

Scilab Textbook Companion for
Modern Physics
by R. A. Serway¹

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Book Description

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Scilab numbering policy used in this document and the relation to the above book.

Exa Example (Solved example)

Eqn Equation (Particular equation of the above book)

AP Appendix to Example(Scilab Code that is an Appednix to a particular Example of the above book)

For example, Exa 3.51 means solved example 3.51 of this book. Sec 2.3 means a scilab code whose theory is explained in Section 2.3 of the book.

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Chapter 1

Relativity I

Scilab code Exa 1.2 Period of the pendulum wrt different frames of references

```
1 // Scilab code Ex1.2: Pg.18 (2005)
2 clc; clear;
3 c = 3e+08;           // Velocity of light , m/s
4 v = 0.95*c;          // Velocity of observer , m/s
5 T_proper = 3;        // Proper time period of pendulum
                     in rest frame , s
6 gama = 1/(sqrt(1 - (v/c)^2));           // Multiplying
                     factor
7 // From time-dilation formula , we have
8 T = gama*T_proper;                  // Time period
                     of pendulum w.r.t to moving observer , s
9 printf("\nTime period of pendulum w.r.t to moving
                     observer = %3.1f s", T);
10
11 // Result
12 // Time period of pendulum w.r.t to moving observer
                     = 9.6 s
```

Scilab code Exa 1.3 Contraction of spaceship

```
1 // Scilab code Ex1.3: Pg 20 (2005)
2 clc; clear;
3 c = 3e+08; // Velocity of light , m
4 /s
5 L_p = 100; // Proper length of
6 spaceship , m
7 v = 0.99*c; // Velocity of spaceship
8 , m/s
9 // Using length contracction formula ,
10 L = L_p*sqrt(1 - (v/c)^2); // Observed length of spaceship , m
11 printf("Observed length of spaceship = %2d m" , L);
12
13 // Result
14 // Observed length of spaceship = 14 m
```

Scilab code Exa 1.4 Altitude of spaceship wrt different frames of references

```
1 // Scilab code Ex1.4: Pg 20 (2005)
2 clc; clear;
3 c = 3e+08; // Velocity of light , m
4 /s
5 L_p = 435; // Proper altitude of
6 spaceship , m
7 v = 0.970*c; // Velocity of
8 spaceship , m/s
9 // Using length contracction formula ,
10 L = L_p*sqrt(1 - (v/c)^2); // Observed altitude of spaceship , m
11 printf("Observed altitude of spaceship = %2d m" ,
ceil(L));
12
13 // Result
14 // Observed altitude of spaceship = 106 m
```

Scilab code Exa 1.5 Shape of spaceship seen from different frames of references

```
1 // Scilab code Ex1.5: Pg 20 (2005)
2 clc; clear;
3 c = 3e+08; // Velocity of light , m
4 s
5 L_p = 50; // Proper distance
6 between points x & y of spaceship , m
7 v = 0.950*c; // Velocity of
8 spaceship , m/s
9 // Using length contracction formula ,
10 L = L_p*sqrt(1 - (v/c)^2); // Observed distance between points x & y of
11 spaceship , m
12 printf("\nObserved distance between points x and y
13 of spaceship = %4.1f m", L);
14 printf("\nThe spaceship will get contracted in the
15 direction of motion");
16
17 // Result
18 // Observed distance between points x and y of
19 spaceship = 15.6 m
20 // The spaceship will get contracted in the
21 direction of motion
```

Scilab code Exa 1.6 Speed of recession of the galaxy Hydra

```
1 // Scilab code Ex1.6: Pg 25 (2005)
2 clc; clear;
3 // For simplification assume velocity of light equal
4 to unity
5 c = 1 // Velocity of light , m/s
```

```

5 lamda_obs = 474e-09;           // Wavelength measured
   by observer , m
6 lamda_source = 394e-09;        // Wavelength measured
   in the source 's rest frame , m
7 v = ((lamda_obs^2 - lamda_source^2)/(lamda_obs^2 +
   lamda_source^2))*c;           //
   Receding velocity of Hydra , m/s
8 printf("\nReceding velocity of Hydra = %5.3fc m/s" ,
   v);
9
10 // Result
11 // Receding velocity of Hydra = 0.183c m/s

```

Scilab code Exa 1.8 Relative velocity of spaceships

```

1 // Scilab code Ex1.8: Pg 30 (2005)
2 clc; clear;
3 // For simplification assume velocity of light equal
   to unity
4 c = 1;                      // Velocity of light , m/s
5 v = 0.750*c;                // Velocity of spaceship A
   relative to S frame , m/s
6 u_x = (-0.850)*c;           // Velocity of spaceship B
   relative to S frame , m/s
7 // Using Lorentz velocity transformation
8 U_x = (u_x - v)/(1 - u_x*v/c^2);      //
   Velocity of spaceship B with respect to spaceship
   A, m/s
9 printf("\nVelocity of spaceship B with respect to
   spaceship A = %6.4fc m/s" , U_x);
10
11 // Result
12 // Velocity of spaceship B with respect to spaceship
   A = -0.9771c m/s

```

Scilab code Exa 1.9 Velocity of ball wrt stationary observer

```
1 // Scilab code Ex1.9: Pg 30 (2005)
2 clc; clear;
3 // For simplification assume velocity of light equal
   to unity
4 c = 1;                      // Velocity of light , m/s
5 v = 0.800*c;                // Velocity of motorcycle w
   .r.t stationary observer , m/s
6 U_x = 0.700*c;              // Velocity of ball in the
   reference frame of motorcyclist , m/s
7 // Using inverse Lorentz velocity transformation
8 u_x = (U_x + v)/(1 + U_x*v/ c^2);          //
   Velocity of ball relative to stationary observer
   , m/s
9 printf("\nVelocity of ball relative to stationary
   observer = %6.4fc m/s", u_x);
10
11 // Result
12 // Velocity of ball relative to stationary observer
   = 0.9615c m/s
```

Scilab code Exa 1.10 Relative velocity of recession of two gang leaders

```
1 // Scilab code Ex1.10: Pg 30–31 (2005)
2 clc; clear;
3 // For simplification assume velocity of light equal
   to unity
4 c = 1;                      // Velocity of light , m/s
5 ux = 0.75*c;                // Velocity of pack
   leader alpha , m/s
6 gama = 1/(sqrt(1 - (ux/c)^2));
```

```

7 u_x = 0; // Velocity component of
           beta measured in S frame, m/s
8 U_x = (u_x - ux)/(1 - u_x*ux/c^2); // 
           Velocity component of beta along X-axis measured
           in S' frame, (Velocity Addition Rule), m/s
9 u_y = -0.90*c; // Velocity
           component of beta long Y-axis measured in S frame
10 U_y = u_y/(gama*(1 - u_x*ux/c^2)); // Velocity
           component of beta along Y-axis measured in S'
           frame, m/s
11 U = sqrt(U_x^2+U_y^2); // Relative velocity of
           recession of two gang leaders, m/s
12 printf("\nThe relative velocity of recession of two
           gang leaders = %4.2 fc", U);
13
14 // Result
15 // The relative velocity of recession of two gang
           leaders = 0.96c

```

Chapter 2

Relativity II

Scilab code Exa 2.1 Momentum of an electron

```
1 // Scilab code Ex2.1: Pg.44 (2005)
2 clc; clear;
3 c = 3e08; // Velocity of light , m/s
4 u = 0.750*c; // Velocity of electron , m/s
5 m = 9.11e-31; // Rest mass of electron , kg
6 p_r = m*u/(sqrt(1 - (u/c)^2)); // Relativistic
    momentum of electron , kgm/s
7 p = m*u; // Classical momentum of electron , kg-m/s
8 printf("\nThe relativistic momentum of electron = %4
        .2fe-22 kg-m/s", p_r*1e+22);
9 printf("\nThe classical momentum of electron = %4.2
        fe-22 kg-m/s", p*1e+22);
10 printf("\nThe relativistic result is 50 percent
        greater than the classical result.");
11
12 //Result
13 // The relativistic momentum of electron = 3.10e-22
    kg-m/s
14 // The classical momentum of electron = 2.05e-22 kg-
    m/s
15 // The relativistic result is 50 percent greater
```

than the classical result.

Scilab code Exa 2.3 Energy of a speedy electron

```
1 // Scilab code Ex2.3: Pg.47 (2005)
2 clc; clear;
3 c = 3e+08;      // Velocity of light , m/s
4 u = 0.85*c;    // Velocity of electron , m/s
5 E_0 = 0.511;   // Rest energy of electron , MeV
6 E = E_0/(sqrt(1-(u/c)^2)); // Total energy of
                                electron , MeV
7 K = E - E_0;   // Kinetic energy of electron , MeV
8 printf("\nThe total energy of electron = %5.3f MeV",
        E);
9 printf("\nThe kinetic energy of electron = %5.3f MeV
        ", K);
10
11 // Result
12 // The total energy of electron = 0.970 MeV
13 // The kinetic energy of electron = 0.459 MeV
```

Scilab code Exa 2.4 Energy of a speedy proton

```
1 // Scilab code Ex2.4: Pg.47 (2005)
2 clc; clear;
3 c = 3e+08;      // Velocity of light , m/s
4 u = 0.85*c;    // Velocity of electron , m/s
5 m_p = 1.67e-27; // Rest mass of proton , kg
6
7 // Part (a)
8 E_o = m_p*c^2/1.602e-019; // Rest energy of proton
                               , MeV
```

```

9 printf("\nRest energy of proton = %3d MeV", E_o/1e
       +06);
10
11 // Part (b)
12 // Since given that E = 3*E_o = (3*m_p*c^2)/sqrt(1-
       u/c)^2), solving for u
13 u = sqrt(8/9)*c; // Velocity of proton, m/s
14 printf("\nVelocity of proton = %4.2fe+08 m/s", u*1e
       -08);
15
16 // Part (c)
17 // Since K = E - m_p*(c^2) = 3*m_p*(c^2) - m_p*(c^2)
       = 2*m_p*(c^2)
18 K = 2*E_o; // Kinetic energy of proton, MeV
19 printf("\nThe kinetic energy of proton = %4d MeV", K
       *1e-06);
20
21 // Part (d)
22 p = sqrt(8)*(E_o);
23 printf("\nThe momentum of proton = %4d MeV/c", p*1e
       -06);
24
25 // Result
26 // Rest energy of proton = 938 MeV
27 // Velocity of proton = 2.83e+08 m/s
28 // The kinetic energy of proton = 1876 MeV
29 // The momentum of proton = 2653 MeV/c

```

Scilab code Exa 2.5 Increase in mass of colliding balls

```

1 // Scilab code Ex2.5: Pg.49 (2005)
2 clc; clear;
3 u = 450; // Velocity of each ball, m/s
4 m = 5; // Mass of each ball, kg
5 c = 3e+08; // Velocity of light, m/s

```

```

6 // Since  $(u/c)^2 \ll 1$ , therefore, substituting  $1/\sqrt{1 - (u/c)^2} = 1 + (1/2)*(u/c)^2$ 
7 delta_M = m*(u/c)^2; // Increase in the mass of
   balls , kg
8 printf("\nIncrease in the mass of the balls = %3.1fe
   -11 kg" , delta_M*1e+11);
9
10 // Result
11 // Increase in the mass of the balls = 1.1e-11 kg

```

Scilab code Exa 2.7 A Fission Reaction

```

1 // Scilab code Ex2.7: Pg.50 (2005)
2 clc; clear;
3 u = 1.660e-27; // Atomic mass unit
4 M_U = 236.045563; // Atomic mass of Uranium , u
5 M_Rb = 89.914811; // Atomic mass of Rubidium , u
6 M_Cs = 142.927220; // Atomic mass of Caesium , u
7 m_n = 1.008665; // Mass of neutrons , u
8
9 // Part (a)
10 printf("\nU(92,235) --> Rb(37,90) + Cs(55,143) + 3n
   (0,1)");
11 printf("\nSo three neutrons are produced per fission
   .\n");
12
13 // Part (b)
14 delta_M = (M_U - (M_Rb + M_Cs + 3*m_n))*u; // Combined mass of all products , kg
15 printf("\nCombined mass of all products = %6.4fe -28
   kg\n" , delta_M*1e+28);
16
17 // Part (c)
18 // For simplification let velocity of light = 1 m/s
19 c = 1; // Velocity of light , m/s

```

```

20 // Since 1u = 931.5 MeV/(c^2) , therefore
21 Q = (delta_M/u)*931.5*(c^2);      // Energy given out
   per fission event , MeV
22 printf("\nEnergy given out per fission event = %5.1f
   MeV\n", Q);
23
24 // Part (d)
25 N = ((6.02e23)*1000)/236;      // Number of nuclei
   present
26 efficiency = 0.40;
27 E = efficiency*N*Q*(4.45e-20);    // Total energy
   released , kWh
28 printf("\nTotal energy released = %4.2fe+06 kWh\n",
   E*1e-06);
29
30 printf("\nThis amount of energy will keep a 100-W
   lightbulb burning for %d years", E
   *1000/(100*24*365));
31
32 // Result
33 // U(92,235) --> Rb(37,90) + Cs(55,143) + 3n(0,1)
34 // So three neutrons are produced per fission .
35 // Combined mass of all products = 2.9471e-28 kg
36 // Energy given out per fission event = 165.4 MeV
37 // Total energy released = 7.51e+06 kWh
38 // This amount of energy will keep a 100-W lightbulb
   burning for 8571 years

```

Scilab code Exa 2.8 Energy conservation

```

1 // Scilab code Ex2.8: Pg.52 (2005)
2 clc; clear;
3
4 // Part (a)
5 // Since 1 eV = 1.6e-19 J, therefore 3 eV = 3*1.6e

