1 covid19

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1 COVID19

In the previous assignment you have developed a particle based simulation of disease spreading. One infected individual was infecting other individuals when meeting them. The individuals were then recovering after a certain time of being infected and becoming immune to further infections. We now have a look at a model of disease spreading, which describes the same problem in terms of three coupled differential equations. This is the Kermack-McKendrick Model.

```
[12]: import numpy as np
      import matplotlib.pyplot as plt
      from scipy.sparse import diags
      from scipy.integrate import odeint
      import requests
      import io
      import pandas as pd
      %matplotlib inline
      %config InlineBackend.figure_format = 'retina'
      plt.rcParams.update({'font.size': 12,
                            'axes.titlesize': 18.
                            'axes.labelsize': 16,
                            'axes.labelpad': 14,
                            'lines.linewidth': 1,
                            'lines.markersize': 10,
                            'xtick.labelsize' : 16,
                            'ytick.labelsize' : 16,
                            'xtick.top' : True,
                            'xtick.direction' : 'in',
                            'ytick.right' : True,
                            'ytick.direction' : 'in',})
```

1.1 The Kermack-McKendrick Model

The Kermack-McKendrick model (KM) was an early compartmental model. It divides the population into three groups: Susceptible (S), Infected (I), and Removed (R). These are often referred to as **SIR** models. In the simple model below, there is, however, no population removed by death or so, but we will implement the removal as well.

Kermack and McKendrick built their model to deepen their understanding of epidemics where an infected individual enters into a population and suddenly begins spreading the disease. They assumed that the population remains constant (apart from deaths which are represented by the R term in the model), so ignore any effects from birth, non-disease related death, migration in or out of the target area. Moreover, all of the individuals are equally susceptible, so no bonus points for those who are healthier or have better hygiene habits in this model. All of these are limitations of the model, but also make it a great starting point for epidemiology studies.

1.1.1 Model Equation

The differential equations of the model are simple rate equations for the susceptibel x, the infected y and the recovered z population. There are two rate constants in the model, one for getting infected k_I and one for recovering k_R . If you look at the equations below, you see that the number of people removed from the suceptible population per timestep is just the infection rate times the number of people that can be infected x times the number of people who are infected y. That seems to be clear as none can be infected if there none to infect and none the can infect others.

$$\dot{x} = -k_I x y \tag{1}$$

$$\dot{y} = k_I x y - k_B y \tag{2}$$

$$\dot{z} = k_R y \tag{3}$$

1.2 Setup

Here we define again our initial parameters and the initial conditions. We would like to run the simulation for

- n = 1000 people
- i = 1 infected people at the beginning
- N = 200 timesteps of the simulation, i.e. days

We chose the individual rates the following way

- $k_I = 0.0002$ the rate of infection, multiplication with n gives the number of people infected by one individual per day
- $k_R = 0.1$ the rate of recovery, this means here 10 days for recovery
- $k_D = 0.05$ the rate of death, if we include death as well

These rates are just guesses (well I tried earlier), but you might want to map the earlier particle based simulation onto this model to compare the dynamics observed both.

```
[25]: n=1000000 # total people
i=1 # infected people
state=[n-i,i,0] # initial state
N=200

t=np.linspace(0,200,N)
```

1.3 Definition

Our covid19 function now implements the differential equations above into a function. This is rather straight forward, as they are only first order differential equations.

```
[32]: def covid19(state , time):
    k_I=0.5/n ## infection rate
    k_R=1/10 ## recovery rate
    k_D=0.05 ## death rate
    g0 = -k_I*state[0]*state[1]
    g1 = k_I*state[0]*state[1]-k_R*state[1] - k_D*state[1]
    g2 = k_R*state[1]
    return(np.array([g0, g1, g2]))
```

