

We now turn our attention to solving mathematical problems through computer programming. There are many reasons to choose integration as our first application. Integration is well known already from high school mathematics. Most integrals are not tractable by pen and paper, and a computerized solution approach is both very much simpler and much more powerful – you can essentially treat all integrals  $\int_a^b f(x)dx$  in 10 lines of computer code (!). Integration also demonstrates the difference between exact mathematics by pen and paper and *numerical mathematics* on a computer. The latter approaches the result of the former without any worries about rounding errors due to finite precision arithmetics in computers (in contrast to differentiation, where such errors prevent us from getting a result as accurate as we desire on the computer). Finally, integration is thought of as a somewhat difficult mathematical concept to grasp, and programming integration should greatly help with the understanding of what integration is and how it works. Not only shall we understand how to use the computer to integrate, but we shall also learn a series of good habits to ensure your computer work is of the highest scientific quality. In particular, we have a strong focus on how to write Python code that is free of programming mistakes.

Calculating an integral is traditionally done by

$$\int_a^b f(x) dx = F(b) - F(a), \quad (3.1)$$

where

$$f(x) = \frac{dF}{dx}.$$

The major problem with this procedure is that we need to find the *anti-derivative*  $F(x)$  corresponding to a given  $f(x)$ . For some relatively simple integrands  $f(x)$ , finding  $F(x)$  is a doable task, but it can very quickly become challenging, even impossible!

The method (3.1) provides an *exact* or *analytical* value of the integral. If we relax the requirement of the integral being exact, and instead look for *approximate* values, produced by *numerical methods*, integration becomes a very straightforward task for any given  $f(x)$  (!).

The downside of a numerical method is that it can only find an approximate answer. Leaving the exact for the approximate is a mental barrier in the beginning, but remember that most real applications of integration will involve an  $f(x)$  function that contains physical parameters, which are measured with some error. That is,  $f(x)$  is very seldom exact, and then it does not make sense to compute the integral with a smaller error than the one already present in  $f(x)$ .

Another advantage of numerical methods is that we can easily integrate a function  $f(x)$  that is only known as *samples*, i.e., discrete values at some  $x$  points, and not as a continuous function of  $x$  expressed through a formula. This is highly relevant when  $f$  is measured in a physical experiment.

### 3.1 Basic Ideas of Numerical Integration

We consider the integral

$$\int_a^b f(x)dx. \quad (3.2)$$

Most numerical methods for computing this integral split up the original integral into a sum of several integrals, each covering a smaller part of the original integration interval  $[a, b]$ . This re-writing of the integral is based on a selection of *integration points*  $x_i$ ,  $i = 0, 1, \dots, n$  that are distributed on the interval  $[a, b]$ . Integration points may, or may not, be evenly distributed. An even distribution simplifies expressions and is often sufficient, so we will mostly restrict ourselves to that choice. The integration points are then computed as

$$x_i = a + ih, \quad i = 0, 1, \dots, n, \quad (3.3)$$

where

$$h = \frac{b-a}{n}. \quad (3.4)$$

Given the integration points, the original integral is re-written as a sum of integrals, each integral being computed over the sub-interval between two consecutive integration points. The integral in (3.2) is thus expressed as

$$\int_a^b f(x)dx = \int_{x_0}^{x_1} f(x)dx + \int_{x_1}^{x_2} f(x)dx + \dots + \int_{x_{n-1}}^{x_n} f(x)dx. \quad (3.5)$$

Note that  $x_0 = a$  and  $x_n = b$ .

Proceeding from (3.5), the different integration methods will differ in the way they approximate each integral on the right hand side. The fundamental idea is that each term is an integral over a small interval  $[x_i, x_{i+1}]$ , and over this small interval, it makes sense to approximate  $f$  by a simple shape, say a constant, a straight line, or a parabola, which we can easily integrate by hand. The details will become clear in the coming examples.

**Computational example** To understand and compare the numerical integration methods, it is advantageous to use a specific integral for computations and graphical illustrations. In particular, we want to use an integral that we can calculate by hand such that the accuracy of the approximation methods can easily be assessed. Our specific integral is taken from basic physics. Assume that you speed up your car from rest and wonder how far you go in  $T$  seconds. The distance is given by the integral  $\int_0^T v(t)dt$ , where  $v(t)$  is the velocity as a function of time. A rapidly increasing velocity function might be

$$v(t) = 3t^2 e^{t^3}. \quad (3.6)$$

The distance after one second is

$$\int_0^1 v(t)dt, \quad (3.7)$$

which is the integral we aim to compute by numerical methods. Fortunately, the chosen expression of the velocity has a form that makes it easy to calculate the anti-derivative as

$$V(t) = e^{t^3} - 1. \quad (3.8)$$

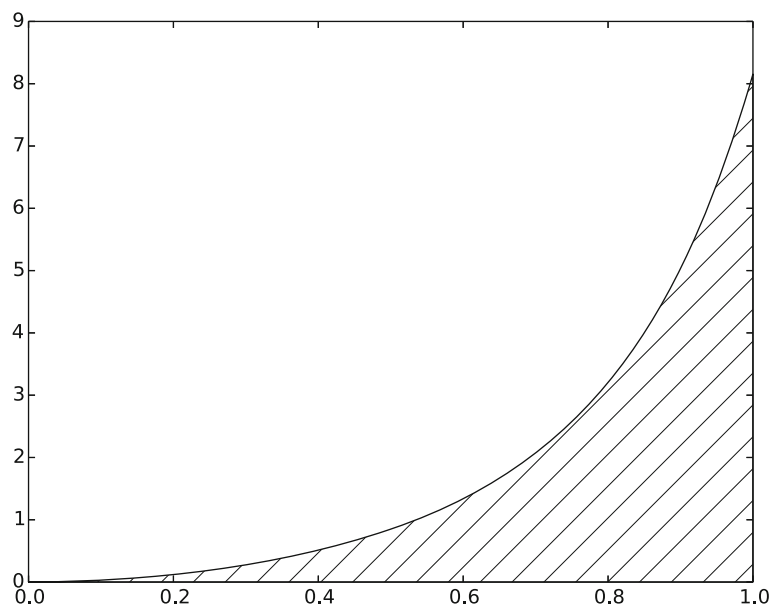
We can therefore compute the exact value of the integral as  $V(1) - V(0) \approx 1.718$  (rounded to 3 decimals for convenience).

---

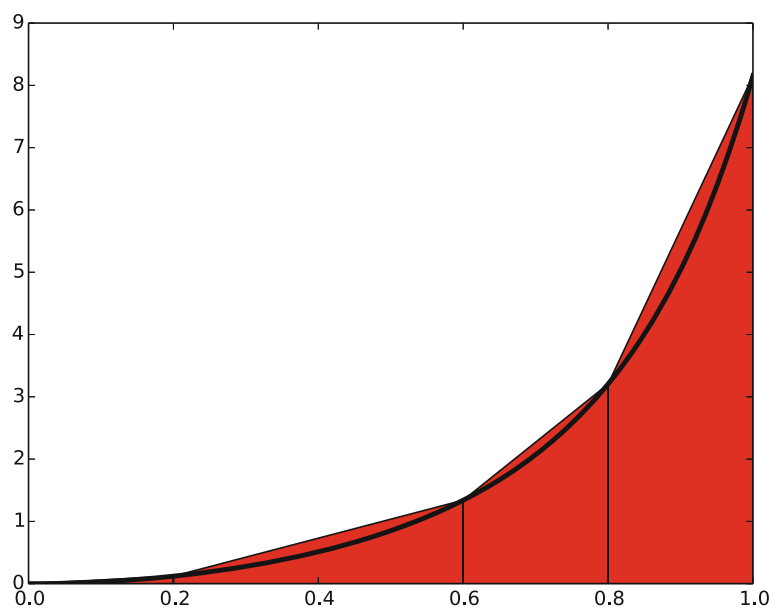
## 3.2 The Composite Trapezoidal Rule

The integral  $\int_a^b f(x)dx$  may be interpreted as the area between the  $x$  axis and the graph  $y = f(x)$  of the integrand. Figure 3.1 illustrates this area for the choice (3.7). Computing the integral  $\int_0^1 f(t)dt$  amounts to computing the area of the hatched region.

If we *replace* the true graph in Fig. 3.1 by a set of straight line segments, we may view the area rather as composed of trapezoids, the areas of which are easy to compute. This is illustrated in Fig. 3.2, where 4 straight line segments give rise to 4 trapezoids, covering the time intervals  $[0, 0.2)$ ,  $[0.2, 0.6)$ ,  $[0.6, 0.8)$  and  $[0.8, 1.0]$ . Note that we have taken the opportunity here to demonstrate the computations with time intervals that differ in size.



**Fig. 3.1** The integral of  $v(t)$  interpreted as the area under the graph of  $v$



**Fig. 3.2** Computing approximately the integral of a function as the sum of the areas of the trapezoids

The areas of the 4 trapezoids shown in Fig. 3.2 now constitute our approximation to the integral (3.7):

$$\begin{aligned} \int_0^1 v(t)dt \approx & h_1 \left( \frac{v(0) + v(0.2)}{2} \right) + h_2 \left( \frac{v(0.2) + v(0.6)}{2} \right) \\ & + h_3 \left( \frac{v(0.6) + v(0.8)}{2} \right) + h_4 \left( \frac{v(0.8) + v(1.0)}{2} \right), \end{aligned} \quad (3.9)$$

where

$$h_1 = (0.2 - 0.0), \quad (3.10)$$

$$h_2 = (0.6 - 0.2), \quad (3.11)$$

$$h_3 = (0.8 - 0.6), \quad (3.12)$$

$$h_4 = (1.0 - 0.8) \quad (3.13)$$

With  $v(t) = 3t^2e^{t^3}$ , each term in (3.9) is readily computed and our approximate computation gives

$$\int_0^1 v(t)dt \approx 1.895. \quad (3.14)$$

Compared to the true answer of 1.718, this is off by about 10 %. However, note that we used just 4 trapezoids to approximate the area. With more trapezoids, the approximation would have become better, since the straight line segments in the upper trapezoid side then would follow the graph more closely. Doing another hand calculation with more trapezoids is not too tempting for a lazy human, though, but it is a perfect job for a computer! Let us therefore derive the expressions for approximating the integral by an arbitrary number of trapezoids.