```

-19

```
6 BE = 3*1.6e-19; // Binding energy of water , J
7 c = 3e+08; // Velocity of light , m/s
8 delta_m = BE/(c^2); // Mass difference of water
molecule & its constituents , kg
9 printf("\nMass difference of water molecule & its
constituents = %3.1fe-36 kg", delta_m*1e+36);
10
11 // Part (b)
12 M = 3.0e-26; // Mass of water molecule , kg
13 M_f = delta_m/M; // Fractional loss of mass per
molecule
14 printf("\nThe fractional loss of mass per molecule =
%3.1fe-10", M_f*1e+10);
15
16 // Part (c)
17 E = M_f*(c^2); // Energy released when 1 g of
water is formed , kJ
18 printf("\nEnergy released when 1 g of water is
formed = %2.0f kJ", E*1e-06);
19
20 // Result
21 // Mass difference of water molecule & its
constituents = 5.3e-36 kg
22 // The fractional loss of mass per molecule = 1.8e
-10
23 // Energy released when 1 g of water is formed = 16
kJ
```

Scilab code Exa 2.9 Mass of Pion

```
1 // Scilab code Ex2.9: Pg.53 (2005)
2 clc; clear;
3 K_mu = 4.6; // Kinetic energy of muon , MeV
4 // For convenience let m_mew*(c^2) = E_mew
```

```
5 E_mu = 106;      // Energy of muon, MeV
6 E_pion = sqrt((E_mu^2) + (K_mu^2) + (2*K_mu*E_mu)) +
           sqrt((K_mu^2) + (2*K_mu*E_mu));
7 m_pion = E_pion; // Mass of pion, MeV/(c^2)
8 printf("\nMass of Pion = %3.0f MeV/(c^2)", m_pion);
9
10 // Result
11 // Mass of Pion = 142 MeV/(c^2)
```

Chapter 3

The Quantum theory of light

Scilab code Exa 3.1 Temperature of sun

```
1 // Scilab code Ex3.1: Pg 69 (2005)
2 clc; clear;
3 e_total = 1400;    // Total power per unit area , W/(m
4 sigma = 5.6e-08;   // Stefan-Boltzmann constant
5 R = 1.5e+11;      // Earth-sun distance , m
6 R_s = 7.0e+08;    // Radius of sun , m
7 // Using Stefan's law & solving for T, we get
8 T = (e_total*R^2/(sigma*R_s^2))^(1/4);    //
9 printf("\nThe temperature of sun = %4d K", T);
10
11 // Result
12 // The temperature of sun = 5820 K
```

Scilab code Exa 3.2 Quantum oscillator vs classical oscillator

```
1 // Scilab code Ex3.2: Pg 75 (2005)
```

```

2 clc; clear;
3
4 // Part (a)
5 h = 6.63e-34; // Plank's constant , Js
6 c = 3e+08; // Velocity of light , m/s
7 lamda_green = 540e-09; // Wavelength of green
    light , nm
8 delta_E_green = h*c/lamda_green/1.602e-19; // Minimum
    energy change in green light , eV
9 lamda_red = 700e-09; // Wavelength of red light ,
    nm
10 delta_E_red = h*c/lamda_red/1.602e-19; // Minimum
    energy change in red light , eV
11
12 printf("\nMinimum energy change in green light = %4
    .2f eV" , delta_E_green);
13 printf("\nMinimum energy change in red light = %4.2f
    eV" , delta_E_red);
14
15 // Part (b)
16 f = 0.50; // Frequency , Hz
17 m = 0.1; // Mass of pendulum , kg
18 l = 1; // Length of pendulum , m
19 theta = %pi/180*10; // Angle , radians
20 g = 9.8; // Acceleration due to gravity , m/s^2
21 E = m*g*l*(1-cos(theta));
22 delta_E = (h*f)/(1.6e-19); // Minimum energy
    change in pendulum , eV
23 delta_E_f = (delta_E*1.6e-19)/E ; // Fractional
    energy change
24 printf("\nFractional energy change = %3.1fe-32" ,
    delta_E_f*1e+32);
25
26 // Result
27 // Minimum energy change in green light = 2.30 eV
28 // Minimum energy change in red light = 1.77 eV
29 // Fractional energy change = 2.2e-32

```

Scilab code Exa 3.3 Stefan law from Planck distribution

```
1 // Scilab code Ex3.3: Pg 80 (2005)
2 clc; clear;
3 k_B = 1.381e-23; // Boltzmann's constant, J/K
4 c = 3e+08; // Velocity of light, m/s
5 h = 6.626e-34; // Plank's constant, Js
6 // Since e_total = sigma*(T^4) = (2*(%pi)^5*(k_B)^4)
   /(15*(c^2)*(h^3))*T^4
7 sigma = (2*(%pi)^5*(k_B)^4)/(15*(c^2)*(h^3));
8 printf("\nThe value of sigma = %3.2fe-08 W/Sq.m/K^4"
       , sigma*1e+08);
9
10 // Result
11 // The value of sigma = 5.67e-08 W/Sq.m/K^4
```

Scilab code Exa 3.4 Time lag between start of illumination and photocurrent generation

```
1 // Scilab code Ex3.4: Pg 83 (2005)
2 clc; clear;
3 phi = 2.38; // Work function for sodium, eV
4 I = 1e-07; // Absorbed light intensity, mJcm^2/s
5 A = %pi*1e-16; // Cross-sectional area, m^2
6 t = phi*1.6e-16/(I*A) // Time lag, days
7 printf("\nTime lag between start of illumination and
       photocurrent generation = %3.1fe+07 s", t*1e-07)
     ;
8
9 // Result
10 // Time lag between start of illumination and
    photocurrent generation = 1.2e+07 s
```

Scilab code Exa 3.5 Time lag between start of illumination and photocurrent generation

```
1 // Scilab code Ex3.5: Pg 85 (2005)
2 clc; clear;
3 e = 1.6e-19;      // Electric charge, C
4 V_s = 4.3;        // Stopping potential, V
5 K_max = e*V_s;    // Maximum kinetic energy attained
                     by photoelectrons, J
6 m_e = 9.11e-31   // Mass of electron, kg
7 // Since K.E = eV_s = 0.5m_e(v_max^2), therefore
8 v_max = sqrt((2*K_max)/m_e);    // Maximum velocity
                     attained by photoelectron, m/s
9 printf("\nMaximum velocity attained by photoelectron
       = %3.1fe+06 m/s", v_max*1e-06);
10
11 // Result
12 // Maximum velocity attained by photoelectron = 1.2e
       +06 m/s
```

Scilab code Exa 3.6 Photoelectric effect for iron

```
1 // Scilab code Ex3.6: Pg 85 (2005)
2 clc; clear;
3
4 // Part (a)
5 I_o = 1;        // Total intensity of light, micro-W/cm
                  ^2
6 I = (0.030)*(0.040)*I_o;    // Intensity available to
                  produce photoelectric effect, nW/cm^2
7 printf("\nIntensity available to produce
       photoelectric effect = %3.1f nW/cm^2", I*1e+03);
8
```

```

9 // Part (b)
10 h = 6.6e-34; // Planck's constant , Js
11 c = 3e+08; // Velocity of light , m/s
12 lamda = 250e-09; // Wavelength , m
13 e_per_sec = (I*lamda*1e-06)/(h*c); // Number of
   electrons emitted per second
14 printf("\nNumber of electrons emitted per second =
   %3.1e", e_per_sec);
15
16 // Part (c)
17 e = 1.6e-019; // Energy equivalent of 1 eV, C
18 i = (e_per_sec)*e; // Electric current in
   phototube , A
19 printf("\nElectric current in phototube = %3.1e A" ,
   i);
20
21 // Part (d)
22 f_o = 1.1e+15; // Cut-off frequency , Hz
23 phi = (h*f_o)/e; // Work function for iron , eV
24 printf("\nWork function for iron = %3.1f eV" , phi);
25
26 // Part (e)
27 V_s = (h*c/(e*lamda))-phi; // Stopping
   voltage , V
28 printf("\nStopping voltage = %4.2f V" , V_s);
29
30 // Result
31 // Intensity available to produce photoelectric
   effect = 1.2 nW/cm^2
32 // Number of electrons emitted per second = 1.5e-09
33 // Electric current in phototube = 2.4e-10 A
34 // Work function for iron = 4.5 eV
35 // Stopping voltage = 0.41 V

```

Scilab code Exa 3.7 Compton shift for carbon

```

1 // Scilab code Ex3.7: Pg 93 (2005)
2 clc; clear;
3 h = 6.63e-34;      // Plank's constant , Js
4 m_e = 9.11e-31;    // Mass of electron , kg
5 c = 3e+08;         // Velocity of light , m/s
6 theta = ((%pi)/180)*45; // Angle , radians
7 delta_lamda = (h/(m_e*c)*(1-cos(theta))); // Compton shift , nm
8 lamda_o = 0.200e-09; // Wavelength of X-ray , nm
9 lamda = delta_lamda+lamda_o // Increased wavelength of scattered X-ray , nm
10 printf("\nIncreased wavelength of scattered X-ray = %8.6f nm", lamda*1e+09);
11
12 // Result
13 // Increased wavelength of scattered X-ray =
0.200711 nm

```

Scilab code Exa 3.8 Xray photons vs visible photons

```

1 // Scilab code Ex3.8: Pg 93 (2005)
2 clc; clear;
3
4 // Part (a)
5 h = 6.63e-34;      // Plank's constant , Js
6 q = 1.6e-19;        // Electric charge , C
7 m_e = 9.11e-31;    // Mass of electron , kg
8 c = 3e+08;          // Velocity of light , m/s
9 theta = ((%pi)/180)*90; // Angle , radians
10 delta_lamda = (h/(m_e*c)*(1-cos(theta)));
                           // Compton shift , Angstrom
11 lamda_C = 0.0106;   // Wavelength of gamma-rays from Cobalt ,
12 f_dl_C = delta_lamda/ lamda_C; // Fractional change in wavelength of gamma rays from cobalt

```

```

13 printf("\nFractional change in wavelength of gamma
       rays from Cobalt = %4.2f", f_dl_C*1e+10);
14 lamda_Mo = 0.712;      // Wavelength of gamma-rays
       from Molybdenum, Angstrom
15 f_dl_Mo = delta_lamda/ lamda_Mo;                      //
       Fractional change in wavelength of gamma rays
       from Molybdenum
16 printf("\nFractional change in wavelength of gamma
       rays from Molybdenum = %6.4f", f_dl_Mo*1e+10);
17 lamda_Hg = 5461;      // Wavelength of gamma-rays from
       Mercury, Angstrom
18 f_dl_Hg = delta_lamda/ lamda_Hg;                      //
       Fractional change in wavelength of gamma rays
       from mercury
19 printf("\nFractional change in wavelength of gamma
       rays from Mercury = %4.2fe-06", f_dl_Hg*1e+16);
20
21 // Part (b)
22 lamda = 0.712e-10;      // Wavelength of X-rays ,
       Angstrom
23 E = (h*c)/(q*lamda);    // Energy of X-rays ' photon ,
       eV
24 printf("\nEnergy of X-rays photon = %5.0f eV\n", E);
25
26 // Result
27 // Fractional change in wavelength of gamma rays
       from Cobalt = 2.29
28 // Fractional change in wavelength of gamma rays
       from Molybdenum = 0.0341
29 // Fractional change in wavelength of gamma rays
       from Mercury = 4.45fe-06
30 // Energy of X-rays photon = 17460 eV

```

Scilab code Exa 3.9 Gravitational redshift for a white dwarf

```
1 // Scilab code Ex3.9: Pg 96 (2005)
2 clc; clear;
3 M = 1.99e+30;      // Mass of sun , kg
4 R_s = 6.37e+06;    // Radius of earth , m
5 G = 6.67e-11;     // Gravitational constant , Nm^2/kg^2
6 lamda = 300e-09;   // Wavelength , m
7 c = 3e+08;         // Velocity of light , m/s
8 delta_lamda = lamda*((G*M)/(R_s*c^2));    //
    Gravitational redshift , angstrom
9 printf("\nGravitational redshift = %3.1f angstrom" ,
    delta_lamda*1e+10);
10
11 // Result
12 // Gravitational redshift = 0.7 angstrom
```

Chapter 4

The particle nature of matter

Scilab code Exa 4.1 Electrolysis of barium chloride

```
1 // Scilab code Ex4.1: Pg 109 (2005)
2 clc; clear;
3 I = 10;      // Electric current , A
4 t = 3600;    // Time , s
5 q = I*t;    // Electric charge liberated , C
6 mm_Ba = 137; // Molar mass of Barium, g
7 mm_Cl = 35.5; // Molar mass of Chlorine , g
8 valence_Ba = 2; // Valence electrons of Barium
9 valence_Cl = 1; // Valence electrons of
                  Chlorine
10 // Using Faradays law of electrolysis , we have
11 m_Ba = (q*mm_Ba)/(96500*valence_Ba); // Mass of
                                             Barium obtained , g
12 m_Cl = (q*mm_Cl)/(96500*valence_Cl); // Mass of
                                             Chlorine obtained , g
13 printf("\nMass of Barium obtained = %4.1f g", m_Ba);
14 printf("\nMass of Chlorine obtained = %4.1f g", m_Cl
      );
15
16 // Result
17 // Mass of Barium obtained = 25.6 g
```

