Ch.8: Random numbers and simple games

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Use of random numbers in programs





Random numbers are used to simulate uncertain events Deterministic problems.

- Some problems in science and technology are described by "exact" mathematics, leading to "precise" results
- Example: throwing a ball up in the air $(y(t) = v_0 t \frac{1}{2}gt^2)$

Stochastic problems.

- Some problems appear physically uncertain
- Examples: rolling a die, molecular motion, games
- Use random numbers to mimic the uncertainty of the experiment.

Drawing random numbers

Python has a random module for drawing random numbers. random() draws random numbers in [0,1):

```
>>> import random
>>> random.random()
0.81550546885338104
>>> random.random()
0.44913326809029852
>>> random.random()
0.88320653116367454
```

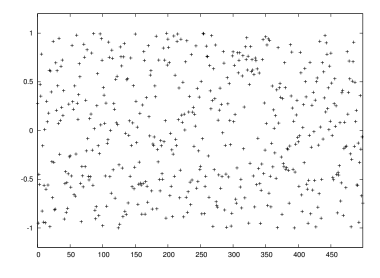
Notice. The sequence of random numbers is produced by a deterministic algorithm - the numbers just appear random.

Distribution of random numbers

- random.random() generates random numbers that are *uniformly distributed* in the interval [0,1)
- \bullet random.uniform(a, b) generates random numbers uniformly distributed in [a,b)
- "Uniformly distributed" means that if we generate a large set of numbers, no part of [a,b) gets more numbers than others

Distribution of random numbers visualized

```
N = 500 # no of samples
x = range(N)
y = [random.uniform(-1,1) for i in x]
from scitools.std import plot
plot(x, y, '+', axis=[0,N-1,-1.2,1.2])
```



Vectorized drawing of random numbers

- random.random() generates one number at a time
- numpy has a random module that efficiently generates a (large) number of random numbers at a time

```
from numpy import random
r = random.random()  # one no between 0 and 1
r = random.random(size=10000)  # array with 10000 numbers
r = random.uniform(-1, 10)  # one no between -1 and 10
r = random.uniform(-1, 10, size=10000)  # array
```

- Vectorized drawing is important for speeding up programs!
- Possible problem: two random modules, one Python "built-in" and one in numpy (np)
- Convention: use random (Python) and np.random

```
random.uniform(-1, 1)  # scalar number
import numpy as np
np.random.uniform(-1, 1, 100000)  # vectorized
```

Drawing integers

- Quite often we want to draw an integer from [a, b] and not a real number
- Python's random module and numpy.random have functions for drawing uniformly distributed integers:

```
import random
r = random.randint(a, b) # a, a+1, ..., b
import numpy as np
r = np.random.randint(a, b+1, N) # b+1 is not included
r = np.random.random_integers(a, b, N) # b is included
```

Example: Rolling a die

Problem.

- Any no of eyes, 1-6, is equally probable when you roll a die
- What is the chance of getting a 6?

Solution by Monte Carlo simulation: Rolling a die is the same as drawing integers in [1, 6].

```
import random
N = 10000
eyes = [random.randint(1, 6) for i in range(N)]
M = 0 # counter for successes: how many times we get 6 eyes
for outcome in eyes:
    if outcome == 6:
        M += 1
print 'Got six %d times out of %d' % (M, N)
print 'Probability:', float(M)/N
Probability: M/N (exact: 1/6)
```

Example: Rolling a die; vectorized version

```
import sys, numpy as np
N = int(sys.argv[1])
eyes = np.random.randint(1, 7, N)
success = eyes == 6  # True/False array
six = np.sum(success)  # treats True as 1, False as 0
print 'Got six %d times out of %d' % (six, N)
print 'Probability:', float(M)/N
```

Impoartant! Use sum from numpy and not Python's built-in sum function! (The latter is slow, often making a vectorized version slower than the scalar version.)