1.4 Solution

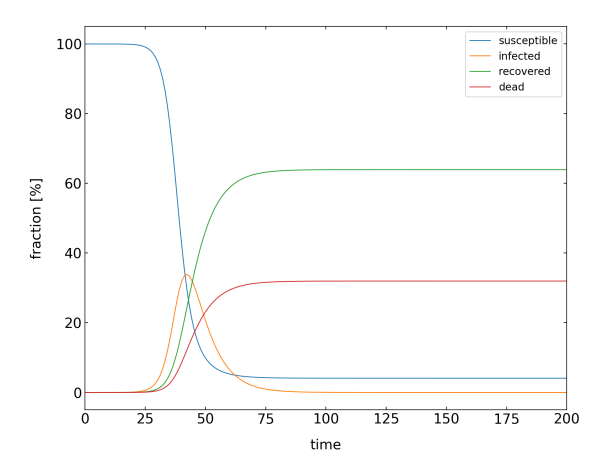
The following code solves the coupled differential equations and sorts the solution into individual arrays for further plotting.

```
[33]: answer = odeint( covid19, state , t )
susceptible=answer[:,0]
infected=answer[:,1]
recovered=answer[:,2]
dead=n-np.sum(answer[:,:],axis=1)
```

1.5 Plotting

We can now plot the solution of the model. You see that after some delay, the number of infected people quickly increases until the recovery sets in. Then the number of infected people decreases again until all are recovered. You may change the above rate of death now to see how it effects the diagram. You can now play around and check out what happens if you increase the recovery or infection rate.

```
[34]: percent=100/n
   plt.figure(figsize=(10,8))
   plt.plot(t,susceptible*percent,label='susceptible')
   plt.plot(t,infected*percent,label='infected')
   plt.plot(t,recovered*percent,label='recovered')
   plt.plot(t,dead*percent,label='dead')
   plt.xlabel('time')
   plt.ylabel('fraction [%]')
   plt.legend()
   plt.xlim(0,200)
   plt.show()
```



1.6 Real COVID19 numbers

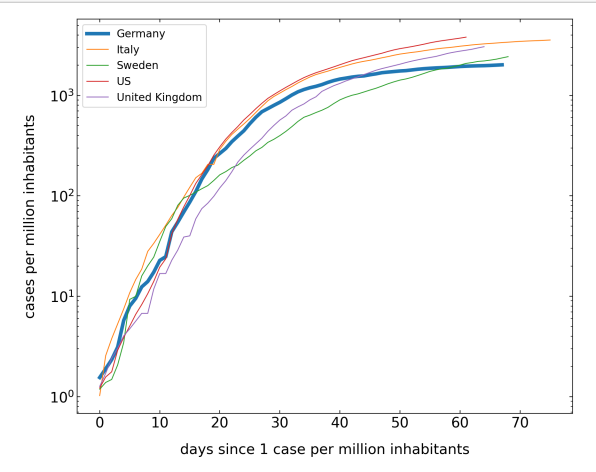
To compare the behavior to the current pandemic, I have downloaded some the current data for 5 different countries and stored them in the covid19.csv file. The data file reports **Date**, **Country**, **Confirmed**, **Recovered** and **Deaths**. To make the data comparable, I have already normalized the cases to the population of the country and multiplied by 1 million. That means the data is showsing the number of cases per million.

1.6.1 Total number of cases

To analyze the current situation we want to plot the total number of cases. To make the data more comparable, we do the following calculations. We first normalize the number of cases to the total population of the country. I have done that already in the dataset as mentioned above. The second issue is, that one should compare the data of different countries, when the total number of cases per million inhabitants crosses one for example. That is the only useful way to compare data. Note that this is a cummulative number.

```
[19]: plt.figure(figsize=(10,8))
    for c in cut.Country.unique():
        df=cut[cut.Country==c]
        cases=df[df.Confirmed>1].Confirmed.values
        if c=='Germany':
            lw=4
        else:
            lw=1
        plt.semilogy(cases,label=c,lw=lw);

plt.legend()
    plt.xlabel('days since 1 case per million inhabitants')
    plt.ylabel('cases per million inhabitants')
    plt.show()
```