18 // Mass of Chlorine obtained = 13.2 g

Scilab code Exa 4.2 Deflection of electron beam by E and B Fields

```
1 // Scilab code Ex4.2: Pg 113 (2005)
2 clc; clear;
3 V = 200;      // Electric potential , V
4 theta = 0.20; // Angle , radians
5 l = 0.050;    // Length of plates , m
6 d = 1.5e-02; // Distance between two plates , m
7 c_m_r = 1.76e+11; // Charge-to-mass ratio , C/kg
8 // Since e/m_e = (V*theta)/(B^2*l*d) , solving for B
9 B = sqrt((V*theta)/(l*d*c_m_r)); // Magnetic
   field , T
10 printf("\nThe magnetic field required to produce the
        deflection of %4.2f rad = %3.1e T", theta, B);
11
12 // Result
13 // The magnetic field required to produce the
   deflection of 0.20 rad = 5.5e-04 T
```

Scilab code Exa 4.3 Experimental determination of e

```
1 // Scilab code Ex4.3: Pg 117 (2005)
2 clc; clear;
3
4 // Part (a)
5 delta_y = 0.600; // Distance of rise or fall of a
   droplet , cm
6 t_av = 21.0; // Average time of fall of droplet ,
   s
7 delta_t = [46.0, 15.5, 28.1, 12.9, 45.3, 20.0]; // 
   Rise time of the droplet in succession , s
```

```

8 v = delta_y/t_av; // Average speed of the falling
droplet , cm/s
9 v_prime = zeros(6);
10 for i = 1:1:6
11     v_prime(i) = delta_y/delta_t(i); // Successive
speeds of the rising drops , cm/s
12 end
13
14 // Calculate charge ratios
15 q1byq2 = (v+v_prime(1))/(v + v_prime(2));
16 q2byq3 = (v+v_prime(2))/(v + v_prime(3));
17 q3byq4 = (v+v_prime(3))/(v + v_prime(4));
18 q4byq5 = (v+v_prime(4))/(v + v_prime(5));
19 q5byq6 = (v+v_prime(5))/(v + v_prime(6));
20 printf("\nq1/q2 = %5.3f", q1byq2);
21 printf("\nq2/q3 = %5.3f", q2byq3);
22 printf("\nq3/q4 = %5.3f", q3byq4);
23 printf("\nq4/q5 = %5.3f", q4byq5);
24 printf("\nq5/q6 = %5.3f", q5byq6);
25 printf("\nThe charge ratios are ratios of small
whole numbers\n");
26
27 // Part (b)
28 eta = 1.83e-05; // Viscosity of
air , kg/ms
29 rho = 858; // Oil density , kg
/m^3
30 g = 9.81; // Acceleration due
to gravity , m/s^2
31 a = sqrt((9*eta*v*1e-02)/(2*rho*g)); // Radius of
oil droplet , m
32 V = 4/3*(pi)*a^3; // Volume of oil
droplet , m^3
33 m = rho*V; // Mass of oil
droplet , kg
34 printf("\nRadius of oil droplet = %4.2e m", a);
35 printf("\nVolume of oil droplet = %4.2e m^3", V);
36 printf("\nMass of oil droplet = %4.2e kg", m);

```

```

37
38 // Part (c)
39 V = 4550; // Potential difference across the
             plates of the capacitor , volt
40 d = 0.0160; // Distance between the plates
41 E = V/d; // Electric field between plates , V/m
42 q = zeros(6), e = zeros(6);
43 for i=1:1:6
44     q(i) = m*g/E*((v+v_prime(i))/v); // Charge on
         first drop , C
45     printf("\nq%d = %4.2e V/m" , i , q(i));
46 end
47 e(1) = q(1)/5;
48 e(2) = q(2)/8;
49 e(3) = q(3)/6;
50 e(4) = q(4)/9;
51 e(5) = q(5)/5;
52 e(6) = q(6)/7;
53 e_tot = 0;
54 for i = 1:1:6
55     e_tot = e_tot + e(i);
56 end
57 e = e_tot/6;
58 printf("\nThe average charge on an electron = %5.3e
          C" , e);
59
60 // Result
61 // q1/q2 = 1.105
62 // q2/q3 = 0.958
63 // q3/q4 = 1.053
64 // q4/q5 = 0.899
65 // q5/q6 = 1.086
66 // The charge ratios are ratios of small whole
      numbers
67
68 // Radius of oil droplet = 1.67e-06 m
69 // Volume of oil droplet = 1.96e-17 m^3
70 // Mass of oil droplet = 1.68e-14 kg

```

```

71
72 // q1 = 8.44e-019 V/m
73 // q2 = 1.36e-018 V/m
74 // q3 = 1.01e-018 V/m
75 // q4 = 1.52e-018 V/m
76 // q5 = 8.48e-019 V/m
77 // q6 = 1.19e-018 V/m
78 // The average charge on an electron = 1.694e-019 C

```

Scilab code Exa 4.4 Collision of alpha particle with proton

```

1 // Scilab code Ex4.4: Pg 121 (2005)
2 clc; clear;
3
4 // Part (b)
5 // For easy calculations , assume all variables to be
   unity
6 m_p = 1;      // Mass of proton , a.m.u
7 m_a = 4*m_p;    // Mass of alpha particle , a.m.u
8 Valpha = 1;    // Velocity of alpha particle before
   collision , m/s
9 v_p = (2*m_a*Valpha)/(m_a + m_p);    // Velocity of
   proton after collision , m/s
10 v_a = ((m_a - m_p)*(Valpha))/(m_a + m_p);   //
   Velocity of alph particle after collision , m/s
11 p_change = ((v_a - Valpha)/(Valpha))*100;   //
   Percentage change in velocity of alpha particle
12 printf("\nVelocity of proton after collision = %4.2
   fVa m/s" , v_p);
13 printf("\nVelocity of alpha particle after collision
   = %4.2fVa m/s" , v_a);
14 printf("\nPercentage change in velocity of alpha
   particle = %2d percent" , p_change);
15
16 // Result

```

```
17 // Velocity of proton after collision = 1.60 V_a m/s
18 // Velocity of alph particle after collision = 0.60
    V_a m/s
19 // Percentage change in velocity of alpha particle =
    -40 percent
```

Scilab code Exa 4.5 Radius of Aluminium Nucleus

```
1 // Scilab code Ex4.5: Pg 124 (2005)
2 clc; clear;
3 Z = 13;      // Atomic number of Aluminium
4 e = 1.6e-19; // Charge on electron , C
5 k = 8.99e+09; // Coulomb constant , Nm^2/C^2
6 K_a = 7.7e+06*e; // Since K_a = (k*Z_e*2*e)/d_min
                    , solving for d_min
7 d_min = (k*2*Z*e^2)/K_a; // Radius of Aluminum , m
8 printf("\nRadius of Aluminum = %3.1e m", d_min);
9
10 // Result
11 // Radius of Aluminum = 4.9e-15 m
```

Scilab code Exa 4.7 Collision of alpha particle with proton

```
1 // Scilab code Ex4.7: Pg 135 (2005)
2 clc; clear;
3 // Part (a)
4 n_i = 2;      // Initial level of electron
5 n_f = 1;      // Final level of electron
6 R = 1.097e+07; // Rydberg constant , per metre
7 c = 3e+08;    // Velocity of light , m/s
8 h = 4.136e-15; // Planck's constant , eV
9 lambda = n_i^2*n_f^2/((n_i^2-n_f^2)*R); //
                    Wavelength of emitted photon , m
```

```

10 f = c/lamda;      // Frequency of emitted photon , Hz
11 E = h*f;          // Energy of emitted photon , eV
12 printf("\nThe wavelength of emitted photon = %5.1f
13 nm" , lamda/1e-09);
13 printf("\nThe frequency of emitted photon = %4.2e Hz
14 " , f);
14 printf("\nEnergy of emitted photon = %4.1f eV" , E);
15
16 // Part (b)
17 mc_square = 938.8e+06;    // Energy of recoil of
18 hydrogen atom , eV
18 K = 0.5*(E^2/mc_square); // Recoil kinetic
19 energy of H atom , eV
19 E_difference = K/E;     // Energy difference
20 printf("\nRecoil kinetic energy of H atom = %4.2e eV
21 " , K);
21 printf("\nThe fraction of energy difference = %3.1e"
22 , E_difference);
22
23 // Result
24 // The wavelength of emitted photon = 121.5 nm
25 // The frequency of emitted photon = 2.47e+15 Hz
26 // Energy of emitted photon = 10.2 eV
27 // Recoil kinetic energy of H atom = 5.55e-08 eV
28 // The fraction of energy difference = 5.4e-09

```

Scilab code Exa 4.8 series for Hydrgen

```

1 // Scilab code Ex4.8: Pg 136 (2005)
2 clc; clear;
3
4 // Part (a)
5 n_i = 3;      // Initial level of electron
6 n_f = 2;      // Final level of electron
7 R = 1.097e+07; // Rydberg constant , per metre

```

```

8 c = 3e+08;           // Velocity of light , m/s
9 h = 6.626e-34;       // Plank's constant , Js
10 lamda_max = (n_i^2*n_f^2)/((n_i^2-n_f^2)*R);      //
    Maximum wavelength of emitted photon , m
11 E_photon = (h*c)/(lamda_max*1.6e-19);                //
    Energy of emitted photon , eV
12 printf("\nThe maximum wavelength of emitted photon = "
    "%5.1f nm" , lamda_max/1e-09);
13 printf("\nEnergy of emitted photon = %4.2f eV" ,
    E_photon);
14
15 // Part (b)
16 n_i = %inf;        // Initial level of electron
17 lamda_min = 1/(R*(1/n_f^2-1/n_i^2));
18 printf("\nThe wavelength corresponding to the series
    limit = %5.1f nm which is in the ultraviolet
    region" , lamda_min/1e-09);
19
20 // Result
21 // The maximum wavelength of emitted photon = 656.3
    nm
22 // Energy of emitted photon =1.89 eV
23 //// The wavelength corresponding to the series
    limit = 364.6 nm which is in the ultraviolet
    region

```

Scilab code Exa 4.9 Hydrogen in its first excited state

```

1 // Scilab code Ex4.9: Pg 137 (2005)
2 clc; clear;
3 k_B = 8.62e-05;      // Boltzmann constant , eV/K
4 delta_E = 10.2;       // Average thermal energy , eV
5 // Since (3/2)*k_B*T = average thermal energy per
    atom = 10.2eV, solving for T
6 T = 10.2/(3/2*k_B);   // Temperature at which H-

```

```

        atoms jump to first excited state , K
7 printf("\nThe temperature at which H-atoms jump to
       first excited state = %5d K", T);
8 N_ratio = 0.10;      // Number ratio of population of
       first excited state relative to the ground state
9 // As N_ratio = exp(-delta_E/(k_B*T)), solving for T
10 T = -delta_E/(k_B*log(N_ratio));      // Temperature
       at which H-atoms jump to first excited state , K
11 printf("\nThe temperature of excitation from
       Boltzmann distribution = %5d K", T);
12
13 // Result
14 // The temperature at which H-atoms jump to first
       excited state = 78886 K
15 // The temperature of excitation from Boltzmann
       distribution = 51389 K

```

Chapter 5

Matter waves

Scilab code Exa 5.1 Wave properties of a baseball

```
1 // Scilab code Ex5.1: Pg 154 (2005)
2 clc; clear;
3 h = 6.63e-34;          // Plank's constant , Js
4 m = 140e-03;           // Mass of baseball , kg
5 v = 27;                // Velocity of baseball , m/s
6 p = m*v;               // Momentum of baseball , kgm/s
7 lamda = h/p;            // de Broglie wavelength associated
                           with baseball , m
8 printf("\nde-Broglie wavelength associated with
baseball = %3.1e m", lamda);
9
10 // Result
11 // de-Broglie wavelength associated with baseball =
1.8e-34 m
```

Scilab code Exa 5.2 de Broglie wavelength of an electron

```
1 // Scilab code Ex5.2: Pg 154 (2005)
```

```

2 clc; clear;
3
4 // Part (b)
5 h = 6.63e-34;      // Plank's constant , Js
6 m_e = 9.11e-31;    // Mass of electron , kg
7 q = 1.6e-19;       // Charge on electron , C
8 V = 50;            // Electric potential applied , V
9 lamda = h/(sqrt(2*m_e*q*V));      // de Broglie
   wavelength of an electron , m
10 printf("\nde Broglie wavelength of an electron = %3
.1f angstrom", lamda/1e-10);
11
12 // Result
13 // de Broglie wavelength of an electron = 1.7
   angstrom

```

Scilab code Exa 5.3 Diffraction of neutrons at the crystal lattice

```

1 // Scilab code Ex5.3: Pg 158 (2005)
2 clc; clear;
3 h = 6.63e-34;      // Plank's constant , J-s
4 lamda = 1e-10;      // de Broglie wavelength of
   neutron , m
5 p = h/lamda;        // Momentum associated with neutron
   , kg-m/s
6 m_n = 1.66e-27;     // Mass of neutron , kg
7 e = 1.6e-19;        // Energy equivalent of 1 eV, J/eV
8 K = p^2/(2*m_n);    // Kinetic energy of neutron , eV
9 printf("\nThe momentum of neutrons = %4.2e kg-m/s",
   p)
10 printf("\nThe kinetic energy of neutrons = %4.2fe-20
   J = %6.4f eV", K*1e+20, K/e);
11
12 // Result
13 // The momentum of neutrons = 6.63e-24 kg-m/s