Debugging programs with random numbers requires fixing the seed of the random sequence

- Debugging programs with random numbers is difficult because the numbers produced vary each time we run the program
- For debugging it is important that a new run reproduces the sequence of random numbers in the last run
- This is possible by fixing the *seed* of the random module: random.seed(121) (int argument)

```
>>> import random
>>> random.seed(2)
>>> ['%.2f' % random.random() for i in range(7)]
['0.96', '0.95', '0.06', '0.08', '0.84', '0.74', '0.67']
>>> ['%.2f' % random.random() for i in range(7)]
['0.31', '0.61', '0.61', '0.58', '0.16', '0.43', '0.39']
```

```
>>> random.seed(2)  # repeat the random sequence
>>> ['%.2f' % random.random() for i in range(7)]
['0.96', '0.95', '0.06', '0.08', '0.84', '0.74', '0.67']
```

By default, the seed is based on the current time

Drawing random elements from a list

There are different methods for picking an element from a list at random, but the main method applies choice(list):

```
>>> awards = ['car', 'computer', 'ball', 'pen']
>>> import random
>>> random.choice(awards)
'car'

Alternatively, we can compute a random index:
>>> index = random.randint(0, len(awards)-1)
>>> awards[index]
'pen'

We can also shuffle the list randomly, and then pick any element:
>>> random.shuffle(awards)
>>> awards[0]
'computer'
```

Example: Drawing cards from a deck; make deck and draw

Make a deck of cards:

Draw a card at random:

```
deck = make_deck()
card = deck[0]
del deck[0]

card = deck.pop(0) # return and remove element with index 0
```

Example: Drawing cards from a deck; draw a hand of cards

Draw a hand of n cards:

```
def deal_hand(n, deck):
    hand = [deck[i] for i in range(n)]
    del deck[:n]
    return hand, deck
```

Note:

- deck is returned since the function changes the list
- deck is changed in-place so the change affects the deck object in the calling code anyway, but returning changed arguments is a Python convention and good habit

Example: Drawing cards from a deck; deal

Deal hands for a set of players:

```
def deal(cards_per_hand, no_of_players):
    deck = make_deck()
    hands = []
    for i in range(no_of_players):
        hand, deck = deal_hand(cards_per_hand, deck)
        hands.append(hand)
    return hands

players = deal(5, 4)
import pprint; pprint.pprint(players)
```

Resulting output:

```
[['D4', 'CQ', 'H10', 'DK', 'CK'], ['D7', 'D6', 'SJ', 'S4', 'C5'], ['C3', 'DQ', 'S3', 'C9', 'DJ'], ['H6', 'H9', 'C6', 'D5', 'S6']]
```

Example: Drawing cards from a deck; analyze results (1)

Analyze the no of pairs or n-of-a-kind in a hand:

Example: Drawing cards from a deck; analyze results (2)

Analyze the no of combinations of the same suit:

```
def same_suit(hand):
    suits = [card[0] for card in hand]
    counter = {} # counter[suit] = how many cards of suit
    for suit in suits:
        # attention only to count > 1:
        count = suits.count(suit)
        if count > 1:
            counter[suit] = count
```

Example: Drawing cards from a deck; analyze results (3)

Analysis of how many cards we have of the same suit or the same rank, with some nicely formatted printout (see the book):

```
The hand D4, CQ, H10, DK, CK
has 1 pairs, 0 3-of-a-kind and
2+2 cards of the same suit.

The hand D7, D6, SJ, S4, C5
has 0 pairs, 0 3-of-a-kind and
2+2 cards of the same suit.

The hand C3, DQ, S3, C9, DJ
has 1 pairs, 0 3-of-a-kind and
2+2 cards of the same suit.

The hand H6, H9, C6, D5, S6
has 0 pairs, 1 3-of-a-kind and
2 cards of the same suit.
```

Class implementation of a deck; class Deck

Class version. We can wrap the previous functions in a class:

- Attribute: the deck
- Methods for shuffling, dealing, putting a card back

Code:

```
del self.deck[:n]

# alternative:
    # hand = [self.pop(0) for i in range(n)]
    return hand

def putback(self, card):
    """Put back a card under the rest."""
    self.deck.append(card)
```

Class implementation of a deck; alternative

Class implementation of a deck; why?