### 3.2.1 The General Formula

For a given function  $f(x)$ , we want to approximate the integral  $\int_a^b f(x)dx$  by  $n$  trapezoids (of equal width). We start out with (3.5) and approximate each integral on the right hand side with a single trapezoid. In detail,

$$\begin{aligned} \int_a^b f(x) dx &= \int_{x_0}^{x_1} f(x)dx + \int_{x_1}^{x_2} f(x)dx + \dots + \int_{x_{n-1}}^{x_n} f(x)dx, \\ &\approx h \frac{f(x_0) + f(x_1)}{2} + h \frac{f(x_1) + f(x_2)}{2} + \dots \\ &\quad + h \frac{f(x_{n-1}) + f(x_n)}{2} \end{aligned} \quad (3.15)$$

By simplifying the right hand side of (3.15) we get

$$\int_a^b f(x) dx \approx \frac{h}{2} [f(x_0) + 2f(x_1) + 2f(x_2) + \dots + 2f(x_{n-1}) + f(x_n)] \quad (3.16)$$

which is more compactly written as

$$\int_a^b f(x) dx \approx h \left[ \frac{1}{2} f(x_0) + \sum_{i=1}^{n-1} f(x_i) + \frac{1}{2} f(x_n) \right]. \quad (3.17)$$

#### Composite integration rules

The word *composite* is often used when a numerical integration method is applied with more than one sub-interval. Strictly speaking then, writing, e.g., “the trapezoidal method”, should imply the use of only a single trapezoid, while “the composite trapezoidal method” is the most correct name when several trapezoids are used. However, this naming convention is not always followed, so saying just “the trapezoidal method” may point to a single trapezoid as well as the composite rule with many trapezoids.

### 3.2.2 Implementation

**Specific or general implementation?** Suppose our primary goal was to compute the specific integral  $\int_0^1 v(t)dt$  with  $v(t) = 3t^2e^{t^3}$ . First we played around with a simple hand calculation to see what the method was about, before we (as one often does in mathematics) developed a general formula (3.17) for the general or “abstract” integral  $\int_a^b f(x)dx$ . To solve our specific problem  $\int_0^1 v(t)dt$  we must then apply the general formula (3.17) to the given data (function and integral limits) in our problem. Although simple in principle, the practical steps are confusing for many because the notation in the abstract problem in (3.17) differs from the notation in our special problem. Clearly, the  $f$ ,  $x$ , and  $h$  in (3.17) correspond to  $v$ ,  $t$ , and perhaps  $\Delta t$  for the trapezoid width in our special problem.

#### The programmer’s dilemma

1. Should we write a special program for the special integral, using the ideas from the general rule (3.17), but replacing  $f$  by  $v$ ,  $x$  by  $t$ , and  $h$  by  $\Delta t$ ?
2. Should we implement the general method (3.17) as it stands in a general function `trapezoid(f, a, b, n)` and solve the specific problem at hand by a specialized call to this function?

#### Alternative 2 is always the best choice!

The first alternative in the box above sounds less abstract and therefore more attractive to many. Nevertheless, as we hope will be evident from the examples, the second alternative is actually the simplest *and* most reliable from both a mathematical and programming point of view. These authors will claim that the second

alternative is the essence of the power of mathematics, while the first alternative is the source of much confusion about mathematics!

**Implementation with functions** For the integral  $\int_a^b f(x)dx$  computed by the formula (3.17) we want the corresponding Python function `trapezoid` to take any  $f$ ,  $a$ ,  $b$ , and  $n$  as input and return the approximation to the integral.

We write a Python function `trapezoidal` in a file `trapezoidal.py` as close as possible to the formula (3.17), making sure variable names correspond to the mathematical notation:

```
def trapezoidal(f, a, b, n):
    h = float(b-a)/n
    result = 0.5*f(a) + 0.5*f(b)
    for i in range(1, n):
        result += f(a + i*h)
    result *= h
    return result
```

**Solving our specific problem in a session** Just having the `trapezoidal` function as the only content of a file `trapezoidal.py` automatically makes that file a module that we can import and test in an interactive session:

```
>>> from trapezoidal import trapezoidal
>>> from math import exp
>>> v = lambda t: 3*(t**2)*exp(t**3)
>>> n = 4
>>> numerical = trapezoidal(v, 0, 1, n)
>>> numerical
1.9227167504675762
```

Let us compute the exact expression and the error in the approximation:

```
>>> V = lambda t: exp(t**3)
>>> exact = V(1) - V(0)
>>> exact - numerical
-0.20443492200853108
```

Is this error convincing? We can try a larger  $n$ :

```
>>> numerical = trapezoidal(v, 0, 1, n=400)
>>> exact - numerical
-2.1236490512777095e-05
```

Fortunately, many more trapezoids give a much smaller error.

**Solving our specific problem in a program** Instead of computing our special problem in an interactive session, we can do it in a program. As always, a chunk of code doing a particular thing is best isolated as a function even if we do not see any future reason to call the function several times and even if we have no need for arguments to parameterize what goes on inside the function. In the present case,

we just put the statements we otherwise would have put in a main program, inside a function:

```
def application():
    from math import exp
    v = lambda t: 3*(t**2)*exp(t**3)
    n = input('n: ')
    numerical = trapezoidal(v, 0, 1, n)

    # Compare with exact result
    V = lambda t: exp(t**3)
    exact = V(1) - V(0)
    error = exact - numerical
    print 'n=%d: %.16f, error: %g' % (n, numerical, error)
```

Now we compute our special problem by calling `application()` as the only statement in the main program. Both the `trapezoidal` and `application` functions reside in the file `trapezoidal.py`, which can be run as

---

Terminal

---

```
Terminal> python trapezoidal.py
n: 4
n=4: 1.9227167504675762, error: -0.204435
```

---

### 3.2.3 Making a Module

When we have the different pieces of our program as a collection of functions, it is very straightforward to create a *module* that can be imported in other programs. That is, having our code as a module, means that the `trapezoidal` function can easily be reused by other programs to solve other problems. The requirements of a module are simple: put everything inside functions and let function calls in the main program be in the so-called *test block*:

```
if __name__ == '__main__':
    application()
```

The if test is true if the module file, `trapezoidal.py`, is run as a program and false if the module is imported in another program. Consequently, when we do `from trapezoidal import trapezoidal` in some file, the test fails and `application()` is not called, i.e., our special problem is not solved and will not print anything on the screen. On the other hand, if we run `trapezoidal.py` in the terminal window, the test condition is positive, `application()` is called, and we get output in the window:

---

Terminal

---

```
Terminal> python trapezoidal.py
n: 400
n=400: 1.7183030649495579, error: -2.12365e-05
```

---



### 3.2.4 Alternative Flat Special-Purpose Implementation

Let us illustrate the implementation implied by alternative 1 in the *Programmer's dilemma* box in Sect. 3.2.2. That is, we make a special-purpose code where we adapt the general formula (3.17) to the specific problem  $\int_0^1 3t^2 e^{t^3} dt$ .

Basically, we use a for loop to compute the sum. Each term with  $f(x)$  in the formula (3.17) is replaced by  $3t^2 e^{t^3}$ ,  $x$  by  $t$ , and  $h$  by  $\Delta t$ <sup>1</sup>. A first try at writing a plain, flat program doing the special calculation is

```
from math import exp

a = 0.0; b = 1.0
n = input('n: ')
dt = float(b - a)/n

# Integral by the trapezoidal method
numerical = 0.5*3*(a**2)*exp(a**3) + 0.5*3*(b**2)*exp(b**3)
for i in range(1, n):
    numerical += 3*((a + i*dt)**2)*exp((a + i*dt)**3)
numerical *= dt

exact_value = exp(1**3) - exp(0**3)
error = abs(exact_value - numerical)
rel_error = (error/exact_value)*100
print 'n=%d: %.16f, error: %g' % (n, numerical, error)
```

The problem with the above code is at least three-fold:

1. We need to reformulate (3.17) for our special problem with a different notation.
2. The integrand  $3t^2 e^{t^3}$  is inserted many times in the code, which quickly leads to errors.
3. A lot of edits are necessary to use the code to compute a different integral – these edits are likely to introduce errors.

The potential errors involved in point 2 serve to illustrate how important it is to use Python functions as mathematical functions. Here we have chosen to use the lambda function to define the integrand as the variable  $v$ :

```
from math import exp

v = lambda t: 3*(t**2)*exp(t**3) # Function to be integrated
a = 0.0; b = 1.0
n = input('n: ')
dt = float(b - a)/n
```

---

<sup>1</sup> Replacing  $h$  by  $\Delta t$  is not strictly required as many use  $h$  as interval also along the time axis. Nevertheless,  $\Delta t$  is an even more popular notation for a small time interval, so we adopt that common notation.

```
# Integral by the trapezoidal method
numerical = 0.5*v(a) + 0.5*v(b)
for i in range(1, n):
    numerical += v(a + i*dt)
numerical *= dt

F = lambda t: exp(t**3)
exact_value = F(b) - F(a)
error = abs(exact_value - numerical)
rel_error = (error/exact_value)*100
print 'n=%d: %.16f, error: %g' % (n, numerical, error)
```

Unfortunately, the two other problems remain and they are fundamental.

Suppose you want to compute another integral, say  $\int_{-1}^{1.1} e^{-x^2} dx$ . How much do we need to change in the previous code to compute the new integral? Not so much:

- the formula for  $v$  must be replaced by a new formula
- the limits  $a$  and  $b$
- the anti-derivative  $V$  is not easily known<sup>2</sup> and can be omitted, and therefore we cannot write out the error
- the notation should be changed to be aligned with the new problem, i.e.,  $t$  and  $dt$  changed to  $x$  and  $h$

These changes are straightforward to implement, but *they are scattered around in the program*, a fact that requires us to be very careful so we do not introduce new programming errors while we modify the code. It is also very easy to forget to make a required change.

With the previous code in `trapezoidal.py`, we can compute the new integral  $\int_{-1}^{1.1} e^{-x^2} dx$  *without touching the mathematical algorithm*. In an interactive session (or in a program) we can just do

```
>>> from trapezoidal import trapezoidal
>>> from math import exp
>>> trapezoidal(lambda x: exp(-x**2), -1, 1.1, 400)
1.5268823686123285
```

When you now look back at the two solutions, the flat special-purpose program and the function-based program with the general-purpose function `trapezoidal`, you hopefully realize that *implementing a general mathematical algorithm in a general function* requires somewhat more abstract thinking, but the resulting code can be used over and over again. Essentially, if you apply the flat special-purpose style, you have to retest the implementation of the algorithm after every change of the program.

---

<sup>2</sup> You cannot integrate  $e^{-x^2}$  by hand, but this particular integral is appearing so often in so many contexts that the integral is a special function, called the [Error function](http://en.wikipedia.org/wiki/Error_function) ([http://en.wikipedia.org/wiki/Error\\_function](http://en.wikipedia.org/wiki/Error_function)) and written `erf(x)`. In a code, you can call `erf(x)`. The `erf` function is found in the `math` module.

The present integral problems result in short code. In more challenging engineering problems the code quickly grows to hundreds and thousands of lines. Without abstractions in terms of general algorithms in general reusable functions, the complexity of the program grows so fast that it will be extremely difficult to make sure that the program works properly.