```

```
14 // The kinetic energy of neutrons = 1.32e-20 J =
0.0828 eV
```

Scilab code Exa 5.8 Uncertainty principle for macroscopic objects

```
1 // Scilab code Ex5.8: Pg 177 (2005)
2 clc; clear;
3 h_cross = 1.05e-34;      // Reduced Plank's constant ,
J-s
4 delta_x = 15;           // Uncertainty in position , m
5 v_x = 2;                // Velocity of ball , m/s
6 m = 100e-03;            // Mass of ball , kg
7 delta_p_x = h_cross/(2*delta_x);    // Uncertainty
in momentum, kg-m/s
8 delta_v_x = delta_p_x/m;        // Minimum spread in
velcoity , m/s
9 U_r = delta_v_x/v_x;        // Relative uncertainty in
velocity of ball
10 printf("\nThe minimum spread in velcoity of ball =
%3.1e m/s" , delta_v_x);
11 printf("\nThe relative uncertainty in velocity of
ball = %4.2e" , U_r);
12
13 // Result
14 // The minimum spread in velcoity of ball = 3.5e-35
m/s
15 // The relative uncertainty in velocity of ball =
1.75e-35
```

Scilab code Exa 5.9 Kinetic energy of electron confined within the nucleus

```
1 // Scilab code Ex5.9: Pg 178 (2005)
2 clc; clear;
```

```

3 delta_x = 1.0e-14/2; //  

    Uncertainty in position of electron , m  

4 q = 1.6e-19; // Charge on  

    electron , C  

5 h_cross = 1.05e-34; // Reduced Plank ' s constant , J-s  

6 c = 3e+08; // Velocity of light , m/s  

7 delta_p_x = (h_cross*c)/(2*delta_x*q); // Uncertainty in momentum , eV/c  

8 E_r = 0.551e+06; // Rest mass energy if electron , eV  

9 E = sqrt((delta_p_x)^2 + (E_r)^2);  

10 K = E - E_r; // Kinetic energy of electron within nucleus , eV  

11 printf("\nKinetic energy of electron within nucleus = %4.1f MeV", K/1e+06);  

12  

13 // Result  

14 // Kinetic energy of electron within nucleus = 19.1 MeV

```

Scilab code Exa 5.10 Width of spectral lines

```

1 // Scilab code Ex5.10: Pg 178 (2005)  

2 clc; clear;  

3  

4 // Part (a)  

5 h_cross = 1.05e-34; // Reduced Plank ' s constant , J-s  

6 h = 6.63e-34; // Plank ' s constant , J-s  

7 delta_t = 1.0e-08; // Average time to measure the excited state , s  

8 delta_E = h_cross/(2*delta_t); // Uncertainty in

```

```

        energy of the excited state , J
9 // Since delta_E = h*delta_f , solving for delta_f
10 delta_f = delta_E/h;      // Line width of emitted
    light , Hz
11 printf("\nLine width of emitted light = %2.0e Hz",
    delta_f);
12
13 // Part (b)
14 c = 3e+08;      // Velocity of light , m/s
15 lamda = 500e-09;      // Wavelength of spectral line
    , m
16 f_o = c/lamda;      // Center frequency of spectral
    line , Hz
17 f_b = delta_f/f_o;      // Fractional broadening of
    spectral line
18 printf("\nFractional broadening of spectral line =
    %3.1e" , f_b);
19
20 // Result
21 // Line width of emitted light = 8.0e+06 Hz
22 // Fractional broadening of spectral line = 1.3e-08

```

Chapter 6

Quantum mechanics in one dimension

Scilab code Exa 6.2 Probability from wave function

```
1 // Scilab code Ex6.2: Pg 193 (2005)
2 clc; clear;
3 x0 = 1;      // For simplicity assume x0 = 1
4 C = 1/sqrt(x0);    // Normalization constant
5 P = 2*C^2*integrate('exp(-2*x/x0)', 'x', 0, x0);
6 printf("\nThe probability that the particle will be
         found in the interval -x0 <= x <= x0 is %6.4f or
         %4.1f percent", P, P*100);
7
8 // Result
9 // The probability that the particle will be found
   in the interval -x0 <= x <= x0 is 0.8647
```

Scilab code Exa 6.4 Dispersion of matter waves

```
1 // Scilab code Ex6.4: Pg 197 (2005)
```

```

2 clc; clear;
3 delta_x0 = 1e-010; // Initial width of the
    localized space, m
4 delta_xt = 10*delta_x0; // Final width at which
    the wave packet is dispersed, m
5 h_cross = 1.055e-034; // Reduced Planck's
    constant, Js
6 m = 9.11e-031; // Mass of the electron, kg
7 // From Dispersion relation, delta_xt^2 - delta_x0^2
    = sqrt(h_cross*t/(2*m*delta_x0)^2), solving for t
8 t = 2*m*sqrt(delta_xt^2 - delta_x0^2)*delta_x0/
    h_cross; // Time which elapses before
    delocalization
9 printf("\nThe time which elapses before the
    localization of electron destroys = %3.1e s", t);
10 m = 1e-03; // Mass of marble, kg
11 delta_x0 = 1e-004; // Initial width of the
    localized space, m
12 delta_xt = 10*delta_x0; // Final width at which
    the wave packet is dispersed, m
13 t = 2*m*sqrt(delta_xt^2 - delta_x0^2)*delta_x0/
    h_cross; // Time which elapses before
    delocalization
14 printf("\nThe time which elapses before the
    localization of marble destroys = %3.1e s", t);
15 printf("\nFor all the practical purposes, the marble
    will remain localized for ever");
16 // Result
17 //

```

Scilab code Exa 6.5 Energy Quantization for Macroscopic Object

```

1 // Scilab code Ex6.5: Pg 202 (2005)
2 clc; clear;
3 h = 6.626e-034; // Planck's constant, Js

```

```

4 m = 1e-06;      // Mass of the object , kg
5 n = 1;          // Quantum number for minimum energy level
6 L = 1e-02;       // Distance between two rigid walls ,
                     m
7 E1 = n^2*h^2/(8*m*L^2); // Minimum energy of the
                           object , J
8 v1 = sqrt(2*E1/m);    // Minimum speed of the object
                           , m/s
9 v = 3.00e-02;        // Given speed of the object , m/s
10 E = 1/2*m*v^2;     // Energy of the object for given
                      speed , J
11 n = sqrt(8*m*L^2*E)/h; // Quantum number
                           corresponding to the given speed
12 printf("\nThe minimum speed of the object = %4.2e m/
           s", v1);
13 printf("\nThe quantum number corresponding to the
           speed of %4.2e m/s is n = %4.2e", v1, n);
14
15 // Result
16 // The minimum speed of the object = 3.31e-26 m/s
17 // The quantum number corresponding to the speed of
           3.31e-26 m/s is n = 9.06e+23

```

Scilab code Exa 6.6 Model of an Atom

```

1 // Scilab code Ex6.6: Pg 203 (2005)
2 clc; clear;
3 c = 1;          // Assume speed of light to be unity , m/
                     s
4 h_cross = 197.3; // Reduced Planck's constant , eV
                     .nm/c^2
5 m_e = 511e+03;   // Mass of an electron , eV/c
                     ^2
6 L = 0.200;       // Length of the box , nm
7 E1 = %pi^2*(h_cross/c)^2/(2*m_e*L^2); // Ground

```

```

        state energy of atomic electron , eV
8 E2 = 2^2*E1;      // Excited state energy of the
                      atomic electron , eV
9 delta_E = E2- E1; // Energy that must be applied
                      to the electron to raise it from ground to the
                      first excited state , eV
10 h = 2*pi*h_cross; // Planck's constant , Js
11 lambda = h*c/delta_E; // Wavelength of the photon
                          to cause the electron transition , nm
12 printf("\nThe energy that must be applied to the
          electron to raise it from ground to the first
          excited state = %4.1f eV", delta_E);
13 printf("\nThe wavelength of the photon to cause this
          electron transition = %4.1f nm", lambda);
14 printf("\nThis wavelength is in the far ultraviolet
          region.");
15
16 // Result
17 // The energy that must be applied to the electron
          to raise it from ground to the first excited
          state = 28.2 eV
18 // The wavelength of the photon to cause this
          electron transition = 44.0 nm
19 // This wavelength is in the far ultraviolet region.

```

Scilab code Exa 6.7 Probabilities for a particle in a Box

```

1 // Scilab code Ex6.7: Pg 205 (2005)
2 clc; clear;
3 L = 1; // For simplicity assume length of finite
          square well to be unity , m
4 P = 2/L*integrate('sin(%pi*x/L)^2', 'x', L/4, 3*L/4)
          ; // Probability that the particle will be found
          in the middle half of the well
5 printf("\nThe probability that the particle will be

```

```

        found in the middle half of the well = %5.3 f" , P)
;
6
7 // Result
8 // The probability that the particle will be found
   in the middle half of the well = 0.818

```

Scilab code Exa 6.8 Ground state energy of an electron confined to a potential well

```

1
2 // Scilab code Ex6.8: Pg 211 (2005)
3 clc; clear;
4 c = 1;           // Assume speed of light to be unity , m/
   s
5 L = 0.200;      // Width of the potential well , nm
6 h_cross = 197.3;    // Reduced Planck's constant , eV
   .nm/c^2
7 m = 511e+03;      // Mass of an electron , eV/c^2
8 U = 100;          // Height of potential well , eV
9 delta = h_cross/sqrt(2*m*U);      // Decay length of
   electron , nm
10 L = L + 2*delta;     // Effective length of the
   infinite potential well , nm
11 E = %pi^2*(h_cross/c)^2/(2*m*L^2);    // Ground state
   energy of the electron with effective length , eV
12 U = U - E;          // New potential energy , eV
13 delta = h_cross/sqrt(2*m*U);      // New decay length
   of electron , nm
14 printf("\nThe ground state energy of an electron
   confined to the potential well = %4.2f eV" , E);
15 printf("\nThe new decay length of the electron = %6
   .4f nm" , delta);
16
17 // Result
18 // The ground state energy of an electron confined

```

```
    to the potential well = 6.58 eV  
19 // The new decay length of the electron = 0.0202 nm
```

Scilab code Exa 6.12 The quantum oscillator in nonclassical region

```
1 // Scilab code Ex6.12: Pg 214 (2005)  
2 clc; clear;  
3 P = 2/sqrt(%pi)*integrate('exp(-z^2)', 'z', 1, 100);  
    // Probability that the quantum oscillator in  
    its ground state will be found in the  
    nonclassical region  
4 printf("\nThe probability that the quantum  
    oscillator in its ground state will be found in  
    the nonclassical region = %5.3f", P);  
5  
6 // Result  
7 // The probability that the quantum oscillator in  
    its ground state will be found in the  
    nonclassical region = 0.157
```

Scilab code Exa 6.13 Quantization of vibrational energy

```
1 // Scilab code Ex6.13: Pg 211 (2005)  
2 clc; clear;  
3 h_cross = 6.582e-016; // Reduced Planck's constant  
    , eV-s  
4 // For spring-mass system  
5 K = 0.100; // Force constant of the spring-mass  
    system , N/m  
6 m = 0.0100; // Mass attached to the spring , kg  
7 omega = sqrt(K/m); // Angular frequency of  
    oscillations , rad/s
```

```

8 delta_E = h_cross*omega;      // Energy spacing
   between quantum levels , eV
9 printf("\nThe energy spacing between quantum levels
   for spring-mass system = %4.2e eV\nwhich is far
   below present limits of detection", delta_E);
10 // For vibrating hydrogen molecule
11 K = 510.5; // Force constant of the hydrogen
   molecule system , N/m
12 mu = 8.37e-028; // Reduced mass of the hydrogen
   molecule , kg
13 omega = sqrt(K/mu); // Angular frequency of
   oscillations , rad/s
14 delta_E = h_cross*omega;      // Energy spacing
   between quantum levels , eV
15 printf("\nThe energy spacing between quantum levels
   for hydrogen molecule = %5.3f eV\nwhich can be
   measured easily", delta_E);
16
17 // Result
18 // The energy spacing between quantum levels for
   spring-mass system = 2.08e-15 eV
19 // which is far below present limits of detection
20 // The energy spacing between quantum levels for
   hydrogen molecule = 0.514 eV
21 // which can be measured easily

```

Scilab code Exa 6.14 Standard Deviations from Averages

```

1 // Scilab code Ex6.14: Pg 219 (2005)
2 clc; clear;
3 x = [2.5, 3.7, 1.4, 7.9, 6.2, 5.4, 8.0, 6.4, 4.1,
   5.4, 7.0, 3.3, 4.2, 8.8, 6.2, 7.1, 5.4, 5.3]; // 
   Data entries
4 sum_x = 0;      // Initialize the accumulator
5 sum_x_sq = 0;    // Initialize the second accumulator

```

```

6 N = 18;      // Total number of data points
7 for i = 1:1:N
8     sum_x = sum_x + x(i);    // Sum of data
9     sum_x_sq = sum_x_sq + x(i)^2; // Sum of square
10    of data
11 end
12 x_av = sum_x/N;      // Average of data
13 x_sq_av = sum_x_sq/N; // Mean square value
14 sigma = sqrt(x_sq_av-x_av^2); // Standard
15    deviation from averages
16
17 // Result
18 // The standard deviation from averages = 1.93

```

Scilab code Exa 6.15 Location of a particle in the box

```

1 // Scilab code Ex6.15: Pg 219 (2005)
2 clc; clear;
3 L = 1; // For simplicity assume length of the box
4 x_av = 2*L/%pi^2*integrate('theta*sin(theta)^2', '
5 theta', 0, %pi); // Average value of x
6 x_sq_av = L^2/%pi^3*(integrate('theta^2', 'theta',
7 0, %pi)-integrate('theta^2*cos(2*theta)', 'theta',
8 0, %pi)); // Average value of x square
9 delta_x = sqrt(x_sq_av - x_av^2); // Uncertainty
10   in the position for this particle, unit
11 printf("\nThe average position of the particle in
12 the box = L/%ld", x_av*4);
13 printf("\nThe uncertainty in the position for the
14 particle = %5.3fL", delta_x);