Warning: To print a Deck instance, Card and Hand must have __repr__ methods that return a "pretty print" string (see the book), because print on list object applies __repr__ to print each element.

Is the class version better than the function version? Yes! The function version has functions updating a global variable deck, as in

```
hand, deck = deal_hand(5, deck)
```

This is often considered bad programming. In the class version we avoid a global variable - the deck is stored and updated inside the class. Errors are less likely to sneak in in the class version.

Probabilities can be computed by Monte Carlo simulation

What is the probability that a certain event A happens? Simulate N events and count how many times M the event A happens. The probability of the event A is then M/N (as $N \to \infty$).

Example: You throw two dice, one black and one green. What is the probability that the number of eyes on the black is larger than that on the green?

A vectorized version can speed up the simulations

```
import sys
N = int(sys.argv[1])  # no of experiments
import numpy as np
r = np.random.random_integers(1, 6, (2, N))

black = r[0,:]  # eyes for all throws with black
green = r[1,:]  # eyes for all throws with green
success = black > green  # success[i] == True if black[i] > green[i]
M = np.sum(success)  # sum up all successes

p = float(M)/N
print 'probability:', p
```

Run 10+ times faster than scalar code

The exact probability can be calculated in this (simple) example

All possible combinations of two dice:

How many of the (black, green) pairs that have the property black > green?

```
success = [black > green for black, green in combinations]
M = sum(success)
print 'probability:', float(M)/len(combinations)
```

How accurate and fast is Monte Carlo simulation?

Programs:

- black_gt_green.py: scalar version
- black_gt_green_vec.py: vectorized version
- black_gt_green_exact.py: exact version

Gamification of this example

Suggested game: Suppose a games is constructed such that you have to pay 1 euro to throw the two dice. You win 2 euros if there are more eyes on the black than on the green die. Should you play this game?

Code:

```
import sys
N = int(sys.argv[1])
                                  # no of experiments
import random
start_capital = 10
money = start_capital
for i in range(N):
    money -= 1
                                   # pay for the game
    black = random.randint(1, 6) # throw black
    green = random.randint(1, 6) # throw brown
    if black > green:
                                  # success?
        money += 2
                                   # get award
net_profit_total = money - start_capital
net_profit_per_game = net_profit_total/float(N)
print 'Net profit per game in the long run:', net_profit_per_game
```

Should we play the game?

```
Terminaldd> python black_gt_green_game.py 1000000 Net profit per game in the long run: -0.167804
```

No!

Vectorization of the game for speeding up the code

Example: Drawing balls from a hat

We have 12 balls in a hat: four black, four red, and four blue

```
hat = []
for color in 'black', 'red', 'blue':
    for i in range(4):
        hat.append(color)
```

Choose two balls at random:

```
import random
index = random.randint(0, len(hat)-1)  # random index
ball1 = hat[index];  del hat[index]
index = random.randint(0, len(hat)-1)  # random index
ball2 = hat[index];  del hat[index]

# or:
random.shuffle(hat)  # random sequence of balls
ball1 = hat.pop(0)
ball2 = hat.pop(0)
```

What is the probability of getting two black balls or more?

```
def draw_ball(hat):
    index = random.randint(0, len(hat)-1)
    color = hat[index]; del hat[index]
    return color, hat # (return hat since it is modified)
# run experiments:
n = input('How many balls are to be drawn? ')
N = input('How many experiments?')
M = 0 # no of successes
for e in range(N):
   hat = new_hat()
    balls = []
                       # the n balls we draw
    for i in range(n):
        color, hat = draw_ball(hat)
        balls.append(color)
    if balls.count('black') >= 2: # two black balls or more?
        M += 1
print 'Probability:', float(M)/N
```