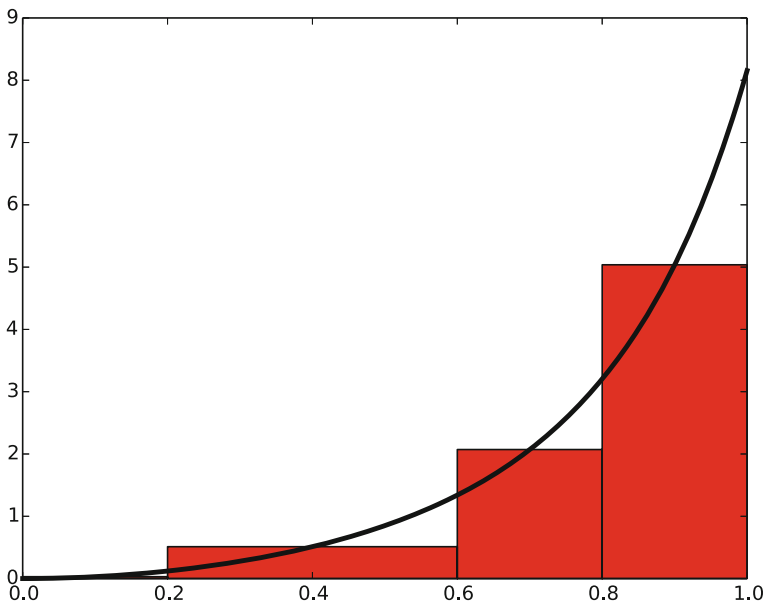
Another advantage of packaging mathematical algorithms in functions is that a function can be reused by anyone to solve a problem by just calling the function with a proper set of arguments. Understanding the function's inner details is not necessary to compute a new integral. Similarly, you can find libraries of functions on the Internet and use these functions to solve your problems without specific knowledge of every mathematical detail in the functions.

This desirable feature has its downside, of course: the user of a function may misuse it, and the function may contain programming errors and lead to wrong answers. Testing the output of downloaded functions is therefore extremely important before relying on the results.

---

### 3.3 The Composite Midpoint Method

**The idea** Rather than approximating the area under a curve by trapezoids, we can use plain rectangles (Fig. 3.3). It may sound less accurate to use horizontal lines and not skew lines following the function to be integrated, but an integration method based on rectangles (the *midpoint method*) is in fact slightly more accurate than the one based on trapezoids!



**Fig. 3.3** Computing approximately the integral of a function as the sum of the areas of the rectangles

In the midpoint method, we construct a rectangle for every sub-interval where the height equals  $f$  at the midpoint of the sub-interval. Let us do this for four rectangles, using the same sub-intervals as we had for hand calculations with the trapezoidal method:  $[0, 0.2)$ ,  $[0.2, 0.6)$ ,  $[0.6, 0.8)$ , and  $[0.8, 1.0]$ . We get

$$\begin{aligned} \int_0^1 f(t)dt &\approx h_1 f\left(\frac{0+0.2}{2}\right) + h_2 f\left(\frac{0.2+0.6}{2}\right) \\ &\quad + h_3 f\left(\frac{0.6+0.8}{2}\right) + h_4 f\left(\frac{0.8+1.0}{2}\right), \end{aligned} \quad (3.18)$$

where  $h_1, h_2, h_3$ , and  $h_4$  are the widths of the sub-intervals, used previously with the trapezoidal method and defined in (3.10)–(3.13).

With  $f(t) = 3t^2e^{t^3}$ , the approximation becomes 1.632. Compared with the true answer (1.718), this is about 5 % too small, but it is better than what we got with the trapezoidal method (10 %) with the same sub-intervals. More rectangles give a better approximation.

### 3.3.1 The General Formula

Let us derive a formula for the midpoint method based on  $n$  rectangles of equal width:

$$\begin{aligned} \int_a^b f(x)dx &= \int_{x_0}^{x_1} f(x)dx + \int_{x_1}^{x_2} f(x)dx + \dots + \int_{x_{n-1}}^{x_n} f(x)dx, \\ &\approx hf\left(\frac{x_0+x_1}{2}\right) + hf\left(\frac{x_1+x_2}{2}\right) + \dots + hf\left(\frac{x_{n-1}+x_n}{2}\right), \end{aligned} \quad (3.19)$$

$$\approx h\left(f\left(\frac{x_0+x_1}{2}\right) + f\left(\frac{x_1+x_2}{2}\right) + \dots + f\left(\frac{x_{n-1}+x_n}{2}\right)\right). \quad (3.20)$$

This sum may be written more compactly as

$$\int_a^b f(x)dx \approx h \sum_{i=0}^{n-1} f(x_i), \quad (3.21)$$

where  $x_i = \left(a + \frac{h}{2}\right) + ih$ .

### 3.3.2 Implementation

We follow the advice and lessons learned from the implementation of the trapezoidal method and make a function `midpoint(f, a, b, n)` (in a file `midpoint.py`) for implementing the general formula (3.21):

```
def midpoint(f, a, b, n):
    h = float(b-a)/n
    result = 0
    for i in range(n):
        result += f((a + h/2.0) + i*h)
    result *= h
    return result
```

We can test the function as we explained for the similar trapezoidal method. The error in our particular problem  $\int_0^1 3t^2 e^{t^3} dt$  with four intervals is now about 0.1 in contrast to 0.2 for the trapezoidal rule. This is in fact not accidental: one can show mathematically that the error of the midpoint method is a bit smaller than for the trapezoidal method. The differences are seldom of any practical importance, and on a laptop we can easily use  $n = 10^6$  and get the answer with an error of about  $10^{-12}$  in a couple of seconds.

### 3.3.3 Comparing the Trapezoidal and the Midpoint Methods

The next example shows how easy we can combine the trapezoidal and midpoint functions to make a comparison of the two methods in the file `compare_integration_methods.py`:

```
from trapezoidal import trapezoidal
from midpoint import midpoint
from math import exp

g = lambda y: exp(-y**2)
a = 0
b = 2
print '      n      midpoint      trapezoidal'
for i in range(1, 21):
    n = 2**i
    m = midpoint(g, a, b, n)
    t = trapezoidal(g, a, b, n)
    print '%7d %.16f %.16f' % (n, m, t)
```

Note the efforts put into nice formatting – the output becomes

n	midpoint	trapezoidal
2	0.8842000076332692	0.8770372606158094
4	0.8827889485397279	0.8806186341245393
8	0.8822686991994210	0.8817037913321336
16	0.8821288703366458	0.8819862452657772

```

32 0.8820933014203766 0.8820575578012112
64 0.8820843709743319 0.8820754296107942
128 0.8820821359746071 0.8820799002925637
256 0.8820815770754198 0.8820810181335849
512 0.8820814373412922 0.8820812976045025
1024 0.8820814024071774 0.8820813674728968
2048 0.8820813936736116 0.8820813849400392
4096 0.8820813914902204 0.8820813893068272
8192 0.8820813909443684 0.8820813903985197
16384 0.8820813908079066 0.8820813906714446
32768 0.8820813907737911 0.8820813907396778
65536 0.8820813907652575 0.8820813907567422
131072 0.8820813907631487 0.8820813907610036
262144 0.8820813907625702 0.8820813907620528
524288 0.8820813907624605 0.8820813907623183
1048576 0.8820813907624268 0.8820813907623890

```

A visual inspection of the numbers shows how fast the digits stabilize in both methods. It appears that 13 digits have stabilized in the last two rows.

#### Remark

The trapezoidal and midpoint methods are just two examples in a jungle of numerical integration rules. Other famous methods are Simpson's rule and Gauss quadrature. They all work in the same way:

$$\int_a^b f(x)dx \approx \sum_{i=0}^{n-1} w_i f(x_i).$$

That is, the integral is approximated by a sum of function evaluations, where each evaluation  $f(x_i)$  is given a weight  $w_i$ . The different methods differ in the way they construct the evaluation points  $x_i$  and the weights  $w_i$ . We have used equally spaced points  $x_i$ , but higher accuracy can be obtained by optimizing the location of  $x_i$ .

## 3.4 Testing

### 3.4.1 Problems with Brief Testing Procedures

Testing of the programs for numerical integration has so far employed two strategies. If we have an exact answer, we compute the error and see that increasing  $n$  decreases the error. When the exact answer is not available, we can (as in the comparison example in the previous section) look at the integral values and see that they stabilize as  $n$  grows. Unfortunately, these are very weak test procedures and not at all satisfactory for claiming that the software we have produced is correctly implemented.

To see this, we can introduce a bug in the application function that calls `trapezoidal`: instead of integrating  $3t^2e^{t^3}$ , we write “accidentally”  $3t^3e^{t^3}$ , but keep the same anti-derivative  $V(t)e^{t^3}$  for computing the error. With the bug and

$n = 4$ , the error is 0.1, but without the bug the error is 0.2! It is of course completely impossible to tell if 0.1 is the right value of the error. Fortunately, increasing  $n$  shows that the error stays about 0.3 in the program with the bug, so the test procedure with increasing  $n$  and checking that the error decreases points to a problem in the code.

Let us look at another bug, this time in the mathematical algorithm: instead of computing  $\frac{1}{2}(f(a) + f(b))$  as we should, we forget the second  $\frac{1}{2}$  and write  $0.5*f(a) + f(b)$ . The error for  $n = 4, 40, 400$  when computing  $\int_{1.1}^{1.9} 3t^2 e^{t^3} dt$  goes like 1400, 107, 10, respectively, which looks promising. The problem is that the right errors should be 369, 4.08, and 0.04. That is, the error should be reduced faster in the correct than in the buggy code. The problem, however, is that it is reduced in both codes, and we may stop further testing and believe everything is correctly implemented.

### Unit testing

A good habit is to test small pieces of a larger code individually, one at a time. This is known as *unit testing*. One identifies a (small) unit of the code, and then one makes a separate test for this unit. The unit test should be stand-alone in the sense that it can be run without the outcome of other tests. Typically, one algorithm in scientific programs is considered as a unit. The challenge with unit tests in numerical computing is to deal with numerical approximation errors. A fortunate side effect of unit testing is that the programmer is forced to use functions to modularize the code into smaller, logical pieces.

## 3.4.2 Proper Test Procedures

There are three serious ways to test the implementation of numerical methods via unit tests:

1. *Comparing with hand-computed results* in a problem with few arithmetic operations, i.e., small  $n$ .
2. *Solving a problem without numerical errors*. We know that the trapezoidal rule must be exact for linear functions. The error produced by the program must then be zero (to machine precision).
3. *Demonstrating correct convergence rates*. A strong test when we can compute exact errors, is to see how fast the error goes to zero as  $n$  grows. In the trapezoidal and midpoint rules it is known that the error depends on  $n$  as  $n^{-2}$  as  $n \rightarrow \infty$ .

**Hand-computed results** Let us use two trapezoids and compute the integral  $\int_0^1 v(t)$ ,  $v(t) = 3t^2 e^{t^3}$ :

$$\frac{1}{2}h(v(0) + v(0.5)) + \frac{1}{2}h(v(0.5) + v(1)) = 2.463642041244344,$$

when  $h = 0.5$  is the width of the two trapezoids. Running the program gives exactly the same result.