```

```
10 // Result
11 // The average position of the particle in the box =
    L/2
12 // The uncertainty in the position for the particle
    = 0.181L
```

Chapter 7

Tunnelling phenomena

Scilab code Exa 7.1 Transmission coefficient for an oxide layer

```
1 // Scilab code Ex7.1: Pg 235 (2005)
2 clc; clear;
3 c = 3e+08;      // Velocity of light , m/s
4 m_e = 511e+03/(c^2);    // Mass of electron , eV
5 U = 3.00;       // Ground state energy neglecting E, eV
6 h_cross = (1.973e+03)/c;    // Reduced planck's
                                constant , eV
7 alpha = sqrt(2*m_e*U)/h_cross;
8 L = 50;         // Thickness of the layer , angstrom
9 T = 1/(1+1/4*10^2/(7*3)*sinh(alpha*L)^2);
10 printf("\nThe transmission coefficient for the layer
           thickness of");
11 printf("\n%2d angstrom = %5.3e", L, T);
12 L = 10;         // // Thickness of the layer , angstrom
13 T = 1/(1+1/4*10^2/(7*3)*sinh(alpha*L)^2);
14 printf("\n%2d angstrom = %5.3e", L, T);
15
16 // Result
17 // The transmission coefficient for the layer
           thickness of
18 // 50 angstrom = 9.628e-39
```

```
19 // 10 angstrom = 6.573e-08
```

Scilab code Exa 7.2 Tunnelling current through an oxide layer

```
1 // Scilab code Ex7.2: Pg 236 (2005)
2 clc; clear;
3 e = 1.60e-19;      // Electrc charge, C
4 i = 1.00e-03;      // Electron current, A
5 N = i/e;          // Electrons per second
6 T = 0.657e-07;    // Fraction of electrons
                     transmitted
7 T_e = N*T;        // Number of electrons transmitted per
                     second
8 T_i = T_e*e;      // Transmitted current, A
9 printf("\nThe transmitted current through the oxide
       layer = %4.1f pA", T_i*1e+12);
10
11 // Result
12 // The transmitted current through the oxide layer =
       65.7 pA
```

Scilab code Exa 7.5 Tunnelling in a parallel plate capacitor

```
1 // Scilab code Ex7.5: Pg 241 (2005)
2 clc; clear;
3 epsilon_c = 5.5e+10;    // Characteristic field
                         strength, V/m
4 epsilon = 1.0e+09;     // Electric field, V/m
5 f = 1.0e+30;          // Collision frequency, s(-1)cm(-2)
6 lamda = f*exp(-epsilon_c/epsilon); // Electron
                     emission rate, electrons/sec
7 e = 1.60e-19;         // Electrc charge, C
8 I = lamda*e;          // Tunelling current, A
```

```

9 printf("\nTunelling current in parallel plate
       capacitor = %4.2f pA", I/1e-12);
10 printf("\n");
11
12 // Result
13 // Tunelling current in parallel plate capacitor =
       0.21 pA

```

Scilab code Exa 7.6 Estimating halflives of Thorium and Polonium

```

1 // Scilab code Ex7.6: Pg 244 (2005)
2 clc; clear;
3 Z_T = 88;      // Atomic number of daughter nucleus
4 E_T = 4.05e+06;    // Energy of ejected alphas , eV
5 R = 9.00e-15;    // Nuclear radius , m
6 r_o = 7.25e-15;    // Bohr radius , m
7 E_o = 0.0993e+06;    // Energy analogous to the
                           Rydberg in Atomic Physics
8 T_T = exp(-4*pi*Z_T*sqrt(E_o/E_T) + 8*sqrt((Z_T*R)/
           r_o));    // Transmission factor in case of
                           Thorium
9 f = 1e+21;      // Frequency of collisions , Hz
10 lamda_T = f*T_T;    // Decay rate in case of Thorium
                         , s^(-1)
11 t_T = 0.693/lamda_T;    // Half-life time of
                           Thorium , s
12 Z_P = 82;      // Atomic number of daughter nucleus
13 E_P = 8.95e+06;    // Energy of ejected alphas , eV
14 R = 9.00e-15;    // Nuclear radius , m
15 r_o = 7.25e-15;    // Bohr radius , m
16 E_o = 0.0993e+06;    // Energy unit , eV
17 T_P = exp(-4*pi*Z_P*sqrt(E_o/E_P) + 8*sqrt((Z_P*R)/
           r_o));    // Transmission factor in case of
                           Polonium
18 f = 1e+21;      // Frequency of collisions , Hz

```

```
19 lamda_P = f*T_P;           // Decay rate in case of
    Thorium , s^(-1)
20 t_P = 0.693/lamda_P;      // Half-life time of
    Polonium , s
21
22 printf("\nHalf-life time of Thorium = %3.1e s = %3.1
    e yrs", t_T, t_T/(365*24*60*60));
23 printf("\nHalf-life time of Polonium = %3.1e s", t_P
    );
24
25 // Result
26 // Half-life time of Thorium = 5.3e+17 s = 1.7e+10
    yrs
27 // Half-life time of Polonium = 8.4e-10 s
```

Chapter 8

Quantum Mechanics in Three Dimensions

Scilab code Exa 8.4 Orbital quantum number for a stone

```
1 // Scilab code Ex8.4: Pg 270 (2005)
2 clc; clear;
3 R = 1.00;      // Radius of circle , m
4 T = 1.00;      // Time period of revolution , s
5 v = (2*pi*R)/T;    // Speed of stone in its orbit ,
m/s
6 m = 1.00;      // Mass of stone , kg
7 L = m*v*R;    // Angular momentum of stone , kg-m^2/s
8 h_cross = 1.055e-34;    // Reduced Planck's constant
, kg-m^2/s
9 l = L/h_cross;   // Orbital quantum number
10 printf("\nThe orbital quantum number for stone = %4.2
fe+34", l*1e-34);
11
12 // Result
13 // Orbital quantum number for stone = 5.96e+34
```

Scilab code Exa 8.6 Space quantisation for an atomic electron

```
1 // Scilab code Ex8.6: Pg 272 (2005)
2 clc; clear;
3 // For simplicity let h_cross = 1
4 h_cross = 1; // Reduced planck's constant
5 l = 3; // Given orbital quantum number
6 L = sqrt(l*(l+1)*h_cross); // Magnitude of total
    angular momentum, in h_cross units
7 m_l = [-3, -2, -1, 0, 1, 2, 3];
8 L_z = m_l*h_cross; // Allowed values of L_z
9 cos_theta = L_z/L;
10 theta = acosd(L_z/L); // Orientations of L_z ,
    degrees
11 for i = 1:1:7
12     if theta(i) > 90 then
13         theta(i) = theta(i)-180;
14     end
15 end
16 printf("\nThe magnitude of total angular momentum =
    2*sqrt(%d)*h_cross\n", L^2/4);
17 printf("\nThe allowed values of L_z in units of
    h_cross are :");
18 disp(L_z);
19 printf("\nThe orientations of L_z in degrees are:");
20 disp(theta);
21
22 // Result
23 // The magnitude of total angular momentum = 2*sqrt
    (2)*h_cross
24
25 // The allowed values of L_z in units of h_cross are
    :
26 // - 3. - 2. - 1. 0. 1. 2. 3.
27
28 // The orientations of L_z in degrees are:
29 // - 30. - 54.73561 - 73.221345 90.
    73.221345 54.73561 30.
```

Scilab code Exa 8.7 Energy of Hydrogen atom at first excited state

```
1 // Scilab code Ex8.7: Pg 281 (2005)
2 clc; clear;
3 k = 9e+09; // Coulomb constant , N/Sq.m/C
4 e = 1.6e-019; // Electronic charge , C
5 a_0 = 0.529e-010; // Bohr's radius , m
6 n = 2; // Principal quantum number
7 l = [0, 1]; // Orbital quantum number
8 m_l = [-1, 0, 1]; // Orbital magnetic quantum
number
9 Z = 1; // Atomic number of hydrogen
10 E2 = -k*e^2/(2*a_0)*Z^2/n^2; // Energy of first
excited level of hydrogen,
11 printf("\nThe energy of first excited level of
hydrogen = %3.1f eV", E2/e);
12
13 // Result
14 // The energy of first excited level of hydrogen =
-3.4 eV
```

Scilab code Exa 8.8 Probabilities for the Electron in Hydrogen

```
1 // Scilab code Ex8.8: Pg 284 (2005)
2 clc; clear;
3 P = 1/2*integrate('z^2*exp(-z)', 'z', 2, 100); // 
Take some large value of upper limit
4 printf("\nP(electron in the ground state of hydrogen
will be found outside the first Bohr radius) =
%4.1f percent", P*100);
5
```

```
6 // Result
7 // P(electron in the ground state of hydrogen will
   be found outside the first Bohr radius) = 67.7
   percent
```

Chapter 9

Atomic Structure

Scilab code Exa 9.1 Magnetic energy of electron in Hydrogen

```
1 // Scilab code Ex9.1: Pg 300 (2005)
2 clc; clear;
3 // Since mu_B = (e*h_cross)/(2*m_e)
4 mu_B = 9.27e-24;      // Bohr magneton , J/T
5 B = 1.00;              // Magnetic flux , T
6 // Since 1 eV = 1.6e-19 J
7 eV = 1.6e-19;          // Energy , J
8 h_cross = 6.58e-16;    // Reduced Plank's constant ,
                           eV-s
9 omega_L = (mu_B*B)/(eV*h_cross);      // Larmor
                                           frequency , rad/s
10 printf("\nLarmour frequency at n = 2 is %4.2fe+10
           rad/s", omega_L*1e-10);
11
12 // Result
13 // Larmour frequency at n = 2 is 8.81e+10 rad/s
```

Scilab code Exa 9.2 Angles between z axis and the spin angular momentum vector

```

1 // Scilab code Ex9.2: Pg 307 (2005)
2 clc; clear;
3 h_cross = 6.58e-16;      // Reduced Plank's constant ,
   eV-s
4 S = h_cross*sqrt(3)/2;    // Spin angular momentum ,
   eV-s
5 S_z = h_cross/2;         // Z-component of spin angular
   momentum , eV-s
6 theta_up = acosd(S_z/S);
7 theta_down = acosd(-S_z/S);
8 printf("\nFor up spin state , theta = %4.2f degrees",
   theta_up);
9 printf("\nFor down spin state , theta = %5.1f degrees",
   theta_down);
10
11 // Result
12 // For up spin state , theta = 54.74 degrees
13 // For down spin state , theta = 125.3 degrees

```

Scilab code Exa 9.3 Zeeman Spectrum of Hydrogen Including Spin

```

1 // Scilab code Ex9.3: Pg 311 (2005)
2 clc; clear;
3 e = 1.6e-019;      // Energy equivalent of 1 eV, J/eV
4 B = 1.00;          // Magnitude of magnetic field , tesla
5 n = 2;             // Initial state of the hydrogen atom
6 mu_B = 9.27e-024; // Bohr's magneton , J/T
7 E_Z = mu_B*B/e;   // Zeeman energy , eV
8 E2 = -13.6/n^2;   // Energy of first excited state
   , eV
9 m_l = [-2, -1, 0, 1, 2]; // Orbital magnetic
   quantum number for l = 2
10 printf("\nThe energies of the electron (in eV) in n
   = 2 state are:\n");
11 for i = 1:1:5

```

```

12     if m_l(i) < 0 then
13         sig = '-';
14     else
15         sig = '+';
16     end
17     printf("(%.4.2f %s %.4.2e)", E2, sig, abs(E_Z*
18     m_l(i)));
19
20 // Result
21 // The energies of the electron (in eV) in n = 2
22 // state are:
23 // (-3.40 - 1.16e-04) (-3.40 - 5.79e-05) (-3.40 +
24 // 0.00e+00) (-3.40 + 5.79e-05) (-3.40 + 1.16e-04)

```

Scilab code Exa 9.4 Spin orbit energy of Sodium doublet

```

1 // Scilab code Ex9.4: Pg 311 (2005)
2 clc; clear;
3 hc = 1240;      // Product of plank's constant &
                  velocity of light , eV
4 lamda_1 = 588.995;    // Wavelength of first doublet
                  of Na lines , nm
5 lamda_2 = 589.592;    // Wavelength of second
                  doublet of Na lines , nm
6 delta_E = hc*(lamda_2 - lamda_1)/(lamda_1*lamda_2);
                  // Spin orbit energy , eV
7 printf("\nSpin orbit energy from doublet spacing =
%4.2fe-03 eV", delta_E*1e+03);
8
9 // Result
10 // Spin orbit energy from doublet spacing = 2.13e-03
eV

```

Scilab code Exa 9.5 Ground state of Helium atom

```
1 // Scilab code Ex9.5: Pg 316 (2005)
2 clc; clear;
3 n = 1;      // Principal quantum number
4 Z = 2;      // Atomic number of Helium
5 E_a = (-13.6*Z^2)/n^2;      // Energy of the
     electron in state 'a', eV
6 E_b = (-13.6*Z^2)/n^2;      // Energy of the
     electron in state 'b', eV
7 E = E_a + E_b;      // Total electronic energy of
     Helium, eV
8 printf("\nTotal electronic energy of Helium = %5.1f
     eV", E);
9
10 // Result
11 // Total electronic energy of Helium = -108.8 eV
```

Scilab code Exa 9.6 Effective atomic number for 3s electron in Na

```
1 // Scilab code Ex9.6: Pg 317 (2005)
2 clc; clear;
3 E_i = 5.14;      // Ionisation energy of Na, eV
4 n = 3;      // Principal quantum number
5 Z_eff = sqrt((n^2*E_i)/13.6);      // Effective atomic
     number
6 printf("\nEffective atomic number for 3s electron in
     Na = %4.2f", Z_eff);
7
8 // Result
9 // Effective atomic number for 3s electron in Na =
     1.84
```