Examples on computing the probabilities

```
Terminal> python balls_in_hat.py
How many balls are to be drawn? 2
How many experiments? 10000
Probability: 0.0914

Terminal> python balls_in_hat.py
How many balls are to be drawn? 8
How many experiments? 10000
Probability: 0.9346

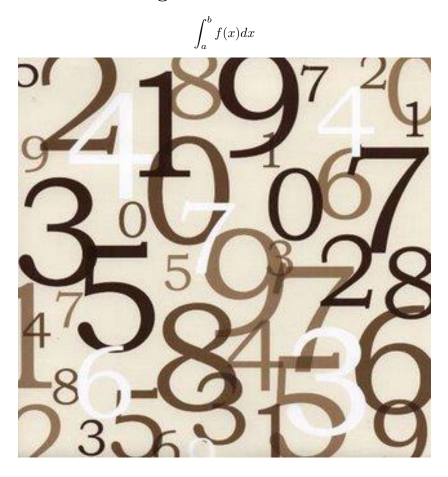
Terminal> python balls_in_hat.py
How many balls are to be drawn? 4
How many experiments? 10000
Probability: 0.4033
```

Guess a number game

Game: Let the computer pick a number at random. You guess at the number, and the computer tells if the number is too high or too low.

Program:

Monte Carlo integration



There is a strong link between an integral and the average of the integrand

Idea: Recall a famous theorem from calculus: Let f_m be the mean value of f(x) on [a,b]. Then

$$\int_{a}^{b} f(x)dx = f_{m}(b-a)$$

Idea: compute f_m by averaging N function values. To choose the N coordinates x_0, \ldots, x_{N-1} we use random numbers in [a, b]. Then

$$f_m = N^{-1} \sum_{j=0}^{N-1} f(x_j)$$

This is called Monte Carlo integration.

Implementation of Monte Carlo integration; scalar version

```
def MCint(f, a, b, n):
    s = 0
    for i in range(n):
        x = random.uniform(a, b)
        s += f(x)
    I = (float(b-a)/n)*s
    return I
```

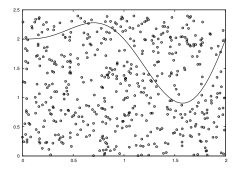
Implementation of Monte Carlo integration; vectorized version

```
def MCint_vec(f, a, b, n):
    x = np.random.uniform(a, b, n)
    s = np.sum(f(x))
    I = (float(b-a)/n)*s
    return I
```

Remark: Monte Carlo integration is slow for $\int f(x)dx$ (slower than the Trapezoidal rule, e.g.), but very efficient for integrating functions of many variables $\int f(x_1, x_2, \dots, x_n)dx_1dx_2 \cdots dx_n$

Dart-inspired Monte Carlo integration

- Choose a box $B = [x_L, x_H] \times [y_L, y_H]$ with some geometric object G inside, what is the area of G?
- Method: draw N points at random inside B, count how many, M, that fall within G, G's area is then $M/N \times \text{area}(B)$
- Special case: G is the geometry between y=f(x) and the x axis for $x\in [a,b]$, i.e., the area of G is $\int_a^b f(x)dx$, and our method gives $\int_a^b f(x)dx\approx \frac{M}{N}m(b-a)$ if B is the box $[a,b]\times [0,m]$



The code for the dart-inspired Monte Carlo integration Scalar code:

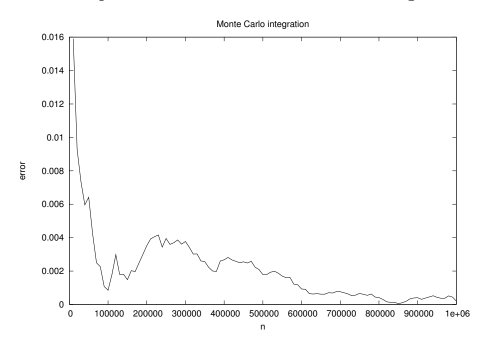
```
def MCint_area(f, a, b, n, fmax):
    below = 0  # counter for no of points below the curve
    for i in range(n):
        x = random.uniform(a, b)
        y = random.uniform(0, fmax)
        if y <= f(x):
            below += 1
        area = below/float(n)*(b-a)*fmax
    return area</pre>
```