**Solving a problem without numerical errors** The best unit tests for numerical algorithms involve mathematical problems where we know the numerical result beforehand. Usually, numerical results contain unknown approximation errors, so knowing the numerical result implies that we have a problem where the approximation errors vanish. This feature may be present in very simple mathematical problems. For example, the trapezoidal method is exact for integration of linear functions  $f(x) = ax + b$ . We can therefore pick some linear function and construct a test function that checks equality between the exact analytical expression for the integral and the number computed by the implementation of the trapezoidal method.

A specific test case can be  $\int_{1.2}^{4.4} (6x - 4)dx$ . This integral involves an “arbitrary” interval  $[1.2, 4.4]$  and an “arbitrary” linear function  $f(x) = 6x - 4$ . By “arbitrary” we mean expressions where we avoid the special numbers 0 and 1 since these have special properties in arithmetic operations (e.g., forgetting to multiply is equivalent to multiplying by 1, and forgetting to add is equivalent to adding 0).

**Demonstrating correct convergence rates** Normally, unit tests must be based on problems where the numerical approximation errors in our implementation remain unknown. However, we often know or may assume a certain *asymptotic* behavior of the error. We can do some experimental runs with the test problem  $\int_0^1 3t^2 e^{t^3} dt$  where  $n$  is doubled in each run:  $n = 4, 8, 16$ . The corresponding errors are then 12 %, 3 % and 0.77 %, respectively. These numbers indicate that the error is roughly reduced by a factor of 4 when doubling  $n$ . Thus, the error converges to zero as  $n^{-2}$  and we say that the *convergence rate* is 2. In fact, this result can also be shown mathematically for the trapezoidal and midpoint method. Numerical integration methods usually have an error that converge to zero as  $n^{-p}$  for some  $p$  that depends on the method. With such a result, it does not matter if we do not know what the actual approximation error is: we know at what rate it is *reduced*, so running the implementation for two or more different  $n$  values will put us in a position to measure the expected rate and see if it is achieved.

The idea of a corresponding unit test is then to run the algorithm for some  $n$  values, compute the error (the absolute value of the difference between the exact analytical result and the one produced by the numerical method), and check that the error has *approximately* correct asymptotic behavior, i.e., that the error is proportional to  $n^{-2}$  in case of the trapezoidal and midpoint method.

Let us develop a more precise method for such unit tests based on convergence rates. We assume that the error  $E$  depends on  $n$  according to

$$E = Cn^r,$$

where  $C$  is an unknown constant and  $r$  is the convergence rate. Consider a set of experiments with various  $n$ :  $n_0, n_1, n_2, \dots, n_q$ . We compute the corresponding errors  $E_0, \dots, E_q$ . For two consecutive experiments, number  $i$  and  $i - 1$ , we have the error model

$$E_i = Cn_i^r, \tag{3.22}$$

$$E_{i-1} = Cn_{i-1}^r. \tag{3.23}$$



These are two equations for two unknowns  $C$  and  $r$ . We can easily eliminate  $C$  by dividing the equations by each other. Then solving for  $r$  gives

$$r_{i-1} = \frac{\ln(E_i/E_{i-1})}{\ln(n_i/n_{i-1})}. \quad (3.24)$$

We have introduced a subscript  $i - 1$  in  $r$  since the estimated value for  $r$  varies with  $i$ . Hopefully,  $r_{i-1}$  approaches the correct convergence rate as the number of intervals increases and  $i \rightarrow q$ .

### 3.4.3 Finite Precision of Floating-Point Numbers

The test procedures above lead to comparison of numbers for checking that calculations were correct. Such comparison is more complicated than what a newcomer might think. Suppose we have a calculation  $a + b$  and want to check that the result is what we expect. We start with  $1 + 2$ :

```
>>> a = 1; b = 2; expected = 3
>>> a + b == expected
True
```

Then we proceed with  $0.1 + 0.2$ :

```
>>> a = 0.1; b = 0.2; expected = 0.3
>>> a + b == expected
False
```

So why is  $0.1 + 0.2 \neq 0.3$ ? The reason is that real numbers cannot in general be exactly represented on a computer. They must instead be approximated by a [float-point number](https://en.wikipedia.org/wiki/Floating_point)<sup>3</sup> that can only store a finite amount of information, usually about 17 digits of a real number. Let us print 0.1, 0.2,  $0.1 + 0.2$ , and 0.3 with 17 decimals:

```
>>> print '%.17f\n%.17f\n%.17f\n%.17f' % (0.1, 0.2, 0.1 + 0.2, 0.3)
0.10000000000000001
0.20000000000000001
0.30000000000000004
0.29999999999999999
```

We see that all of the numbers have an inaccurate digit in the 17th decimal place. Because  $0.1 + 0.2$  evaluates to 0.30000000000000004 and 0.3 is represented as 0.29999999999999999, these two numbers are not equal. In general, real numbers in Python have (at most) 16 correct decimals.

When we compute with real numbers, these numbers are inaccurately represented on the computer, and arithmetic operations with inaccurate numbers lead to small rounding errors in the final results. Depending on the type of numerical algorithm, the rounding errors may or may not accumulate.

<sup>3</sup> [https://en.wikipedia.org/wiki/Floating\\_point](https://en.wikipedia.org/wiki/Floating_point)

If we cannot make tests like  $0.1 + 0.2 == 0.3$ , what should we then do? The answer is that we must accept some small inaccuracy and make a test with a *tolerance*. Here is the recipe:

```
>>> a = 0.1; b = 0.2; expected = 0.3
>>> computed = a + b
>>> diff = abs(expected - computed)
>>> tol = 1E-15
>>> diff < tol
True
```

Here we have set the tolerance for comparison to  $10^{-15}$ , but calculating  $0.3 - (0.1 + 0.2)$  shows that it equals  $-5.55\text{e-}17$ , so a lower tolerance could be used in this particular example. However, in other calculations we have little idea about how accurate the answer is (there could be accumulation of rounding errors in more complicated algorithms), so  $10^{-15}$  or  $10^{-14}$  are robust values. As we demonstrate below, these tolerances depend on the magnitude of the numbers in the calculations.

Doing an experiment with  $10^k + 0.3 - (10^k + 0.1 + 0.2)$  for  $k = 1, \dots, 10$  shows that the answer (which should be zero) is around  $10^{16-k}$ . This means that the tolerance must be larger if we compute with larger numbers. Setting a proper tolerance therefore requires some experiments to see what level of accuracy one can expect. A way out of this difficulty is to work with *relative* instead of *absolute* differences. In a relative difference we divide by one of the operands, e.g.,

$$a = 10^k + 0.3, \quad b = (10^k + 0.1 + 0.2), \quad c = \frac{a - b}{a}.$$

Computing this  $c$  for various  $k$  shows a value around  $10^{-16}$ . A safer procedure is thus to use *relative differences*.

### 3.4.4 Constructing Unit Tests and Writing Test Functions

Python has several frameworks for automatically running and checking a potentially very large number of tests for parts of your software by one command. This is an extremely useful feature during program development: whenever you have done some changes to one or more files, launch the test command and make sure nothing is broken because of your edits.

The test frameworks `nose` and `py.test` are particularly attractive as they are very easy to use. Tests are placed in special *test functions* that the frameworks can recognize and run for you. The requirements to a test function are simple:

- the name must start with `test_`
- the test function cannot have any arguments
- the tests inside test functions must be boolean expressions
- a boolean expression `b` must be tested with `assert b, msg`, where `msg` is an optional object (string or number) to be written out when `b` is false

Suppose we have written a function

```
def add(a, b):
    return a + b
```

A corresponding test function can then be

```
def test_add():
    expected = 1 + 1
    computed = add(1, 1)
    assert computed == expected, '1+1=%g' % computed
```

Test functions can be in any program file or in separate files, typically with names starting with `test`. You can also collect tests in subdirectories: running `py.test -s -v` will actually run all tests in all `test*.py` files in all subdirectories, while `nosetests -s -v` restricts the attention to subdirectories whose names start with `test` or end with `_test` or `_tests`.

As long as we add integers, the equality test in the `test_add` function is appropriate, but if we try to call `add(0.1, 0.2)` instead, we will face the rounding error problems explained in Sect. 3.4.3, and we must use a test with tolerance instead:

```
def test_add():
    expected = 0.3
    computed = add(0.1, 0.2)
    tol = 1E-14
    diff = abs(expected - computed)
    assert diff < tol, 'diff=%g' % diff
```

Below we shall write test functions for each of the three test procedures we suggested: comparison with hand calculations, checking problems that can be exactly solved, and checking convergence rates. We stick to testing the trapezoidal integration code and collect all test functions in one common file by the name `test_trapezoidal.py`.

**Hand-computed numerical results** Our previous hand calculations for two trapezoids can be checked against the trapezoidal function inside a test function (in a file `test_trapezoidal.py`):

```
from trapezoidal import trapezoidal

def test_trapezoidal_one_exact_result():
    """Compare one hand-computed result."""
    from math import exp
    v = lambda t: 3*(t**2)*exp(t**3)
    n = 2
    computed = trapezoidal(v, 0, 1, n)
    expected = 2.463642041244344
    error = abs(expected - computed)
    tol = 1E-14
    success = error < tol
    msg = 'error=%g > tol=%g' % (error, tol)
    assert success, msg
```

Note the importance of checking `err` against `exact` with a tolerance: rounding errors from the arithmetics inside `trapezoidal` will not make the result exactly like the hand-computed one. The size of the tolerance is here set to  $10^{-14}$ , which is a kind of all-round value for computations with numbers not deviating much from unity.

**Solving a problem without numerical errors** We know that the trapezoidal rule is exact for linear integrands. Choosing the integral  $\int_{1.2}^{4.4} (6x - 4)dx$  as test case, the corresponding test function for this unit test may look like

```
def test_trapezoidal_linear():
    """Check that linear functions are integrated exactly."""
    f = lambda x: 6*x - 4
    F = lambda x: 3*x**2 - 4*x # Anti-derivative
    a = 1.2; b = 4.4
    expected = F(b) - F(a)
    tol = 1E-14
    for n in 2, 20, 21:
        computed = trapezoidal(f, a, b, n)
        error = abs(expected - computed)
        success = error < tol
        msg = 'n=%d, err=%g' % (n, error)
        assert success, msg
```

**Demonstrating correct convergence rates** In the present example with integration, it is known that the approximation errors in the trapezoidal rule are proportional to  $n^{-2}$ ,  $n$  being the number of subintervals used in the composite rule.