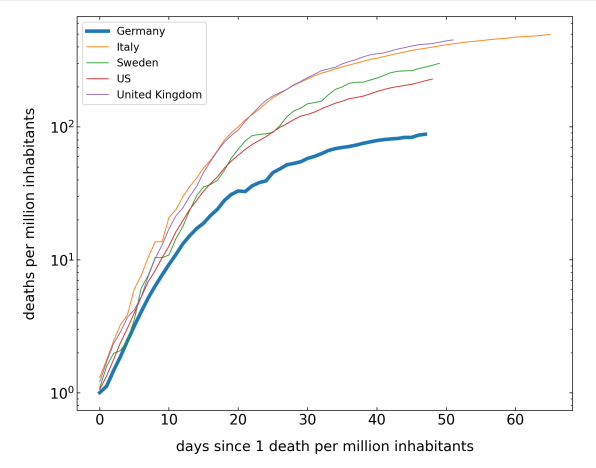
1.6.2 Number of Deaths

The graph below does the same analysis as the one for the confimed cases, but only for the number of death. While the differences in the cumulative numbers of confirmed cases are small, the

number of death per million in the countries show already some differences in the total death. Yet the kinetics is still similar.

```
[23]: plt.figure(figsize=(10,8))
    for c in cut.Country.unique():
        df=cut[cut.Country==c]
        cases=df[df.Deaths>1].Deaths.values
        if c=='Germany':
            lw=4
        else:
            lw=1
        plt.semilogy(cases,label=c,lw=lw);

plt.legend()
    plt.xlabel('days since 1 death per million inhabitants')
    plt.ylabel('deaths per million inhabitants')
    plt.show()
```

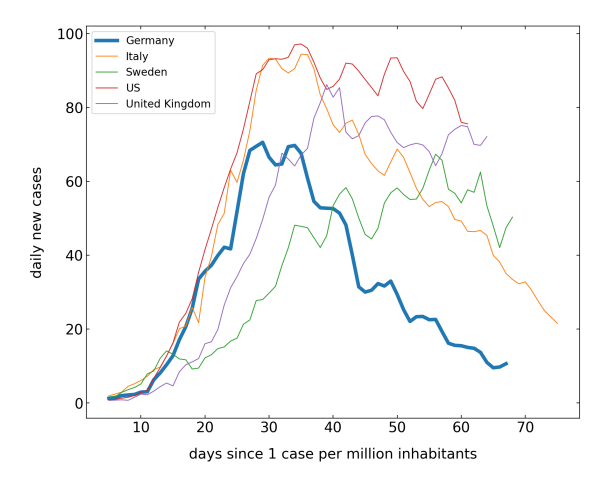


1.6.3 New cases per day

The next plot is comparing the new cases that appear per million people each day. Have a look at how this is done with the help of pandas. The data would be rather "noisy" as this is just the derivative of the confirmed cases curve. To smooth that out, I have included a rolling average. This rolling average takes the average of 5 following days and the moves one day forward each time. You may recognize the different dynamics in the countries

```
[24]: plt.figure(figsize=(10,8))
    for c in cut.Country.unique():
        df=cut[cut.Country==c]
        cases=df[df.Confirmed>1].Confirmed
        new_cases=cases.diff(1).rolling(5).mean()
        if c=='Germany':
            lw=4
        else:
            lw=1
        #plt.plot(cases.diff().values,label=c);
        plt.plot(new_cases.values,label=c,lw=lw)

plt.legend()
    plt.xlabel('days since 1 case per million inhabitants')
    plt.ylabel('daily new cases')
    plt.show()
```



1.6.4 Current cases

The last plot is the number of current active cases. This can be calculated from the total number of cases when we subtract the recovered and the death. This curve should roughly correspond in its shape to the model above. Think about, how the model is different from the real data.

```
[334]: plt.figure(figsize=(10,8))
for c in cut.Country.unique():
    df=cut[cut.Country==c]
    cases=df[df.Confirmed>1].Confirmed.values-df[df.Confirmed>1].Recovered.
    values-df[df.Confirmed>1].Deaths.values
    if c=='Germany':
        lw=4
    else:
        lw=1
    plt.plot(cases,label=c,lw=lw);

plt.legend()
plt.xlabel('days since 1 case per million inhabitants')
```

```
plt.ylabel('active cases per million inhabitants')
plt.show()
```