Vectorized code:

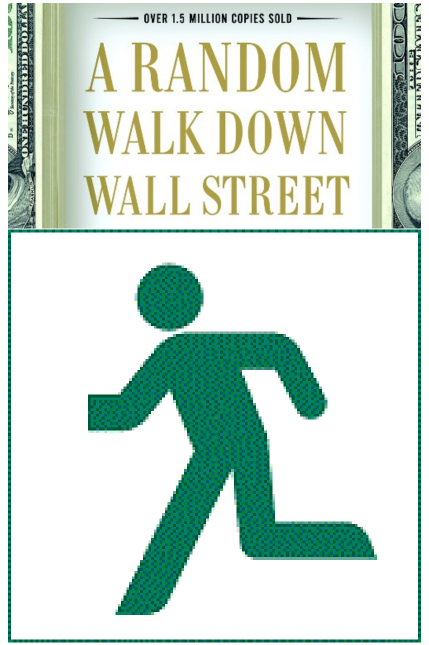
```
from numpy import *

def MCint_area_vec(f, a, b, n, fmax):
    x = np.random.uniform(a, b, n)
    y = np.random.uniform(0, fmax, n)
    below = y[y < f(x)].size
    area = below/float(n)*(b-a)*fmax
    return area</pre>
```

The development of the error in Monte Carlo integration



Random walk



Random walk in one space dimension

Basics of random walk in 1D:

- One particle moves to the left and right with equal probability
- n particles start at x = 0 at time t = 0 how do the particles get distributed over time?

Applications:

- molecular motion
- heat transport
- quantum mechanics
- polymer chains
- population genetics
- brain research
- hazard games
- pricing of financial instruments

Program for 1D random walk

```
from scitools.std import plot
import random
                          # no of particles
# no of steps
np = 4
ns = 100
positions = zeros(np) # all particles start at x=0 HEAD = 1; TAIL = 2 # constants
xmax = sqrt(ns); xmin = -xmax # extent of plot axis
for step in range(ns):
    for p in range(np):
         coin = random_.randint(1,2) # flip coin
         if coin == HE\overline{A}D:
             positions[p] += 1  # step to the right
         elif coin == TAIL:
             positions[p] -= 1
                                    # step to the left
    plot(positions, y, 'ko3', axis=[xmin, xmax, -0.2, 0.2])
    time.sleep(0.2)
                                     # pause between moves
```

Random walk as a difference equation

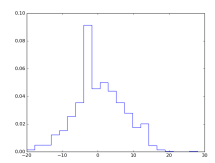
Let x_n be the position of one particle at time n. Updating rule:

$$x_n = x_{n-1} + s$$

where s = 1 or s = -1, both with probability 1/2.

Computing statistics of the random walk

Scientists are not interested in just looking at movies of random walks - they are interested in statistics (mean position, "width" of the cluster of particles, how particles are distributed)



Vectorized implementation of 1D random walk

First we draw all moves at all times:

```
moves = numpy.random.random_integers(1, 2, size=np*ns)
moves = 2*moves - 3 # -1, 1 instead of 1, 2
moves.shape = (ns, np)

Evolution through time:

positions = numpy.zeros(np)
for step in range(ns):
    positions += moves[step, :]

# can do some statistics:
    print numpy.mean(positions), numpy.std(positions)
```

Now to more exciting stuff: 2D random walk

Let each particle move north, south, west, or east - each with probability 1/4

```
def random_walk_2D(np, ns, plot_step):
    xpositions = numpy.zeros(np)
    ypositions = numpy.zeros(np)
    NORTH = 1;    SOUTH = 2;    WEST = 3;    EAST = 4
```

```
for step in range(ns):
    for i in range(len(xpositions)):
        direction = random.randint(1, 4)
        if direction == NORTH:
            ypositions[i] += 1
        elif direction == SOUTH:
            ypositions[i] -= 1
        elif direction == EAST:
            xpositions[i] += 1
        elif direction == WEST:
            xpositions[i] -= 1
    return xpositions, ypositions
```