Computing convergence rates requires somewhat more tedious programming than the previous tests, but can be applied to more general integrands. The algorithm typically goes like

- for  $i = 0, 1, 2, \dots, q$ 
  - $n_i = 2^{i+1}$
  - Compute integral with  $n_i$  intervals
  - Compute the error  $E_i$
  - Estimate  $r_i$  from (3.24) if  $i > 0$

The corresponding code may look like

```
def convergence_rates(f, F, a, b, num_experiments=14):
    from math import log
    from numpy import zeros
    expected = F(b) - F(a)
    n = zeros(num_experiments, dtype=int)
    E = zeros(num_experiments)
    r = zeros(num_experiments-1)
    for i in range(num_experiments):
        n[i] = 2**(i+1)
        computed = trapezoidal(f, a, b, n[i])
        E[i] = abs(expected - computed)
        if i > 0:
            r_im1 = log(E[i]/E[i-1])/log(float(n[i])/n[i-1])
            r[i-1] = float('%.2f' % r_im1) # Truncate to two decimals
    return r
```

Making a test function is a matter of choosing  $f$ ,  $F$ ,  $a$ , and  $b$ , and then checking the value of  $r_i$  for the largest  $i$ :

```
def test_trapezoidal_conv_rate():
    """Check empirical convergence rates against the expected -2."""
    from math import exp
    v = lambda t: 3*(t**2)*exp(t**3)
    V = lambda t: exp(t**3)
    a = 1.1; b = 1.9
    r = convergence_rates(v, V, a, b, 14)
    print r
    tol = 0.01
    msg = str(r[-4:]) # show last 4 estimated rates
    assert (abs(r[-1]) - 2) < tol, msg
```

Running the test shows that all  $r_i$ , except the first one, equal the target limit 2 within two decimals. This observation suggests a tolerance of  $10^{-2}$ .

#### Remark about version control of files

Having a suite of test functions for automatically checking that your software works is considered as a fundamental requirement for reliable computing. Equally important is a system that can keep track of different versions of the files and the tests, known as a *version control system*. Today's most popular version control system is [Git](https://en.wikipedia.org/wiki/Git_(software))<sup>4</sup>, which the authors strongly recommend the reader to use for programming and writing reports. The combination of Git and cloud storage such as GitHub is a very common way of organizing scientific or engineering work. We have a [quick intro](http://hplgit.github.io/teamods/bitgit/Langtangen_bitgit-bootstrap.html)<sup>5</sup> to Git and GitHub that gets you up and running within minutes.

The typical workflow with Git goes as follows.

1. Before you start working with files, make sure you have the latest version of them by running `git pull`.
2. Edit files, remove or create files (new files must be registered by `git add`).
3. When a natural piece of work is done, *commit* your changes by the `git commit` command.
4. Implement your changes also in the cloud by doing `git push`.

A nice feature of Git is that people can edit the same file at the same time and very often Git will be able to automatically merge the changes (!). Therefore, version control is crucial when you work with others or when you do your work on different types of computers. Another key feature is that anyone can at any time view the history of a file, see who did what when, and roll back the entire file collection to a previous commit. This feature is, of course, fundamental for reliable work.

<sup>4</sup> [https://en.wikipedia.org/wiki/Git\\_\(software\)](https://en.wikipedia.org/wiki/Git_(software))

<sup>5</sup> [http://hplgit.github.io/teamods/bitgit/Langtangen\\_bitgit-bootstrap.html](http://hplgit.github.io/teamods/bitgit/Langtangen_bitgit-bootstrap.html)

### 3.5 Vectorization

The functions `midpoint` and `trapezoid` usually run fast in Python and compute an integral to a satisfactory precision within a fraction of a second. However, long loops in Python may run slowly in more complicated implementations. To increase the speed, the loops can be replaced by vectorized code. The integration functions constitute a simple and good example to illustrate how to vectorize loops.

We have already seen simple examples on vectorization in Sect. 1.4 when we could evaluate a mathematical function  $f(x)$  for a large number of  $x$  values stored in an array. Basically, we can write

```
def f(x):
    return exp(-x)*sin(x) + 5*x

from numpy import exp, sin, linspace
x = linspace(0, 4, 101) # coordinates from 100 intervals on [0, 4]
y = f(x)                # all points evaluated at once
```

The result `y` is the array that would be computed if we ran a `for` loop over the individual `x` values and called `f` for each value. Vectorization essentially eliminates this loop in Python (i.e., the looping over `x` and application of `f` to each `x` value are instead performed in a library with fast, compiled code).

**Vectorizing the midpoint rule** The aim of vectorizing the midpoint and trapezoidal functions is also to remove the explicit loop in Python. We start with vectorizing the midpoint function since trapezoid is not equally straightforward. The fundamental ideas of the vectorized algorithm are to

1. compute all the evaluation points in one array `x`
2. call `f(x)` to produce an array of corresponding function values
3. use the `sum` function to sum the `f(x)` values

The evaluation points in the midpoint method are  $x_i = a + (i + \frac{1}{2})h$ ,  $i = 0, \dots, n-1$ . That is,  $n$  uniformly distributed coordinates between  $a + h/2$  and  $b - h/2$ . Such coordinates can be calculated by `x = linspace(a+h/2, b-h/2, n)`. Given that the Python implementation `f` of the mathematical function  $f$  works with an array argument, which is very often the case in Python, `f(x)` will produce all the function values in an array. The array elements are then summed up by `sum`: `sum(f(x))`. This sum is to be multiplied by the rectangle width  $h$  to produce the integral value. The complete function is listed below.

```
from numpy import linspace, sum

def midpoint(f, a, b, n):
    h = float(b-a)/n
    x = linspace(a + h/2, b - h/2, n)
    return h*sum(f(x))
```

The code is found in the file `integration_methods_vec.py`.

Let us test the code interactively in a Python shell to compute  $\int_0^1 3t^2 dt$ . The file with the code above has the name `integration_methods_vec.py` and is a valid module from which we can import the vectorized function:

```
>>> from integration_methods_vec import midpoint
>>> from numpy import exp
>>> v = lambda t: 3*t**2*exp(t**3)
>>> midpoint(v, 0, 1, 10)
1.7014827690091872
```

Note the necessity to use `exp` from `numpy`: our `v` function will be called with `x` as an array, and the `exp` function must be capable of working with an array.

The vectorized code performs all loops very efficiently in compiled code, resulting in much faster execution. Moreover, many readers of the code will also say that the algorithm looks clearer than in the loop-based implementation.

**Vectorizing the trapezoidal rule** We can use the same approach to vectorize the trapezoid function. However, the trapezoidal rule performs a sum where the end points have different weight. If we do `sum(f(x))`, we get the end points `f(a)` and `f(b)` with weight unity instead of one half. A remedy is to subtract the error from `sum(f(x))`: `sum(f(x)) - 0.5*f(a) - 0.5*f(b)`. The vectorized version of the trapezoidal method then becomes

```
def trapezoidal(f, a, b, n):
    h = float(b-a)/n
    x = linspace(a, b, n+1)
    s = sum(f(x)) - 0.5*f(a) - 0.5*f(b)
    return h*s
```

---

## 3.6 Measuring Computational Speed

Now that we have created faster, vectorized versions of functions in the previous section, it is interesting to measure how much faster they are. The purpose of the present section is therefore to explain how we can record the CPU time consumed by a function so we can answer this question. There are many techniques for measuring the CPU time in Python, and here we shall just explain the simplest and most convenient one: the `%timeit` command in IPython. The following interactive session should illustrate a competition where the vectorized versions of the functions are supposed to win:

```
In [1]: from integration_methods_vec import midpoint as midpoint_vec

In [3]: from midpoint import midpoint

In [4]: from numpy import exp

In [5]: v = lambda t: 3*t**2*exp(t**3)
```

```

In [6]: %timeit midpoint_vec(v, 0, 1, 1000000)
1 loops, best of 3: 379 ms per loop

In [7]: %timeit midpoint(v, 0, 1, 1000000)
1 loops, best of 3: 8.17 s per loop

In [8]: 8.17/(379*0.001)      # efficiency factor
Out[8]: 21.556728232189972

```

We see that the vectorized version is about 20 times faster: 379 ms versus 8.17 s. The results for the trapezoidal method are very similar, and the factor of about 20 is independent of the number of intervals.

## 3.7 Double and Triple Integrals

### 3.7.1 The Midpoint Rule for a Double Integral

Given a double integral over a rectangular domain  $[a, b] \times [c, d]$ ,

$$\int_a^b \int_c^d f(x, y) dy dx,$$

how can we approximate this integral by numerical methods?

**Derivation via one-dimensional integrals** Since we know how to deal with integrals in one variable, a fruitful approach is to view the double integral as two integrals, each in one variable, which can be approximated numerically by previous one-dimensional formulas. To this end, we introduce a help function  $g(x)$  and write

$$\int_a^b \int_c^d f(x, y) dy dx = \int_a^b g(x) dx, \quad g(x) = \int_c^d f(x, y) dy.$$

Each of the integrals

$$\int_a^b g(x) dx, \quad g(x) = \int_c^d f(x, y) dy$$

can be discretized by any numerical integration rule for an integral in one variable. Let us use the midpoint method (3.21) and start with  $g(x) = \int_c^d f(x, y) dy$ . We introduce  $n_y$  intervals on  $[c, d]$  with length  $h_y$ . The midpoint rule for this integral then becomes

$$g(x) = \int_c^d f(x, y) dy \approx h_y \sum_{j=0}^{n_y-1} f(x, y_j), \quad y_j = c + \frac{1}{2}h_y + jh_y.$$



The expression looks somewhat different from (3.21), but that is because of the notation: since we integrate in the  $y$  direction and will have to work with both  $x$  and  $y$  as coordinates, we must use  $n_y$  for  $n$ ,  $h_y$  for  $h$ , and the counter  $i$  is more naturally called  $j$  when integrating in  $y$ . Integrals in the  $x$  direction will use  $h_x$  and  $n_x$  for  $h$  and  $n$ , and  $i$  as counter.

The double integral is  $\int_a^b g(x)dx$ , which can be approximated by the midpoint method:

$$\int_a^b g(x)dx \approx h_x \sum_{i=0}^{n_x-1} g(x_i), \quad x_i = a + \frac{1}{2}h_x + ih_x.$$

Putting the formulas together, we arrive at the *composite midpoint method for a double integral*:

$$\begin{aligned} \int_a^b \int_c^d f(x, y)dydx &\approx h_x \sum_{i=0}^{n_x-1} h_y \sum_{j=0}^{n_y-1} f(x_i, y_j) \\ &= h_x h_y \sum_{i=0}^{n_x-1} \sum_{j=0}^{n_y-1} f\left(a + \frac{h_x}{2} + ih_x, c + \frac{h_y}{2} + jh_y\right). \end{aligned} \quad (3.25)$$

**Direct derivation** The formula (3.25) can also be derived directly in the two-dimensional case by applying the idea of the midpoint method. We divide the rectangle  $[a, b] \times [c, d]$  into  $n_x \times n_y$  equal-sized cells. The idea of the midpoint method is to approximate  $f$  by a constant over each cell, and evaluate the constant at the midpoint. Cell  $(i, j)$  occupies the area

$$[a + ih_x, a + (i + 1)h_x] \times [c + jh_y, c + (j + 1)h_y],$$

and the midpoint is  $(x_i, y_j)$  with

$$x_i = a + ih_x + \frac{1}{2}h_x, \quad y_j = c + jh_y + \frac{1}{2}h_y.$$

The integral over the cell is therefore  $h_x h_y f(x_i, y_j)$ , and the total double integral is the sum over all cells, which is nothing but formula (3.25).