Chapter 10

Statistical Physics

Scilab code Exa 10.1 Population of excited states with respect to ground states in

```
1 // Scilab code Ex10.1: Pg 340 (2005)
2 clc; clear;
3 // Part (a)
4 E1 = -13.6;      // Energy of ground state , eV
5 E2 = -3.40;      // Energy of first excited state , eV
6 E3 = -1.51;      // Energy of second excited state , eV
7 g1 = 2;          // Degeneracy for ground state
8 g2 = 8;          // Degeneracy for first excited state
9 g3 = 18;          // Degeneracy for second excited state
10 kB = 8.617e-05; // Boltzmann constant , eV/K
11 Ta = 300;        // Temperature , K
12 // As n_2/n_1 = (g_2*A*e^(-E_2/(kB*T)))/(g_1*A*e^(-E_1/(kB*T))), on simplifying we get
13 N21 = (g2/g1)*exp((E1 - E2)/(kB*Ta));      // The
           population of first excited state w.r.t ground
           state
14 printf("\nThe population of first excited state w.r.
           t. ground state at %3d K = %ld", Ta, N21);
15
16 // Part (b)
17 Tb = 20000;      // Temperature , K
```

```

18 n21 = (g2/g1)*exp((E1 - E2)/(kB*Tb)); // The
      population of first excited state w.r.t ground
      state
19 n31 = (g3/g1)*exp((E1 - E3)/(kB*Tb)); // The
      population of second excited state w.r.t ground
      state
20 printf("\nThe population of first excited state w.r.
      t. ground state at %4d K = %6.4f", Tb, n21);
21 printf("\nThe population of second excited state w.r.
      t ground state at %4d K = %6.4f", Tb, n31);
22
23 // Part (c)
24 E_strength = (g3/g2)*exp((E2 - E3)/(kB*Tb)); // Emission strength
25 printf("\nEmission strength of spectral lines = %3.2
      f", E_strength);
26
27 // Result
28 // The population of first excited state w.r.t.
      ground state at 300 K = 0
29 // The population of first excited state w.r.t.
      ground state at 20000 K = 0.0108
30 // The population of second excited state w.r.t
      ground state at 20000 K = 0.0081
31 // Emission strength of spectral lines = 0.75

```

Scilab code Exa 10.2 Validity of Maxwell Boltzmann Statistics

```

1 // Scilab code Ex10.2: Pg 345 (2005)
2 clc; clear;
3 // Part (a)
4 N = 6.02e+23; // Number of molecules at STP
5 m = 3.34e-27; // Mass of H-molecule , kg
6 h_cross = 1.055e-34; // Reduced Plank's constant ,
      J-s

```

```

7 V = 22.4e-03;      // Volume occupied by molecules at
                      // STP, m^3
8 T = 273;           // Absolute temperature, K
9 k_B = 13.8e-24;    // Boltzmann constant, J/K
10 x_H = N/V*h_cross^3/(8*(m*k_B*T)^(3/2));   //
                           Particle concentration at STP
11 printf("\nx_H = %4.2e", x_H);
12 if (x_H < 1)
13 printf("\n\nThe criterion for the validity of
          Maxwell Boltzmann Statistics is satisfied in
          hydrogen.");
14
15 // Part (b)
16 d_Ag = 10.5;       // Density of silver, g/m^3
17 M_Ag = 107.9;      // Molar weight of silver, g
18 NV_Ag = (d_Ag/M_Ag)*(6.02e+023)*1e+06;    // Density
          of free electrons in silver, electrons/m^3
19 me = 9.109e-031;   // Mass of an electron, kg
20 T = 300;           // Room temperature, K
21 x_Ag = ((NV_Ag)*h_cross^3)/(8*(me*k_B*T)^(3/2));
                           // Particle concentration at STP
22 printf("\nx_Ag = %4.2f", x_Ag);
23 if (x_Ag > 1)
24 printf("\n\nThe criterion for the validity of
          Maxwell Boltzmann Statistics does not hold for
          electrons in silver");
25
26 // Result
27 // x_H = 8.84e-08
28 // The criterion for the validity of
          Maxwell Boltzmann Statistics is satisfied in
          hydrogen.
29 // x_Ag = 37.13
30 // The criterion for the validity of
          Maxwell Boltzmann Statistics does not hold for
          electrons in silver

```

Scilab code Exa 10.3 Photons in a box

```
1 // Scilab code Ex10.3: Pg 352 (2005)
2 clc; clear;
3 // Part (b)
4 I = integrate('z^2/(exp(z)-1)', 'z', 0, 100); // Integral value
5 k_B = 8.62e-05; // Boltzmann constant, eV/K
6 T = 3000; // Temperature, K
7 h = 4.136e-15; // Plank's constant, eV
8 c = 3e+10; // Velocity of light, cm/s
9 N_V = 8*pi*((k_B*T)/(h*c))^3*I; // Number of photons/cc
10 printf("\nThe density of photons inside the cavity = %4.2fe+11 photons/cc", N_V*1e-11);
11
12 // Result
13 // The density of photons inside the cavity = 5.47e+11 photons/cc
```

Scilab code Exa 10.4 Specific Heat of Diamond

```
1 // Scilab code Ex10.4: Pg 356 (2005)
2 clc; clear;
3
4 // Part (a)
5 k_B = 8.62e-05; // Boltzmann constant, eV/K
6 T_E = 1300; // Temperature, K
7 h_cross = 6.58e-16; // Reduced plank's constant, eV-s
8 omega = (k_B*T_E)/h_cross; // Frequency of vibration of carbon atom in diamond, Hz
```

```

9 spacing = (h_cross*omega); // Spacing between
    adjacent oscillator energy level , eV
10 printf("\nFrequency of vibration of carbon atom in
    diamond = %4.2e Hz", omega);
11 printf("\nSpacing between adjacent oscillator energy
    level = %5.3f eV", spacing);
12
13 // Part (b)
14 T_R = 300; // Room temperature , K
15 p = exp((h_cross*omega)/(k_B*T_R)); // For
    simplification
16 E_R = (h_cross*omega)/(p-1); // Average energy of
    oscillator at room temperature , eV
17 T = 1500; // Temperature , K
18 q = exp((h_cross*omega)/(k_B*T)); // For
    simplification
19 E_bar = (h_cross*omega)/(q-1); // Average energy
    at 1500 K, eV
20 printf("\nAverage energy of oscillator at room
    temperature = %7.5f eV", E_R);
21 printf("\nAverage oscillator energy at %4d K = %7.5f
    eV", T, E_bar);
22
23
24 // Result
25 // Frequency of vibration of carbon atom in diamond
    = 1.70e+14 Hz
26 // Spacing between adjacent oscillator energy level
    = 0.112 eV
27 // Average energy of oscillator at room temperature
    = 0.00149 eV
28 // Average oscillator energy at 1500 K = 0.0813 eV

```

Scilab code Exa 10.5 Fermi Energy of Gold

```

1 // Scilab code Ex10.5: Pg 360 (2005)
2 clc; clear;
3
4 // Part (a)
5 h = 6.625e-34; // Plank's constant , J-s
6 m_e = 9.11e-31; // Mass of electron , kg
7 density = 19.32/(1e-02)^3; // Density of gold , g/m
^3
8 weight = 197; // Molar weight , g/mol
9 N_V = (density/weight)*6.02e+23; // Number of
electrons per mole
10 E_F = (h^2/(2*m_e*1.6e-19))*((3*(N_V))/(8*pi))
^(2/3); // Fermi energy of Gold at 0 K
11 printf("\nFermi energy of Gold at 0 K = %4.2f eV",
E_F);
12
13 // Part (b)
14 v_F = sqrt((2*E_F*1.6e-19)/m_e); // Fermi speed of
Gold at 0 K
15 printf("\nFermi speed of Gold at 0 K = %4.2fe+06 m/s
", v_F*1e-06);
16
17 // Part (c)
18 k_B = 8.62e-05; // Boltzmann constant , eV/K
19 T_F = (E_F)/(k_B); // Fermi temperature for Gold
at 0 K, K
20 printf("\nFermi temperature for Gold at 0 K = %5d K"
, T_F);
21
22 // Result
23 // Fermi energy of Gold at 0 K = 5.53 eV
24 // Fermi speed of Gold at 0 K = 1.39fe+06 m/s
25 // Fermi temperature for Gold at 0 K = 64201 K

```

Chapter 11

Molecular Structure

Scilab code Exa 11.1 Rotation of CO molecule

```
1 // Scilab code Ex11.1: Pg 380 (2005)
2 clc; clear;
3 // Part (a)
4 f = 1.15e+11; // Frequency of transitions , Hz
5 omega = 2*(%pi)*f; // Angular frequency of
    absorbed radiations , Hz
6 h_cross = 1.055e-34; // Reduced planks constant , J
    -s
7 // Since E = (h_cross)^2/I_CM = h_cross*omega ,
    solving for I_CM
8 I_CM = h_cross/omega; // Moment of inertia of
    molecule about its center of mass , kg-m^2
9 printf("\nThe moment of inertia of molecule about
    its center of mass = %4.2e kg-m^2" , I_CM);
10
11 // Part (b)
12 m_O = 16; // Mass of oxygen atom , a.m.u
13 m_C = 12; // Mass of carbon atom , a.m.u
14 mu = (m_O * m_C *0.166e-26)/(m_O + m_C); // 
    Reduced mass , kg
15 // Since I_CM = mew*R_o^2, solving for R_o
```

```

16 R_0 = sqrt(I_CM/mu); // Bond length of carbon
monoxide molecule, m
17 printf("\nThe bond length of carbon monoxide
molecule = %5.3f nm", R_0/1e-09);
18
19 // Result
20 // The moment of inertia of molecule about its
center of mass = 1.46e-046 kg-m^2
21 // The bond length of carbon monoxide molecule =
0.113 nm

```

Scilab code Exa 11.2 Variation of CO molecule

```

1 // Scilab code Ex11.2: Pg 383 (2005)
2 clc; clear;
3
4 // Part (a)
5 f = 6.42e+13; // Frequency of absorption, Hz
6 omega = 2*(pi)*f; // Angular frequency of
absorbed radiations, Hz
7 mu = 1.14e-26; // Reduced mass of CO molecule, kg
8 K = mu*(omega^2); // Effective force constant of
CO molecule, N/m
9 printf("\nThe effective force constant of CO
molecule = %4.2e N/m", K);
10
11 // Part (b)
12 h_cross = 1.055e-34; // Reduced Planck's constant,
J-s
13 A = sqrt(h_cross/(mu*omega)); // Amplitude of
vibrations, m
14 printf("\nThe amplitude of vibrations = %7.5f nm", A
/1e-09);
15
16 // Result

```

```
17 // The effective force constant of CO molecule =  
    1.85e+003 N/m  
18 // The amplitude of vibrations = 0.00479 nm
```

Chapter 12

The Solid State

Scilab code Exa 12.1 Classical free electron model

```
1 // Scilab code Ex12.1: Pg 418 (2005)
2 clc; clear;
3 // Part (a)
4 k_B = 1.38e-23;    // Boltzmann constat , J/K
5 m_e = 9.11e-31;    // Mass of electron , kg
6 T = 300;           // Temperature , K
7 N_A = 6.023e+023; // Avogadro 's number
8 v_rms = sqrt((3*k_B*T)/m_e);          // Root mean
   square velocity of electrons , m/s
9 I = 10;            // Electric current , A
10 A = 4e-06;         // Area of cross-section of copper
   wire , m^2
11 J = I/A;           // Current density , A-m^(-2)
12 d = 8.96;          // Density of copper at room
   temperature , g/cc
13 M = 63.5;          // Atomic mass of Cu, g
14 n = d*N_A/M*1e+06; // Number of electrons per
   metre cube
15 e = 1.6e-19;        // Charge on electron , C
16 v_d = J/(n*e);     // Drift velocity , m/s
17 v_d_rms = v_d/v_rms; // Ratio of drift speed to
```

```

    rms speed
18 printf("\nThe ratio of drift speed to rms speed is =
        %3.1e", v_d_rms);
19
20 // Part (b)
21 L = 2.6e-10;
22 tau = L/v_rms; // Average time between two
                  collisions , s
23 printf("\nAverage time between two collisions = %2.2
        e s", tau);
24
25 // Part (c)
26 sigma = (n*e^2*L)/sqrt(3*k_B*T*m_e); // 
                  Conductivity of copper , per ohm-m
27 printf("\nConductivity of copper at room temperature
        = %3.1e per ohm-m", sigma);
28
29
30 // Result
31 // The ratio of drift speed to rms speed is = 1.6e
        -009
32 // Average time between two collisions = 2.23e-015 s
33 // Conductivity of copper at room temperature = 5.3e
        +006 per ohm-m

```

Scilab code Exa 12.2 Conduction in diamond

```

1 // Scilab code Ex12.2: Pg 429 (2005)
2 clc; clear;
3 V = 7; // Energy gap , V
4 L = 5e-08; // Mean free path , m
5 E = V/L; // Electric field , V/m
6 printf("\nThe electric field strength required to
        produce conduction in diamond = %3.1fe+08 V/m", E
        *1e-08);

```

```
7 printf("\n");
8
9 // Result
10 // The electric field strength required to produce
    conduction in diamond = 1.4e+08 V/m
```