Vectorized implementation of 2D random walk

```
def random_walk_2D(np, ns, plot_step):
    xpositions = zeros(np)
    ypositions = zeros(np)
    moves = numpy.random.random_integers(1, 4, size=ns*np)
    moves.shape = (ns, np)
    NORTH = 1;    SOUTH = 2;    WEST = 3;    EAST = 4

    for step in range(ns):
        this_move = moves[step,:]
        ypositions += where(this_move == NORTH, 1, 0)
        xpositions -= where(this_move == SOUTH, 1, 0)
        xpositions -= where(this_move == EAST, 1, 0)
        xpositions -= where(this_move == WEST, 1, 0)
    return xpositions, ypositions
```

Visualization of 2D random walk

- We plot every plot_step step
- One plot on the screen + one hardcopy for movie file
- Extent of axis: it can be shown that after n_s steps, the typical width of the cluster of particles (standard deviation) is of order $\sqrt{n_s}$, so we can set min/max axis extent as, e.g.,

Class implementation of 2D random walk

- Can classes be used to implement a random walk?
- Yes, it sounds natural with class Particle, holding the position of a particle as attributes and with a method move for moving the particle one step
- Class Particles holds a list of Particle instances and has a method move for moving all particles one step and a method moves for moving all particles through all steps
- Additional methods in class Particles can plot and compute statistics
- Downside: with class Particle the code is scalar a vectorized version must use arrays inside class Particles instead of a list of Particle instances
- The implementation is an exercise

i = random.randint(a, b)

Summary of drawing random numbers (scalar code)

```
Draw a uniformly distributed random number in [0, 1):
```

```
import random
r = random.random()

Draw a uniformly distributed random number in [a, b):
r = random.uniform(a, b)

Draw a uniformly distributed random integer in [a, b]:
```

Summary of drawing random numbers (vectorized code)

Draw n uniformly distributed random numbers in [0, 1):

```
import numpy as np
r = np.random.random(n)

Draw n uniformly distributed random numbers in [a,b):
r = np.random.uniform(a, b, n)

Draw n uniformly distributed random integers in [a,b]:
i = np.random.randint(a, b+1, n)
i = np.random.random_integers(a, b, n)
```

Summary of probability computations

- Probability: perform N experiments, count M successes, then success has probability M/N (N must be large)
- Monte Carlo simulation: let a program do N experiments and count M (simple method for probability problems)

Example: investment with random interest rate

Recall difference equation for the development of an investment x_0 with annual interest rate p:

$$x_n = x_{n-1} + \frac{p}{100}x_{n-1}$$
, given x_0

But:

- ullet In reality, p is uncertain in the future
- Let us model this uncertainty by letting p be random

Assume the interest is added every month:

$$x_n = x_{n-1} + \frac{p}{100 \cdot 12} x_{n-1}$$

where n counts months

The model for changing the interest rate

p changes from one month to the next by γ :

$$p_n = p_{n-1} + \gamma$$

where γ is random

- With probability $1/M, \gamma \neq 0$ (i.e., the annual interest rate changes on average every M months)
- If $\gamma \neq 0$, $\gamma = \pm m$, each with probability 1/2
- It does not make sense to have $p_n < 1$ or $p_n > 15$

The complete mathematical model

$$x_{n} = x_{n-1} + \frac{p_{n-1}}{12 \cdot 100} x_{n-1}, \quad i = 1, \dots, N$$

$$r_{1} = \text{random number in } 1, \dots, M$$

$$r_{2} = \text{random number in } 1, 2$$

$$\gamma = \begin{cases} m, & \text{if } r_{1} = 1 \text{ and } r_{2} = 1, \\ -m, & \text{if } r_{1} = 1 \text{ and } r_{2} = 2, \\ 0, & \text{if } r_{1} \neq 1 \end{cases}$$

$$p_{n} = p_{n-1} + \begin{cases} \gamma, & \text{if } p_{n} + \gamma \in [1, 15], \\ 0, & \text{otherwise} \end{cases}$$

A particular realization $x_n, p_n, n = 0, 1, ..., N$, is called a *path* (through time) or a realization. We are interested in the statistics of many paths.