**Programming a double sum** The formula (3.25) involves a double sum, which is normally implemented as a double for loop. A Python function implementing (3.25) may look like

```
def midpoint_double1(f, a, b, c, d, nx, ny):
    hx = (b - a)/float(nx)
    hy = (d - c)/float(ny)
    I = 0
    for i in range(nx):
        for j in range(ny):
            xi = a + hx/2 + i*hx
            yj = c + hy/2 + j*hy
            I += hx*hy*f(xi, yj)
    return I
```

If this function is stored in a module file `midpoint_double.py`, we can compute some integral, e.g.,  $\int_2^3 \int_0^2 (2x + y) dx dy = 9$  in an interactive shell and demonstrate that the function computes the right number:

```
>>> from midpoint_double import midpoint_double1
>>> def f(x, y):
...     return 2*x + y
...
>>> midpoint_double1(f, 0, 2, 2, 3, 5, 5)
9.0
```

**Reusing code for one-dimensional integrals** It is very natural to write a two-dimensional midpoint method as we did in function `midpoint_double1` when we have the formula (3.25). However, we could alternatively ask, much as we did in the mathematics, can we reuse a well-tested implementation for one-dimensional integrals to compute double integrals? That is, can we use function `midpoint`

```
def midpoint(f, a, b, n):
    h = float(b-a)/n
    result = 0
    for i in range(n):
        result += f((a + h/2.0) + i*h)
    result *= h
    return result
```

from Sect. 3.3.2 “twice”? The answer is yes, if we think as we did in the mathematics: compute the double integral as a midpoint rule for integrating  $g(x)$  and define  $g(x_i)$  in terms of a midpoint rule over  $f$  in the  $y$  coordinate. The corresponding function has very short code:

```
def midpoint_double2(f, a, b, c, d, nx, ny):
    def g(x):
        return midpoint(lambda y: f(x, y), c, d, ny)

    return midpoint(g, a, b, nx)
```

The important advantage of this implementation is that we reuse a well-tested function for the standard one-dimensional midpoint rule and that we apply the one-dimensional rule exactly as in the mathematics.

**Verification via test functions** How can we test that our functions for the double integral work? The best unit test is to find a problem where the numerical approximation error vanishes because then we know exactly what the numerical answer should be. The midpoint rule is exact for linear functions, regardless of how many subinterval we use. Also, any linear two-dimensional function  $f(x, y) = px + qy + r$  will be integrated exactly by the two-dimensional midpoint rule. We may pick  $f(x, y) = 2x + y$  and create a proper *test function* that can automatically verify our two alternative implementations of the two-dimensional midpoint rule. To compute the integral of  $f(x, y)$  we take advantage of SymPy to eliminate the possibility of errors in hand calculations. The test function becomes

```
def test_midpoint_double():
    """Test that a linear function is integrated exactly."""
    def f(x, y):
        return 2*x + y

    a = 0; b = 2; c = 2; d = 3
    import sympy
    x, y = sympy.symbols('x y')
    I_expected = sympy.integrate(f(x, y), (x, a, b), (y, c, d))
    # Test three cases: nx < ny, nx = ny, nx > ny
    for nx, ny in (3, 5), (4, 4), (5, 3):
        I_computed1 = midpoint_double1(f, a, b, c, d, nx, ny)
        I_computed2 = midpoint_double2(f, a, b, c, d, nx, ny)
        tol = 1E-14
        #print I_expected, I_computed1, I_computed2
        assert abs(I_computed1 - I_expected) < tol
        assert abs(I_computed2 - I_expected) < tol
```

#### Let test functions speak up?

If we call the above `test_midpoint_double` function and nothing happens, our implementations are correct. However, it is somewhat annoying to have a function that is completely silent when it works – are we sure all things are properly computed? During development it is therefore highly recommended to insert a print statement such that we can monitor the calculations and be convinced that the test function does what we want. Since a test function should not have any print statement, we simply comment it out as we have done in the function listed above.

The trapezoidal method can be used as alternative for the midpoint method. The derivation of a formula for the double integral and the implementations follow exactly the same ideas as we explained with the midpoint method, but there are more terms to write in the formulas. Exercise 3.13 asks you to carry out the details. That exercise is a very good test on your understanding of the mathematical and programming ideas in the present section.

### 3.7.2 The Midpoint Rule for a Triple Integral

**Theory** Once a method that works for a one-dimensional problem is generalized to two dimensions, it is usually quite straightforward to extend the method to three dimensions. This will now be demonstrated for integrals. We have the triple integral

$$\int_a^b \int_c^d \int_e^f g(x, y, z) dz dy dx$$

and want to approximate the integral by a midpoint rule. Following the ideas for the double integral, we split this integral into one-dimensional integrals:

$$\begin{aligned} p(x, y) &= \int_e^f g(x, y, z) dz \\ q(x) &= \int_c^d p(x, y) dy \\ \int_a^b \int_c^d \int_e^f g(x, y, z) dz dy dx &= \int_a^b q(x) dx \end{aligned}$$

For each of these one-dimensional integrals we apply the midpoint rule:

$$\begin{aligned} p(x, y) &= \int_e^f g(x, y, z) dz \approx \sum_{k=0}^{n_z-1} g(x, y, z_k), \\ q(x) &= \int_c^d p(x, y) dy \approx \sum_{j=0}^{n_y-1} p(x, y_j), \\ \int_a^b \int_c^d \int_e^f g(x, y, z) dz dy dx &= \int_a^b q(x) dx \approx \sum_{i=0}^{n_x-1} q(x_i), \end{aligned}$$

where

$$z_k = e + \frac{1}{2}h_z + kh_z, \quad y_j = c + \frac{1}{2}h_y + jh_y, \quad x_i = a + \frac{1}{2}h_x + ih_x.$$

Starting with the formula for  $\int_a^b \int_c^d \int_e^f g(x, y, z) dz dy dx$  and inserting the two previous formulas gives

$$\begin{aligned} &\int_a^b \int_c^d \int_e^f g(x, y, z) dz dy dx \\ &\approx h_x h_y h_z \sum_{i=0}^{n_x-1} \sum_{j=0}^{n_y-1} \sum_{k=0}^{n_z-1} g \left( a + \frac{1}{2}h_x + ih_x, c + \frac{1}{2}h_y + jh_y, e + \frac{1}{2}h_z + kh_z \right). \end{aligned} \tag{3.26}$$

Note that we may apply the ideas under *Direct derivation* at the end of Sect. 3.7.1 to arrive at (3.26) directly: divide the domain into  $n_x \times n_y \times n_z$  cells of volumes  $h_x h_y h_z$ ; approximate  $g$  by a constant, evaluated at the midpoint  $(x_i, y_j, z_k)$ , in each cell; and sum the cell integrals  $h_x h_y h_z g(x_i, y_j, z_k)$ .

**Implementation** We follow the ideas for the implementations of the midpoint rule for a double integral. The corresponding functions are shown below and found in the file `midpoint_triple.py`.

```
def midpoint_triple1(g, a, b, c, d, e, f, nx, ny, nz):
    hx = (b - a)/float(nx)
    hy = (d - c)/float(ny)
    hz = (f - e)/float(nz)
    I = 0
    for i in range(nx):
        for j in range(ny):
            for k in range(nz):
                xi = a + hx/2 + i*hx
                yj = c + hy/2 + j*hy
                zk = e + hz/2 + k*hz
                I += hx*hy*hz*g(xi, yj, zk)
    return I

def midpoint(f, a, b, n):
    h = float(b-a)/n
    result = 0
    for i in range(n):
        result += f((a + h/2.0) + i*h)
    result *= h
    return result

def midpoint_triple2(g, a, b, c, d, e, f, nx, ny, nz):
    def p(x, y):
        return midpoint(lambda z: g(x, y, z), e, f, nz)

    def q(x):
        return midpoint(lambda y: p(x, y), c, d, ny)

    return midpoint(q, a, b, nx)

def test_midpoint_triple():
    """Test that a linear function is integrated exactly."""
    def g(x, y, z):
        return 2*x + y - 4*z

    a = 0; b = 2; c = 2; d = 3; e = -1; f = 2
    import sympy
    x, y, z = sympy.symbols('x y z')
    I_expected = sympy.integrate(
        g(x, y, z), (x, a, b), (y, c, d), (z, e, f))
    for nx, ny, nz in (3, 5, 2), (4, 4, 4), (5, 3, 6):
        I_computed1 = midpoint_triple1(
            g, a, b, c, d, e, f, nx, ny, nz)
        I_computed2 = midpoint_triple2(
            g, a, b, c, d, e, f, nx, ny, nz)
        tol = 1E-14
        print I_expected, I_computed1, I_computed2
        assert abs(I_computed1 - I_expected) < tol
        assert abs(I_computed2 - I_expected) < tol

if __name__ == '__main__':
    test_midpoint_triple()
```

### 3.7.3 Monte Carlo Integration for Complex-Shaped Domains

Repeated use of one-dimensional integration rules to handle double and triple integrals constitute a working strategy only if the integration domain is a rectangle or box. For any other shape of domain, completely different methods must be used. A common approach for two- and three-dimensional domains is to divide the domain into many small triangles or tetrahedra and use numerical integration methods for each triangle or tetrahedron. The overall algorithm and implementation is too complicated to be addressed in this book. Instead, we shall employ an alternative, very simple and general method, called Monte Carlo integration. It can be implemented in half a page of code, but requires orders of magnitude more function evaluations in double integrals compared to the midpoint rule.

However, Monte Carlo integration is much more computationally efficient than the midpoint rule when computing higher-dimensional integrals in more than three variables over hypercube domains. Our ideas for double and triple integrals can easily be generalized to handle an integral in  $m$  variables. A midpoint formula then involves  $m$  sums. With  $n$  cells in each coordinate direction, the formula requires  $n^m$  function evaluations. That is, the computational work explodes as an exponential function of the number of space dimensions. Monte Carlo integration, on the other hand, does not suffer from this explosion of computational work and is the preferred method for computing higher-dimensional integrals. So, it makes sense in a chapter on numerical integration to address Monte Carlo methods, both for handling complex domains and for handling integrals with many variables.

**The Monte Carlo integration algorithm** The idea of Monte Carlo integration of  $\int_a^b f(x)dx$  is to use the mean-value theorem from calculus, which states that the integral  $\int_a^b f(x)dx$  equals the length of the integration domain, here  $b-a$ , times the *average* value of  $f$ ,  $\bar{f}$ , in  $[a, b]$ . The average value can be computed by sampling  $f$  at a set of *random* points inside the domain and take the mean of the function values. In higher dimensions, an integral is estimated as the area/volume of the domain times the average value, and again one can evaluate the integrand at a set of random points in the domain and compute the mean value of those evaluations.

