta_{V_s}$

- Using an exponential ensures we get non-negative values
- lacktriangledown The scaling factors $oldsymbol{\sigma}_{ au}$ and $oldsymbol{\sigma}_{V_s}$ are user-provided
 - They lead to reasonable guesses for typical NN weight initiliazers

There are just a few mild downsides:

- The exponential may lead to numerical issues in edge cases
- lacktriangle We need to have a rough idea of the scale of au and V_s





RC Circuit Layer

The layer for the RC circuit gradient is in the RCNablaLayer class

```
class RCNablaLayer(keras.layers.Layer):
    def build(self, input_shape):
        self.logtau = self.add_weight(
                shape=(1, ),
                initializer = 'random_normal', trainable=True,
        self.logvs = self.add_weight(
                shape=(1, ),
                initializer = 'random_normal', trainable=True,
```

■ In the __init__ method we take care of weight initialization





RC Circuit Layer

The layer for the RC circuit gradient is in the RCNablaLayer class

```
class RCNablaLayer(keras.layers.Layer):
    . . .
    def get_tau(self):
        return tf.math.exp(self.logtau) * self.tau_ref
    def get_vs(self):
        return tf.math.exp(self.logvs) * self.vs_ref
    def call(self, inputs):
        y, t = inputs # unpack the inputs
        return 1. / self.get_tau() * (self.get_vs() - y)
```

- ullet We use dedicated method to obtain au and V_s
- In the call method we compute the (ODE) gradient

Euler Method Model

The model for the Euler method is in the ODEEulerModel class

```
class ODEEulerModel(keras.Model):
    def __init__(self, f, **params): ...
    def call(self, inputs, training=False):
        y, T = inputs # unpack
        res = [y] # initial state
        for i in range(T.shape[1]-1):
            t, nt = T[:, i:i+1], T[:, i+1:i+2] # t_k and t_{k+1}
            dy = self.f([y, t], training=training) # gradient
            y = y + (nt - t) * dy # next state
            res.append(y) # store result
        res = tf.stack(res, axis=1) # concatenate
        return res
```

The __call__ method implements the method using tensor operators





Euler Method Model

The model for the Euler method is in the ODEEulerModel class

```
class ODEEulerModel(keras.Model):
    ...

def train_step(self, data):
    (y0, T), yt = data # unpack
    with tf.GradientTape() as tape:
        y = self.call([y0, T], training=True) # ODE integration
        # Loss computation
        mask = ~keras.ops.isnan(yt)
        loss = self.compute_loss(x=None, y=yt[mask], y_pred=y[mask])
    ...
```

- The loss is computed as usual on all available measurements
- We can exclude points by setting the corresponding target to NaN





Training Set

We have a single sequence of measurements

...Therefore, just a training set (no validation, no test)

Our first input is the initial state:

```
In [8]: tr_y0 = np.array(data.iloc[0]).reshape(1, -1); display(tr_y0)
array([[0.]])
```

The second is the sequence of evaluation points (time steps)





Training Set

We have a single sequence of measurements

...Therefore, just a training set (no validation, no test)

Then we need to prepare our ground truth

■ This is the sequence of all measurements, with the first state "masked"

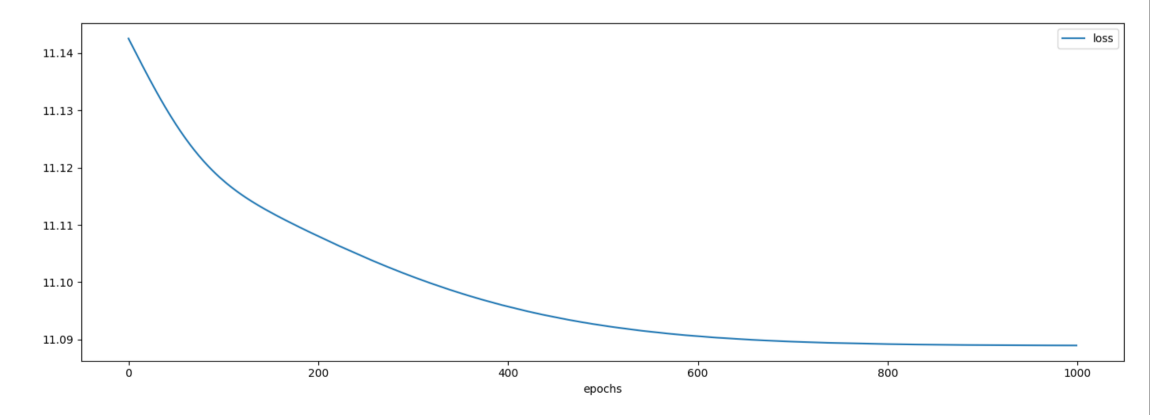




Training Process

We can now build and train the model

```
In [14]: %%time
    dRC = util.RCNablaLayer(tau_ref=10, vs_ref=10)
    euler = util.ODEEulerModel(dRC)
    history = util.train_ml_model(euler, [tr_y0, tr_T], tr_y, validation_split=0.0, epochs=1000,
    util.plot_training_history(history, figsize=figsize)
```







Model loss: 11.0890 (training)

CPU times: user 7.96 s, sys: 2.36 s, total: 10.3 s

Some Considerations

It seems to be working! But there are a few issues

First, the convergence is slow

Stopping before ~500 epochs leads to less stable results

Second, we cannot use a validation set:

■ This is due to the fact that we have a single sequence

Third, we are still not getting the correct parameters:

```
In [15]: print(f'tau: {tau:.2f} (real), {dRC.get_tau().numpy()[0]:.2f} (estimated)')
    print(f'Vs: {Vs:.2f} (real), {dRC.get_vs().numpy()[0]:.2f} (estimated)')

    tau: 8.00 (real), 8.51 (estimated)
    Vs: 12.00 (real), 12.00 (estimated)
```

In the next section, we will see how to address these issues