Note: this is almost a random walk for the interest rate

Remark: The development of p is like a random walk, but the "particle" moves at each time level with probability 1/M (not 1 - always - as in a normal random walk).

Simulating the investment development; one path

```
def simulate_one_path(N, x0, p0, M, m):
    x = zeros(N+1)
    p = zeros(N+1)
    index_set = range(0, N+1)

    x[0] = x0
    p[0] = p0

for n in index_set[1:]:
        x[n] = x[n-1] + p[n-1]/(100.0*12)*x[n-1]

# update interest rate p:
    r = random.randint(1, M)
    if r == 1:
        # adjust gamma:
        r = random.randint(1, 2)
        gamma = m if r == 1 else -m
    else:
        gamma = 0
        pn = p[n-1] + gamma
        p[n] = pn if 1 <= pn <= 15 else p[n-1]
    return x, p</pre>
```

Simulating the investment development; N paths

Compute N paths (investment developments x_n) and their mean path (mean development)

```
def simulate_n_paths(n, N, L, p0, M, m):
    xm = zeros(N+1)
    pm = zeros(N+1)
    for i in range(n):
        x, p = simulate_one_path(N, L, p0, M, m)
        # accumulate paths:
        xm += x
        pm += p
# compute average:
    xm /= float(n)
    pm /= float(n)
    return xm, pm
```

Can also compute the standard deviation path ("width" of the N paths), see the book for details

Input and graphics

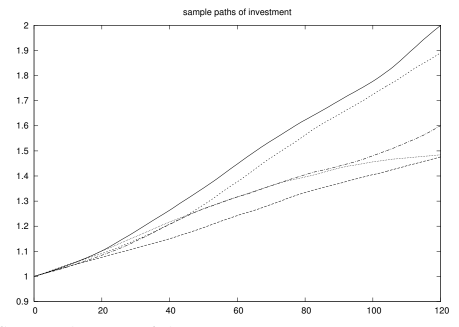
Here is a list of variables that constitute the input:

We may add some graphics in the program:

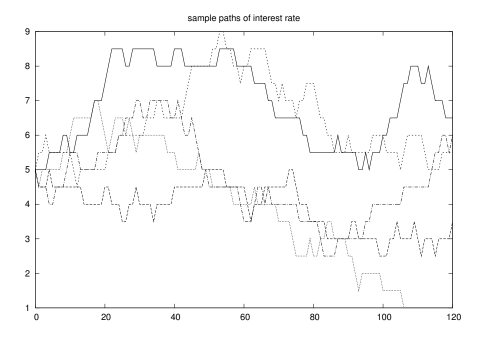
- plot some realizations of x_n and p_n
- plot the mean x_n with plus/minus one standard deviation
- plot the mean p_n with plus/minus one standard deviation

See the book for graphics details (good example on updating several different plots simultaneously in a simulation)

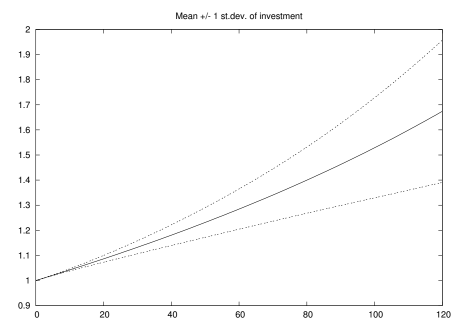
Some realizations of the investment



Some realizations of the interest rate



The mean and uncertainty of the investment over time



The mean and uncertainty of the interest rate over time