Let us introduce some quantities to help us make the specification of the integration algorithm more precise. Suppose we have some two-dimensional integral

$$\int_{\Omega} f(x, y) dx dy,$$

where  $\Omega$  is a two-dimensional domain defined via a help function  $g(x, y)$ :

$$\Omega = \{(x, y) \mid g(x, y) \geq 0\}$$

That is, points  $(x, y)$  for which  $g(x, y) \geq 0$  lie inside  $\Omega$ , and points for which  $g(x, y) < 0$  are outside  $\Omega$ . The boundary of the domain  $\partial\Omega$  is given by the implicit curve  $g(x, y) = 0$ . Such formulations of geometries have been very common during the last couple of decades, and one refers to  $g$  as a *level-set function* and the

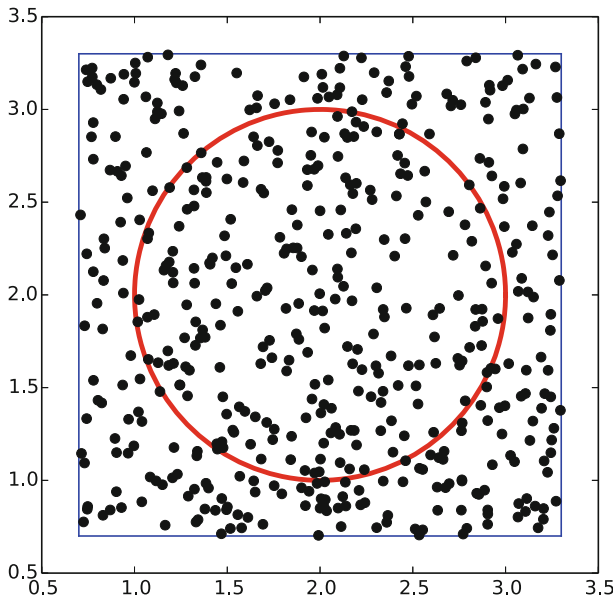
boundary  $g = 0$  as the zero-level contour of the level-set function. For simple geometries one can easily construct  $g$  by hand, while in more complicated industrial applications one must resort to mathematical models for constructing  $g$ .

Let  $A(\Omega)$  be the area of a domain  $\Omega$ . We can estimate the integral by this Monte Carlo integration method:

1. embed the geometry  $\Omega$  in a rectangular area  $R$
2. draw a large number of *random* points  $(x, y)$  in  $R$
3. count the fraction  $q$  of points that are inside  $\Omega$
4. approximate  $A(\Omega)/A(R)$  by  $q$ , i.e., set  $A(\Omega) = qA(R)$
5. evaluate the mean of  $f$ ,  $\bar{f}$ , at the points inside  $\Omega$
6. estimate the integral as  $A(\Omega)\bar{f}$

Note that  $A(R)$  is trivial to compute since  $R$  is a rectangle, while  $A(\Omega)$  is unknown. However, if we assume that the fraction of  $A(R)$  occupied by  $A(\Omega)$  is the same as the fraction of random points inside  $\Omega$ , we get a simple estimate for  $A(\Omega)$ .

To get an idea of the method, consider a circular domain  $\Omega$  embedded in a rectangle as shown below. A collection of random points is illustrated by black dots.



**Implementation** A Python function implementing  $\int_{\Omega} f(x, y) dx dy$  can be written like this:

```
import numpy as np

def MonteCarlo_double(f, g, x0, x1, y0, y1, n):
    """
    Monte Carlo integration of f over a domain g>=0, embedded
    in a rectangle [x0,x1]x[y0,y1]. n^2 is the number of
    random points.
    """
```

```

# Draw n**2 random points in the rectangle
x = np.random.uniform(x0, x1, n)
y = np.random.uniform(y0, y1, n)
# Compute sum of f values inside the integration domain
f_mean = 0
num_inside = 0 # number of x,y points inside domain (g>=0)
for i in range(len(x)):
    for j in range(len(y)):
        if g(x[i], y[j]) >= 0:
            num_inside += 1
            f_mean += f(x[i], y[j])
f_mean = f_mean/float(num_inside)
area = num_inside/float(n**2)*(x1 - x0)*(y1 - y0)
return area*f_mean

```

(See the file `MC_double.py`.)

**Verification** A simple test case is to check the area of a rectangle  $[0, 2] \times [3, 4.5]$  embedded in a rectangle  $[0, 3] \times [2, 5]$ . The right answer is 3, but Monte Carlo integration is, unfortunately, never exact so it is impossible to predict the output of the algorithm. All we know is that the estimated integral should approach 3 as the number of random points goes to infinity. Also, for a fixed number of points, we can run the algorithm several times and get different numbers that fluctuate around the exact value, since different sample points are used in different calls to the Monte Carlo integration algorithm.

The area of the rectangle can be computed by the integral  $\int_0^2 \int_3^{4.5} dy dx$ , so in this case we identify  $f(x, y) = 1$ , and the  $g$  function can be specified as (e.g.) 1 if  $(x, y)$  is inside  $[0, 2] \times [3, 4.5]$  and  $-1$  otherwise. Here is an example on how we can utilize the `MonteCarlo_double` function to compute the area for different number of samples:

```

>>> from MC_double import MonteCarlo_double
>>> def g(x, y):
...     return (1 if (0 <= x <= 2 and 3 <= y <= 4.5) else -1)
...
>>> MonteCarlo_double(lambda x, y: 1, g, 0, 3, 2, 5, 100)
2.9484
>>> MonteCarlo_double(lambda x, y: 1, g, 0, 3, 2, 5, 1000)
2.947032
>>> MonteCarlo_double(lambda x, y: 1, g, 0, 3, 2, 5, 1000)
3.0234600000000005
>>> MonteCarlo_double(lambda x, y: 1, g, 0, 3, 2, 5, 2000)
2.9984580000000003
>>> MonteCarlo_double(lambda x, y: 1, g, 0, 3, 2, 5, 2000)
3.1903469999999996
>>> MonteCarlo_double(lambda x, y: 1, g, 0, 3, 2, 5, 5000)
2.986515

```

We see that the values fluctuate around 3, a fact that supports a correct implementation, but in principle, bugs could be hidden behind the inaccurate answers.

It is mathematically known that the standard deviation of the Monte Carlo estimate of an integral converges as  $n^{-1/2}$ , where  $n$  is the number of samples. This



kind of convergence rate estimate could be used to verify the implementation, but this topic is beyond the scope of this book.

**Test function for function with random numbers** To make a test function, we need a unit test that has identical behavior each time we run the test. This seems difficult when random numbers are involved, because these numbers are different every time we run the algorithm, and each run hence produces a (slightly) different result. A standard way to test algorithms involving random numbers is to *fix the seed* of the random number generator. Then the sequence of numbers is the same every time we run the algorithm. Assuming that the `MonteCarlo_double` function works, we fix the seed, observe a certain result, and take this result as the correct result. Provided the test function always uses this seed, we should get exactly this result every time the `MonteCarlo_double` function is called. Our test function can then be written as shown below.

```
def test_MonteCarlo_double_rectangle_area():
    """Check the area of a rectangle."""
    def g(x, y):
        return (1 if (0 <= x <= 2 and 3 <= y <= 4.5) else -1)

    x0 = 0; x1 = 3; y0 = 2; y1 = 5 # embedded rectangle
    n = 1000
    np.random.seed(8) # must fix the seed!
    I_expected = 3.121092 # computed with this seed
    I_computed = MonteCarlo_double(
        lambda x, y: 1, g, x0, x1, y0, y1, n)
    assert abs(I_expected - I_computed) < 1E-14
```

(See the file `MC_double.py`.)

**Integral over a circle** The test above involves a trivial function  $f(x, y) = 1$ . We should also test a non-constant  $f$  function and a more complicated domain. Let  $\Omega$  be a circle at the origin with radius 2, and let  $f = \sqrt{x^2 + y^2}$ . This choice makes it possible to compute an exact result: in polar coordinates,  $\int_{\Omega} f(x, y) dx dy$  simplifies to  $2\pi \int_0^2 r^2 dr = 16\pi/3$ . We must be prepared for quite crude approximations that fluctuate around this exact result. As in the test case above, we experience better results with larger number of points. When we have such evidence for a working implementation, we can turn the test into a proper test function. Here is an example:

```
def test_MonteCarlo_double_circle_r():
    """Check the integral of r over a circle with radius 2."""
    def g(x, y):
        xc, yc = 0, 0 # center
        R = 2 # radius
        return R**2 - ((x-xc)**2 + (y-yc)**2)

    # Exact: integral of r*r*dr over circle with radius R becomes
    # 2*pi*1/3*R**3
    import sympy
    r = sympy.symbols('r')
    I_exact = sympy.integrate(2*sympy.pi*r*r, (r, 0, 2))
```

```

print 'Exact integral:', I_exact.evalf()
x0 = -2; x1 = 2; y0 = -2; y1 = 2
n = 1000
np.random.seed(6)
I_expected = 16.7970837117376384 # Computed with this seed
I_computed = MonteCarlo_double(
    lambda x, y: np.sqrt(x**2 + y**2),
    g, x0, x1, y0, y1, n)
print 'MC approximation %d samples): %.16f' % (n**2, I_computed)
assert abs(I_expected - I_computed) < 1E-15

```

(See the file `MC_double.py`.)

### 3.8 Exercises

#### Exercise 3.1: Hand calculations for the trapezoidal method

Compute by hand the area composed of two trapezoids (of equal width) that approximates the integral  $\int_1^3 2x^3 dx$ . Make a test function that calls the trapezoidal function in `trapezoidal.py` and compares the return value with the hand-calculated value.

Filename: `trapezoidal_test_func.py`.

#### Exercise 3.2: Hand calculations for the midpoint method

Compute by hand the area composed of two rectangles (of equal width) that approximates the integral  $\int_1^3 2x^3 dx$ . Make a test function that calls the midpoint function in `midpoint.py` and compares the return value with the hand-calculated value.

Filename: `midpoint_test_func.py`.

#### Exercise 3.3: Compute a simple integral

Apply the trapezoidal and midpoint functions to compute the integral  $\int_2^6 x(x-1)dx$  with 2 and 100 subintervals. Compute the error too.

Filename: `integrate_parabola.py`.

#### Exercise 3.4: Hand-calculations with sine integrals

We consider integrating the sine function:  $\int_0^b \sin(x)dx$ .

- Let  $b = \pi$  and use two intervals in the trapezoidal and midpoint method. Compute the integral by hand and illustrate how the two numerical methods approximates the integral. Compare with the exact value.
- Do a) when  $b = 2\pi$ .

Filename: `integrate_sine.pdf`.

#### Exercise 3.5: Make test functions for the midpoint method

Modify the file `test_trapezoidal.py` such that the three tests are applied to the function `midpoint` implementing the midpoint method for integration.

Filename: `test_midpoint.py`.

**Exercise 3.6: Explore rounding errors with large numbers**

The trapezoidal method integrates linear functions exactly, and this property was used in the test function `test_trapezoidal_linear` in the file `test_trapezoidal.py`. Change the function used in Sect. 3.4.2 to  $f(x) = 6 \cdot 10^8 x - 4 \cdot 10^6$  and rerun the test. What happens? How must you change the test to make it useful? How does the convergence rate test behave? Any need for adjustment?