Scilab code Exa 12.3 Forward and reverse currents in diode

```
1 // Scilab code Ex12.3: Pg 436 (2005)
2 clc; clear;
3 e_V = 1;      // Energy applied to diode , eV
4 k_B_T = 0.025; // Product of Boltzmann constant
                  and temperature , eV
5 // For simplicity let (q*V)/(k_B*T) = x
6 x = (e_V/(k_B_T));
7 I_f_r = (exp(x)-1)/(exp(-x)-1); // Ratio of
        forward current to reverse current in diode
8 printf("\nThe ratio of forward current to reverse
        current in diode = %3.1fe+17", I_f_r*1e-17);
9
10 // Result
11 // The ratio of forward current to reverse current
    in diode = -2.4e+17
```

Chapter 13

Nuclear Structure

Scilab code Exa 13.1 The Atomic Mass Unit

```
1 // Scilab code Ex13.1: Pg 466 (2005)
2 clc; clear;
3 M = 0.012; // Atomic mass of carbon , kg
4 N_A = 6.02e+023; // Avogadro's number
5 m = M/N_A; // Mass of one Carbon-12 atom , kg
6 // As m = 12*u, twelve mass units , solving for u
7 u = m/12; // The atomic mass unit , kg
8 printf("\nThe atomic mass unit = %4.2e kg", u);
9
10 // Result
11 // The atomic mass unit = 1.66e-27 kg
```

Scilab code Exa 13.2 The Volume and Density of Nucleus

```
1 // Scilab code Ex13.2: Pg 468 (2005)
2 clc; clear;
3 r0 = 1.2e-015; // Nuclear mean radius , m
4 m = 1.67e-027; // Mass of the nucleon , kg
```

```

5 rho_0 = 3*m/(4*pi*r0^3); // Density of the
    nucleus , kg per metre cube
6 printf("\nThe mass of the nucleus = Am approx.");
7 printf("\nThe volume of the nucleus = 4/3*pi*r0^3*A"
    );
8 printf("\nThe density of the nucleus = %3.1e kg per
    metre cube", rho_0);
9
10 // Result
11 // The mass of the nucleus = Am approx.
12 // The volume of the nucleus = 4/3*pi*r0^3*A
13 // The density of the nucleus = 2.3e+17 kg per metre
    cube

```

Scilab code Exa 13.3 Binding energy of the Deuteron

```

1 // Scilab code Ex13.3: Pg 473 (2005)
2 clc; clear;
3 M2 = 2.014102; // Atomic mass of deuteron , u
4 M_H = 1.007825; // Atomic mass of hydrogen , u
5 m_n = 1.008665; // Mass of a neutron , u
6 E_b = (M_H + m_n - M2)*931.494; // Binding
    energy of the deuteron , MeV/u
7 printf("\nThe binding energy of the Deuteron = %5.3f
    MeV", E_b);
8
9 // Result
10 // The binding energy of the Deuteron = 2.224 MeV

```

Scilab code Exa 13.4 Left out sample during radioactive decay

```

1 // Scilab code Ex13.4: Pg 482 (2005)
2 clc; clear;

```

```

3 T = 5730; // Half life of the carbon-14 isotope ,
years
4 N0 = 1000; // Initial number of carbon-14 isotope
5 t = 22920; // Time of decay , years
6 n = t/T; // Total number of half lives
7 N = (1/2)^n*N0; // Sample remains after 22920 years
8 printf("\nNumber of C-14 isotopes remained after %d
years = %d", t, N);
9
10 // Result
11 // Number of C-14 isotopes remained after 22920
years = 62

```

Scilab code Exa 13.5 The Activity of Radium

```

1 // Scilab code Ex13.5: Pg 483 (2005)
2 clc; clear;
3 T_half = 1.6e+03*3.16e+07; // Half life of
radioactive nucleus Ra-226, s
4 lambda = 0.693/T_half; // Decay constant of Ra-226,
per second
5 N0 = 3.0e+016; // Number of radioactive nuclei at t
= 0
6 R0 = lambda*N0; // Activity of sample at t = 0,
decays/s
7 t = 2.0e+003*3.16e+07; // Time during which the
radioactive disintegration takes place , s
8 R = R0*exp(-1*lambda*t); // Decay rate after 2.0e
+003 years , decay/s
9 printf("\nThe decay constant of Ra-226 = %3.1e per
second", lambda);
10 printf("\nThe activity of sample at t = 0 = %4.1f
micro-Ci", R0/(3.7e+010*1e-006));
11 printf("\nThe activity of sample after %3.1e years =
%3.1e decays/s", t, R);

```

```
12
13 // Result
14 // The decay constant of Ra-226 = 1.4e-11 per second
15 // The activity of sample at t = 0 = 11.1 micro-Ci
16 // The activity of sample after 6.3e+10 years = 1.7e
+05 decays/s
```

Scilab code Exa 13.6 The Activity of Carbon

```
1 // Scilab code Ex13.6: Pg 483 (2005)
2 clc; clear;
3 M = 11.0;      // Atomic mass of C-11 isotope , g
4 NA = 6.02e+023; // Avogadro's number
5 m = 3.50e-06;   // Given mass of Cabon-11, g
6
7 // Part (a)
8 N = m/M*NA;      // Number of C-11 atoms in 3.50
                     micro-g of sample
9 printf("\nThe number of C-11 atoms in %4.2f micro-g
       of sample = %4.2e nuclei", m/1e-06, N);
10
11 // Part (b)
12 T_half = 20.4*60;    // Half life of radioactive
                         nucleus C-11, s
13 lambda = 0.693/T_half; // Decay constant of C-11,
                         per second
14 R0 = lambda*N; // Activity of sample at t = 0,
                     decays/s
15 t = 8.00*60*60;    // Time during which the
                     radioactive disintegration takes place, s
16 R = R0*exp(-1*lambda*t); // Decay rate after 2.0e
+003 years , decay/s
17
18 printf("\nThe activity of C-11 sample at t = 0 is %4
       .2e decays/s", R0);
```

```

19 printf("\nThe activity of sample after %4.2f hours =  

        %4.2e decays/s", t/3600, R);  

20  

21 // Result  

22 // The number of C-11 atoms in 3.50 micro-g of  

    sample = 1.92e+17 nuclei  

23 // The activity of C-11 sample at t = 0 is 1.08e+14  

    decays/s  

24 // The activity of sample after 8.00 hours = 8.99e  

    +06 decays/s

```

Scilab code Exa 13.7 The Radiactive Isotope of Iodine

```

1 // Scilab code Ex13.7: Pg 484 (2005)  

2 clc; clear;  

3 R0 = 5; // Activity of I-131 isotope at the time of  

    shipment, mCi  

4 R = 4.2; // Activity of I-131 isotope at the time  

    of receipt by the medical laboratory, mCi  

5 T_half = 8.04; // Half life of radioactive  

    nucleus I-131, days  

6 lambda = 0.693/T_half; // Decay constant of C-11,  

    per second  

7 // As log(R/R0) = -lambda*t, solving for t  

8 t = -1/lambda*log(R/R0); // Time that has elapsed  

    between two measurements, days  

9 printf("\nThe time that has elapsed between two  

    measurements = %4.2f days", t);  

10  

11 // Result  

12 // The time that has elapsed between two  

    measurements = 2.02 days

```

Scilab code Exa 13.8 Energy Liberated during Decay of Radium

```
1 // Scilab code Ex13.8: Pg 486 (2005)
2 clc; clear;
3 M_X = 226.025406; // Atomic mass of Ra-226, u
4 M_Y = 222.017574; // Atomic mass of Rn-222, u
5 M_alpha = 4.002603; // Mass of alpha particle , u
6 Q = (M_X - M_Y - M_alpha)*931.494; // Q-value for
    Radium Decay, MeV/u
7 printf("\nThe Q-value for Radium Decay = %4.2f MeV",
       Q);
8
9 // Result
10 // The Q-value for Radium Decay = 4.87 MeV
```

Scilab code Exa 13.9 Probability of Alpha Decay

```
1 // Scilab code Ex13.9: Pg 487 (2005)
2 clc; clear;
3 Z = 86; // Atomic number of radon
4 A = 222; // Mass number of radon
5 k = 9e+09; // Coulomb constant , N-metre square per
    C-square
6 e = 1.6e-019; // Charge on an electron , C
7 r0 = 7.25e-015; // Bohr radius for alpha particle , m
8 E0 = k*e^2/(2*r0*1e+06*e); // Rydberg energy , MeV
9 R = 1.2e-015*A^(1/3); // Radius of radon nucleus ,
    fm
10 E = 5; // Disintegration energy during alpha decay ,
    MeV
11 T_E = exp(-4*pi*Z*sqrt(E0/E)+8*sqrt(Z*R/r0)); // 
    Decay probability for alpha disintegration
12 printf("\nThe decay probability for alpha
    disintegration at %d MeV energy = %4.2e", E, T_E)
    ;
```

```
13
14 // Result
15 // The decay probability for alpha disintegration at
  5 MeV energy = 1.29e-34
```

Scilab code Exa 13.11 Radioactive Dating

```
1 // Scilab code Ex13.11: Pg 490 (2005)
2 clc; clear;
3 T_half = 5370*3.6e+07; // Half life of C-14, s
4 lambda = 0.693/T_half; // // Decay constant for C
  -14 disintegration , per sec
5 N_C12 = 6.02e+023/12*25; // Number of C-12 nuclei
  in 25.0 g of carbon
6 N0_C14 = 1.3e-012*N_C12; // Number of C-14 nuclei
  in 25.0 g of carbon before decay
7 R0 = N0_C14*3.83e-012*60; // Initial activity of the
  sample , decays/min
8 R = 250; // Present activity of the sample
9 // As R = R0*exp(-lambda*t), solving for t
10 t = -1/lambda*log(R/R0); // Time during which the
  tree dies , s
11 printf("\nThe lifetime of the tree = %3.1e yr", t
  /(365*24*60*60));
12
13 // Result
14 // The lifetime of the tree = 3.6e+03 yr
```

Chapter 14

Nuclear Physics Applications

Scilab code Exa 14.1 Energy released in Fission

```
1 // Scilab code Ex14.1: Pg 513 (2005)
2 clc; clear;
3 // Part (a)
4 u = 931.5;      // Atomic mass unit , Mev
5 M_Li = 7.016003;    // Mass of Lithium , kg
6 M_H = 1.007825;    // Mass of Hydrogen , kg
7 M_He = 4.002603;   // Mass of Helium , kg
8 Q = (M_Li + M_H - 2*M_He)*u;      // Q-value of the
                                          reaction , MeV
9 // Part (b)
10 K_incident = 0.6;     // Kinetic energy of the
                           incident protons , MeV
11 K_products = Q + K_incident;    // Kinetic energy of
                                         the products
12 printf("\nThe Q value of the reaction = %4.1f MeV" ,
        Q);
13 printf("\nThe kinetic energy of the products (two
           alpha particles) = %4.1f MeV" , K_products);
14
15 // Result
16 // The Q value of the reaction = 17.3 MeV
```

```
17 // The kinetic energy of the products (two alpha  
    particles) = 17.9 MeV
```

Scilab code Exa 14.2 Neutron capture by Al

```
1 // Scilab code Ex14.2: Pg 509 (2005)  
2 clc; clear;  
3 roh = 2.7e+06; // Density of Al, g/cm^3  
4 A = 27; // Mass number of Al  
5 n = (6.02e+23*roh)/A; // Number of nuclei/m^3  
6 sigma = 2.0e-31; // Effective area of nucleus  
    normal to motion, m^2  
7 R_0 = 5.0e+12; // Rate of incident particles  
    per unit area, neutrons/cm^2-s  
8 x = 0.30e-03; // Thickness of foil, m  
9 R = (R_0*sigma*n*x) // Number of neutrons captured  
    by foil, neutrons/cm^2-s  
10 printf("\nThe number of neutrons captured by foil =  
    %3.1fe+07 neutrons/Sq.cm-s", R*1e-07);  
11  
12 // Result  
13 // The number of neutrons captured by foil = 1.8e+07  
    neutrons/Sq.cm-s
```

Scilab code Exa 14.4 Energy released in the Fission of U235

```
1 // Scilab code Ex14.4: Pg 513 (2005)  
2 clc; clear;  
3 m = 1; // Mass of Uranium taken, kg  
4 Q = 208; // Disintegration energy per event, MeV  
5 A = 235; // Mass number of Uranium  
6 N = (6.02e+23*m)/A; // Number of nuclei  
7 E = N*Q; // Disintegration energy, MeV
```

```

8 printf("\nThe total energy released if %1d kg of
    Uranium undergoes fission = %4.2fe+26 MeV" , m, E
    *1e-23);
9
10 // Result
11 // The total energy released if 1 kg of Uranium
    undergoes fission = 5.33e+26 MeV

```

Scilab code Exa 14.5 A Rough Mechanism for Fission Process

```

1 // Scilab code Ex14.5: Pg 513 (2005)
2 clc; clear;
3 A_Ba = 141;      // Mass number of Barium
4 A_Kr = 92;       // Mass number of Barium
5 r_0 = 1.2e-15;   // Separation constant , m
6 r_Ba = r_0*A_Ba^(1/3);   // Nuclear radius of Barium
, m
7 r_Kr = r_0*A_Kr^(1/3);   // Nuclear radius of
Krypton , m
8 r = r_Ba + r_Kr;   // Separation between two atoms ,
m
9 Z_1 = 56;      // Atomic number of Barium
10 Z_2 = 36;     // Atomic number of Barium
11 k = 1.440e-09; // Coulomb constant , eV-m
12 U = k*Z_1*Z_2/r           // Coulomb Potential
energy of two charges , MeV
13 printf("\nThe Coulomb potential energy for two
charges = %3d MeV" , U/1e+06);
14 printf("\nThis shows that the fission mechanism is
plausible");
15
16 // Result
17 // The Coulomb potential energy for two charges =
248 MeV
18 // This shows that the fission mechanism is