Filename: `test_trapezoidal2.py`.

**Exercise 3.7: Write test functions for  $\int_0^4 \sqrt{x} dx$** 

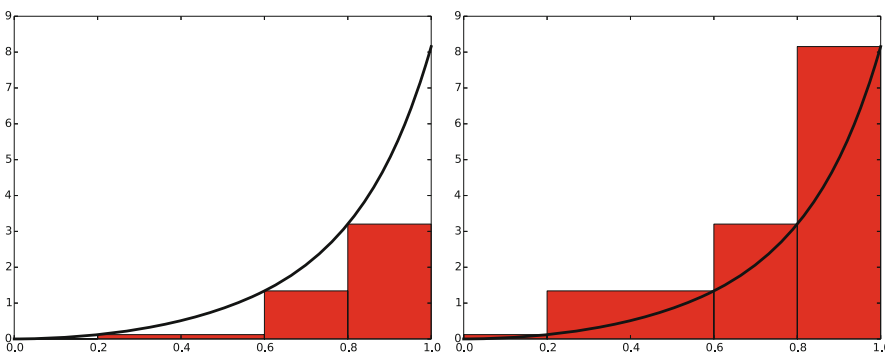
We want to test how the trapezoidal function works for the integral  $\int_0^4 \sqrt{x} dx$ . Two of the tests in `test_trapezoidal.py` are meaningful for this integral. Compute by hand the result of using 2 or 3 trapezoids and modify the `test_trapezoidal_one_exact_result` function accordingly. Then modify `test_trapezoidal_conv_rate` to handle the square root integral.

Filename: `test_trapezoidal3.py`.

*Remarks* The convergence rate test fails. Printing out `r` shows that the actual convergence rate for this integral is  $-1.5$  and not  $-2$ . The reason is that the [error in the trapezoidal method](#)<sup>6</sup> is  $-(b-a)^3 n^{-2} f''(\xi)$  for some (unknown)  $\xi \in [a, b]$ . With  $f(x) = \sqrt{x}$ ,  $f''(\xi) \rightarrow -\infty$  as  $\xi \rightarrow 0$ , pointing to a potential problem in the size of the error. Running a test with  $a > 0$ , say  $\int_{0.1}^4 \sqrt{x} dx$  shows that the convergence rate is indeed restored to  $-2$ .

**Exercise 3.8: Rectangle methods**

The midpoint method divides the interval of integration into equal-sized subintervals and approximates the integral in each subinterval by a rectangle whose height equals the function value at the midpoint of the subinterval. Instead, one might use either the left or right end of the subinterval as illustrated in Fig. 3.4. This defines a *rectangle method* of integration. The height of the rectangle can be based on the left or right end or the midpoint.



**Fig. 3.4** Illustration of the rectangle method with evaluating the rectangle height by either the left or right point

<sup>6</sup> [http://en.wikipedia.org/wiki/Trapezoidal\\_rule#Error\\_analysis](http://en.wikipedia.org/wiki/Trapezoidal_rule#Error_analysis)

- a) Write a function `rectangle(f, a, b, n, height='left')` for computing an integral  $\int_a^b f(x)dx$  by the rectangle method with height computed based on the value of `height`, which is either `left`, `right`, or `mid`.
- b) Write three test functions for the three unit test procedures described in Sect. 3.4.2. Make sure you test for `height` equal to `left`, `right`, and `mid`. You may call the midpoint function for checking the result when `height=mid`.

*Hint* Edit `test_trapezoidal.py`.

Filename: `rectangle_methods.py`.

### Exercise 3.9: Adaptive integration

Suppose we want to use the trapezoidal or midpoint method to compute an integral  $\int_a^b f(x)dx$  with an error less than a prescribed tolerance  $\epsilon$ . What is the appropriate size of  $n$ ?

To answer this question, we may enter an iterative procedure where we compare the results produced by  $n$  and  $2n$  intervals, and if the difference is smaller than  $\epsilon$ , the value corresponding to  $2n$  is returned. Otherwise, we halve  $n$  and repeat the procedure.

*Hint* It may be a good idea to organize your code so that the function `adaptive_integration` can be used easily in future programs you write.

- a) Write a function

```
adaptive_integration(f, a, b, eps, method=midpoint)
```

that implements the idea above (`eps` corresponds to the tolerance  $\epsilon$ , and `method` can be `midpoint` or `trapezoidal`).

- b) Test the method on  $\int_0^2 x^2 dx$  and  $\int_0^2 \sqrt{x} dx$  for  $\epsilon = 10^{-1}, 10^{-10}$  and write out the exact error.
- c) Make a plot of  $n$  versus  $\epsilon \in [10^{-1}, 10^{-10}]$  for  $\int_0^2 \sqrt{x} dx$ . Use logarithmic scale for  $\epsilon$ .

Filename: `adaptive_integration.py`.

*Remarks* The type of method explored in this exercise is called *adaptive*, because it tries to adapt the value of  $n$  to meet a given error criterion. The true error can very seldom be computed (since we do not know the exact answer to the computational problem), so one has to find other indicators of the error, such as the one here where the changes in the integral value, as the number of intervals is doubled, is taken to reflect the error.

### Exercise 3.10: Integrating $x$ raised to $x$

Consider the integral

$$I = \int_0^4 x^x dx.$$

The integrand  $x^x$  does not have an anti-derivative that can be expressed in terms of standard functions (visit <http://wolframalpha.com> and type `integral(x**x,x)` to convince yourself that our claim is right. Note that Wolfram alpha does give you an answer, but that answer is an approximation, it is *not* exact. This is because Wolfram alpha too uses numerical methods to arrive at the answer, just as you will in this exercise). Therefore, we are forced to compute the integral by numerical methods. Compute a result that is right to four digits.

*Hint* Use ideas from Exercise 3.9.

Filename: `integrate_x2x.py`.

### Exercise 3.11: Integrate products of sine functions

In this exercise we shall integrate

$$I_{j,k} = \int_{-\pi}^{\pi} \sin(jx) \sin(kx) dx,$$

where  $j$  and  $k$  are integers.

- Plot  $\sin(x) \sin(2x)$  and  $\sin(2x) \sin(3x)$  for  $x \in ]-\pi, \pi]$  in separate plots. Explain why you expect  $\int_{-\pi}^{\pi} \sin x \sin 2x dx = 0$  and  $\int_{-\pi}^{\pi} \sin 2x \sin 3x dx = 0$ .
- Use the trapezoidal rule to compute  $I_{j,k}$  for  $j = 1, \dots, 10$  and  $k = 1, \dots, 10$ .

Filename: `products_sines.py`.

### Exercise 3.12: Revisit fit of sines to a function

This is a continuation of Exercise 2.18. The task is to approximate a given function  $f(t)$  on  $[-\pi, \pi]$  by a sum of sines,

$$S_N(t) = \sum_{n=1}^N b_n \sin(nt). \quad (3.27)$$

We are now interested in computing the unknown coefficients  $b_n$  such that  $S_N(t)$  is in some sense the *best approximation* to  $f(t)$ . One common way of doing this is to first set up a general expression for the *approximation error*, measured by “summing up” the squared deviation of  $S_N$  from  $f$ :

$$E = \int_{-\pi}^{\pi} (S_N(t) - f(t))^2 dt.$$

We may view  $E$  as a function of  $b_1, \dots, b_N$ . Minimizing  $E$  with respect to  $b_1, \dots, b_N$  will give us a *best approximation*, in the sense that we adjust  $b_1, \dots, b_N$  such that  $S_N$  deviates as little as possible from  $f$ .

Minimization of a function of  $N$  variables,  $E(b_1, \dots, b_N)$  is mathematically performed by requiring all the partial derivatives to be zero:

$$\begin{aligned}\frac{\partial E}{\partial b_1} &= 0, \\ \frac{\partial E}{\partial b_2} &= 0, \\ &\vdots \\ \frac{\partial E}{\partial b_N} &= 0.\end{aligned}$$

- a) Compute the partial derivative  $\partial E / \partial b_1$  and generalize to the arbitrary case  $\partial E / \partial b_n$ ,  $1 \leq n \leq N$ .
- b) Show that

$$b_n = \frac{1}{\pi} \int_{-\pi}^{\pi} f(t) \sin(nt) dt.$$

- c) Write a function `integrate_coeffs(f, N, M)` that computes  $b_1, \dots, b_N$  by numerical integration, using  $M$  intervals in the trapezoidal rule.
- d) A remarkable property of the trapezoidal rule is that it is exact for integrals  $\int_{-\pi}^{\pi} \sin nt dt$  (when subintervals are of equal size). Use this property to create a function `test_integrate_coeff` to verify the implementation of `integrate_coeffs`.
- e) Implement the choice  $f(t) = \frac{1}{\pi}t$  as a Python function `f(t)` and call `integrate_coeffs(f, 3, 100)` to see what the optimal choice of  $b_1, b_2, b_3$  is.
- f) Make a function `plot_approx(f, N, M, filename)` where you plot `f(t)` together with the best approximation  $S_N$  as computed above, using  $M$  intervals for numerical integration. Save the plot to a file with name `filename`.
- g) Run `plot_approx(f, N, M, filename)` for  $f(t) = \frac{1}{\pi}t$  for  $N = 3, 6, 12, 24$ . Observe how the approximation improves.
- h) Run `plot_approx` for  $f(t) = e^{-(t-\pi)}$  and  $N = 100$ . Observe a fundamental problem: regardless of  $N$ ,  $S_N(-\pi) = 0$ , not  $e^{2\pi} \approx 535$ . (There are ways to fix this issue.)

Filename: `autofit_sines.py`.

### Exercise 3.13: Derive the trapezoidal rule for a double integral

Use ideas in Sect. 3.7.1 to derive a formula for computing a double integral  $\int_a^b \int_c^d f(x, y) dy dx$  by the trapezoidal rule. Implement and test this rule.

Filename: `trapezoidal_double.py`.

### Exercise 3.14: Compute the area of a triangle by Monte Carlo integration

Use the Monte Carlo method from Sect. 3.7.3 to compute the area of a triangle with vertices at  $(-1, 0)$ ,  $(1, 0)$ , and  $(3, 0)$ .

Filename: `MC_triangle.py`.

**Open Access** This chapter is distributed under the terms of the Creative Commons Attribution-NonCommercial 4.0 International License (<http://creativecommons.org/licenses/by-nc/4.0/>), which permits any noncommercial use, duplication, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, a link is provided to the Creative Commons license and any changes made are indicated.

The images or other third party material in this chapter are included in the work's Creative Commons license, unless indicated otherwise in the credit line; if such material is not included in the work's Creative Commons license and the respective action is not permitted by statutory regulation, users will need to obtain permission from the license holder to duplicate, adapt or reproduce the material.

Programming for Computations - Python  
A Gentle Introduction to Numerical Simulations with  
Python

Linge, S.; Langtangen, H.P.

2016, XVI, 232 p. 45 illus., Hardcover

ISBN: 978-3-319-32427-2