```

plausible

Scilab code Exa 14.6 The Fusion of Two Deutrons

```
1 // Scilab code Ex14.6: Pg 519 (2005)
2 clc; clear;
3 // Part (a)
4 e = 1.6e-19;      // Charge on electron , C
5 k = 8.99e-09;     // Coulomb constant , N-m^2/C^2
6 r = 1.0e-14;      // Distance between two deuterons , m
7 // We have U = (k*q1*q2)/r, for deuterons q1 = q2 = e
    , therefore we get
8 U = (k*e^2)/r;    // Potential energy of deuterons , J
9 E_C = 1.1e-014;    // The coulomb energy per deuteron
    , J
10 k_B = 1.38e-023;   // Boltzmann constant , J/mol/K
11 T = 2/3*E_C/k_B;  // Effective temperature
    required for deuteron to overcome the potential
    barrier , K
12 printf("\nThe potential energy of two deuterons
        separated by the distance of %1.0de-14 m = %4.2f
        MeV" , r*1e+14 , (U*1e+12)/e);
13 printf("\nThe effective temperature required for
        deuteron to overcome the potential barrier = %3.1e
        K" , T);
14
15 // Result
16 // The potential energy of two deuterons separated by
    the distance of 1e-14 m = 0.14 MeV
17 // The effective temperature required for deuteron to
    overcome the potential barrier = 5.3e+008 K
18 // Result
19 // The potential energy of two deuterons separated by
    the distance of 1e-14 m = 0.14 MeV
```

Scilab code Exa 14.7 Half value thickness

```
1 // Scilab code Ex14.7: Pg 530 (2005)
2 clc; clear;
3 mew = 55e-02;      // Linear absorption coefficient ,
per m
4 // In equation I(x) = I_o*exp(-mew*x) , replacing I(x)
) by I_o/2 & solving for x, we get
5 x = log(2)/mew;    // Half value thickness , m
6 printf("\nThe half value thickness for lead = %4.2fe
-02 cm", x);
7
8 // Result
9 // The half value thickness for lead = %1.26e-02 cm
```

Chapter 15

Elementary Particle

Scilab code Exa 15.2 Checking Baryon Numbers

```
1 // Scilab code Ex15.2: Pg 560 (2005)
2 clc; clear;
3 // Data for Reaction 1
4 R1 = cell(6,2); // Declare a 6X2 cell
5 R1(1,1).entries = 'p';
6 R1(2,1).entries = 'n';
7 R1(3,1).entries = 'p';
8 R1(4,1).entries = 'p';
9 R1(5,1).entries = 'n';
10 R1(6,1).entries = 'p-bar';
11 R1(1,2).entries = 1;
12 R1(2,2).entries = 1;
13 R1(3,2).entries = 1;
14 R1(4,2).entries = 1;
15 R1(5,2).entries = 1;
16 R1(6,2).entries = -1;
17 // Data for reaction 2
18 R2 = cell(5,2); // Declare a 5X2 cell
19 R2(1,1).entries = 'p';
20 R2(2,1).entries = 'n';
21 R2(3,1).entries = 'p';
```

```

22 R2(4,1).entries = 'p';
23 R2(5,1).entries = 'p-bar';
24 R2(1,2).entries = 1;
25 R2(2,2).entries = 1;
26 R2(3,2).entries = 1;
27 R2(4,2).entries = 1;
28 R2(5,2).entries = -1;
29 // Check baryon number conservation for first
   reaction
30 if (R1(1,2).entries+R1(2,2).entries == (R1(3,2).
   entries+R1(4,2).entries+R1(5,2).entries+R1(6,2).
   entries) then
31   printf("\nThe reaction %s + %s --> %s + %s + %s
         + %s can occur (B is conserved)", R1(1,1).
         entries, R1(2,1).entries, R1(3,1).entries, R1
         (4,1).entries, R1(5,1).entries, R1(6,1).
         entries);
32 else
33   printf("\nThe reaction %s + %s --> %s + %s + %s
         + %s cannot occur (B is not conserved)", R1
         (1,1).entries, R1(2,1).entries, R1(3,1).
         entries, R1(4,1).entries, R1(5,1).entries, R1
         (6,1).entries);
34 end
35 // Check baryon number conservation for second
   reaction
36 if R2(1,2).entries+R2(2,2).entries == R2(3,2).
   entries+R2(4,2).entries+R2(5,2).entries then
37   printf("\nThe reaction %s + %s --> %s + %s + %s
         can occur (B is conserved)", R2(1,1).entries,
         R2(2,1).entries, R2(3,1).entries, R2(4,1).
         entries, R2(5,1).entries);
38 else
39   printf("\nThe reaction %s + %s --> %s + %s + %s
         cannot occur (B is not conserved)", R2(1,1).
         entries, R2(2,1).entries, R2(3,1).entries, R2
         (4,1).entries, R2(5,1).entries);
40 end

```

```
41
42 // Result
43 // The reaction p + n --> p + p + n + p_bar can
   occur (B is conserved)
44 // The reaction p + n --> p + p + p_bar cannot occur
   (B is not conserved)
```

Scilab code Exa 15.3 Checking Lepton Numbers

```
1 // Scilab code Ex15.3: Pg 561 (2005)
2 clc; clear;
3 // Data for Reaction 1
4 R1 = cell(4,3); // Declare a 4X3 cell
5 R1(1,1).entries = 'mu';
6 R1(2,1).entries = 'e-';
7 R1(3,1).entries = 'nue_bar';
8 R1(4,1).entries = 'nu_mu';
9 R1(1,2).entries = 1; // Muon number for mu
10 R1(2,2).entries = 0; // Muon number for e-
11 R1(3,2).entries = 0; // Muon number for nue_bar
12 R1(4,2).entries = 1; // Muon number for nu_mu
13 R1(1,3).entries = 0; // Lepton number for mu
14 R1(2,3).entries = 1; // Lepton number for e-
15 R1(3,3).entries = -1; // Lepton number for
   nue_bar
16 R1(4,3).entries = 0; // Lepton number for nu_mu
17 // Data for Reaction 2
18 R2 = cell(4,3); // Declare a 4X3 cell
19 R2(1,1).entries = 'Pi+';
20 R2(2,1).entries = 'mu+';
21 R2(3,1).entries = 'nu_mu';
22 R2(4,1).entries = 'nu_e';
23 R2(1,2).entries = 0; // Muon number for Pi+
24 R2(2,2).entries = -1; // Muon number for mu+
25 R2(3,2).entries = 1; // Muon number for nu_mu
```

```

26 R2(4,2).entries = 0;           // Muon number for nu_e
27 R2(1,3).entries = 0;           // Lepton number for Pi+
28 R2(2,3).entries = 0;           // Lepton number for mu+
29 R2(3,3).entries = 0;           // Lepton number for nu_mu
30 R2(4,3).entries = 1;           // Lepton number for nu_e
31 // Check lepton number conservation for first
   reaction
32 if (R1(1,2).entries== R1(2,2).entries+R1(3,2).
   entries+R1(4,2).entries) & (R1(1,3).entries == R1
   (2,3).entries+R1(3,3).entries+R1(4,3).entries)
   then
33     printf("\nThe reaction %s --> %s + %s + %s can
          occur (Both L_mu and L_e are conserved)", R1
          (1,1).entries, R1(2,1).entries, R1(3,1).
          entries, R1(4,1).entries);
34 else
35     printf("\nThe reaction %s + %s --> %s + %s + %s
          + %s cannot occur (L_mu and L_e are not
          conserved)", R1(1,1).entries, R1(2,1).entries
          , R1(3,1).entries, R1(4,1).entries);
36 end
37 // Check lepton number conservation for second
   reaction
38 if (R2(1,2).entries== R2(2,2).entries+R2(3,2).
   entries+R2(4,2).entries) & (R2(1,3).entries == R2
   (2,3).entries+R2(3,3).entries+R2(4,3).entries)
   then
39     printf("\nThe reaction %s --> %s + %s + %s can
          occur (Both L_mu and L_e are conserved)", R2
          (1,1).entries, R2(2,1).entries, R2(3,1).
          entries, R2(4,1).entries);
40 else
41     printf("\nThe reaction %s --> %s + %s + %s
          cannot occur (L_mu is conserved but L_e is
          not conserved)", R2(1,1).entries, R2(2,1).
          entries, R2(3,1).entries, R2(4,1).entries);
42 end
43

```

```

44 // Result
45 // The reaction mu --> e- + nue_bar + nu_mu can
   occur (Both L_mu and L_e are conserved)
46 // The reaction Pi+ --> mu+ + nu_mu + nu_e cannot
   occur (L_mu is conserved but L_e is not
   conserved0

```

Scilab code Exa 15.4 Conservation of strangeness

```

1 // Scilab code Ex15.4: Pg 563 (2005)
2 clc; clear;
3 // Data for Reaction 1
4 R1 = cell(4,2); // Declare a 4X2 cell
5 R1(1,1).entries = 'Pi0';
6 R1(2,1).entries = 'n';
7 R1(3,1).entries = 'K+';
8 R1(4,1).entries = 'sigma-';
9 R1(1,2).entries = 0; // Strangeness number for
   Pi0
10 R1(2,2).entries = 0; // Strangeness number for n
11 R1(3,2).entries = 1; // Strangeness number for K+
12 R1(4,2).entries = -1; // Strangeness number for
   sigma-
13 // Data for Reaction 2
14 R2 = cell(4,2); // Declare a 4X2 cell
15 R2(1,1).entries = 'Pi-';
16 R2(2,1).entries = 'p';
17 R2(3,1).entries = 'Pi-';
18 R2(4,1).entries = 'sigma+';
19 R2(1,2).entries = 0; // Strangeness number for Pi
   -
20 R2(2,2).entries = -1; // Strangeness number for p
21 R2(3,2).entries = 1; // Strangeness number for pi
   -
22 R2(4,2).entries = 0; // Strangeness number for

```

```

        sigma+
23 // Check strangeness number conservation for first
   reaction
24 if R1(1,2).entries + R1(2,2).entries == R1(3,2).
   entries+R1(4,2).entries then
25   printf("\nThe reaction %s + %s --> %s + %s can
          occur (Strangeness is conserved)", R1(1,1).
          entries, R1(2,1).entries, R1(3,1).entries, R1
          (4,1).entries);
26 else
27   printf("\nThe reaction %s + %s --> %s + %s
          cannot occur (Strangeness is not conserved)", R1(1,1).
          entries, R1(2,1).entries, R1(3,1).
          entries, R1(4,1).entries);
28 end
29 // Check strangeness number conservation for second
   reaction
30 if R2(1,2).entries + R2(2,2).entries == R2(3,2).
   entries+R2(4,2).entries then
31   printf("\nThe reaction %s + %s --> %s + %s can
          occur (Strangeness is conserved)", R2(1,1).
          entries, R2(2,1).entries, R2(3,1).entries, R2
          (4,1).entries);
32 else
33   printf("\nThe reaction %s + %s --> %s + %s
          cannot occur (Strangeness is not conserved)", R2(1,1).
          entries, R2(2,1).entries, R2(3,1).
          entries, R2(4,1).entries);
34 end
35
36 // Result
37 // The reaction Pi0 + n --> K+ + sigma- can occur (
   Strangeness is conserved)
38 // The reaction Pi- + p --> Pi- + sigma+ cannot
   occur (Strangeness is not conserved)

```

Scilab code Exa 15.5 Making virtual particle real

```
1 // Scilab code Ex15.5: Pg 570 (2005)
2 clc; clear;
3 m_pi = 135;      // Mass of pion , MeV/c^2
4 m_p = 938.3;    // Mass op proton , MeV/c^2
5 // For simplification , let velocity of light be
   unity
6 c = 1;          // Velocity of light , m/s
7 // Simplifying K_th = (m_3 + m_4 + m_5 + ....)^2*c
   ^2 - (m_1 + m_2)^2*c^2, we get
8 K_th = 2*m_pi*c^(2) + ((m_pi*c)^2/(2*m_p));      //
   Required kinetic energy of proton , MeV
9 printf("\nRequired kinetic energy of proton = %3d
   MeV" , ceil(K_th));
10
11 // Result
12 // Required kinetic energy of proton = 280 MeV
```

Chapter 16

Cosmology

Scilab code Exa 16.1 Hubbles law

```
1 // Scilab code Ex16.1: Pg 15 (2005)
2 clc; clear;
3 c = 3e+05;      // Velocity of light , km/s
4 v = c/4;        // Recessional velocity , km/s
5 H_0 = 20e-06;   // Hubble's constant , km/s/
                  lightyear
6 // From Hubble's law , v = H_0*R_max, solving for
   R_max
7 R_max = v/H_0;  // Maximum distance at which Hubble'
                  s law applies without relativistic correction ,
                  lightyears
8 printf("\nThe maximum distance at which Hubbles law
         applies without relativistic correction = %1.0 e
         ly" , R_max);
9 printf("\n");
10
11 // Result
12 // The maximum distance at which Hubbles law applies
   without relativistic correction = 4e+09 ly
```

Scilab code Exa 16.2 Critical density of universe

```
1 // Scilab code Ex16.2: Pg 22 (2005)
2 clc; clear;
3 H = 23e-03/(9.46e15);      // Hubble's constant , km/s/
ly
4 G = 6.67e-11;           // Gravitational constant , N-m^2/
kg^2
5 // Since H^2 = (8*%pi*G*p_c)/3, solving for p_c
6 p_c = (3*H^2)/(8*%pi*G);    // Critical mass density
of universe , kg/m^3
7 printf("\nCritical mass density of universe = %4.2e
kg per metre cube", p_c);
8
9
10 // Result
11 // Critical mass density of universe = %1.06e-27 kg/
m^3
```
