

Scilab Textbook Companion for  
Engineering Physics - I  
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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

## Ultrasonics

Scilab code Exa 1.a.1 calculating fundamental frequency

```
1 // Chapter 1 addl_Example 1
2 //


---


3 clc;
4 clear;
5
6 //input data
7
8 P           = 1;           // for fundamental mode
9 t           = 1.5*10^-3;    // thickness of quartz
10 crystal
11 E           = 7.9*10^10;   // young's modulus in N/
12 m^2
13 p           = 2650;        // density in kg/m^3
14
15 //Calculations
16 f           = (P/(2*t))*sqrt(E/p);      // frequency
of the oscillator circuit
```

```
17 //Output
18 mprintf('The Fundamental Frequency of the Quartz
           crystal = %3.4f MHz', f/10^6);
19
20 //
```

---

### Scilab code Exa 1.a.2 Finding Thickness

```
1 // Chapter 1 addl_Example 2
2 //
3
4
5
6 //input data
7
8 v      = 5000;          // velocity of ultrasonics
   in m/s
9 df     = 60*10^3;        // difference b/w two
   adjacent harmonic freq. in Hz
10
11 //Calculations
12
13 d      = v/(2*df) ;    // thickness of steel
   plate
14
15 //Output
16 mprintf('The thickness of steel plate = %f m',d);
17
18 //
```

---

---

### Scilab code Exa 1.a.3 Finding depth of submerged submarine

```
1 // Chapter 1 addl_Example 3
2 //
3 clc;
4 clear;
5
6 //input data
7
8 v      = 1440;           // velocity of ultrasonics
9 t      = 0.33;           // time taken b/w tx and rx
10     in sec
11 //Calculations
12
13 d      = v*t;           // distance travelled by
14 D      = d/2;           // depth of submerged
15     submarine in m
16 //output
17 mprintf('Depth of submerged submarine = %3.1f m',D);
18
19 //
```

---

---

### Scilab code Exa 1.a.4 Finding velocity of ultrasonics

```

1 // Chapter 1 addl_Example 4
2 //


---


3 clc;
4 clear;
5
6 //input data
7
8 d = 0.55*10^-3; // distance b/w two
      antinodes
9 f = 1.5*10^6; // freq of the
      crystal
10
11 //Calculations
12
13 lamda = 2*d; // wavelength
14 v = f*lamda; // velocity of
      ultronics
15
16 //Output
17 fprintf('Velocity of waves in sea water = %3.0f m/s',
      ,v);
18
19 //


---



```

**Scilab code Exa 1.a.5 Finding Youngs modulus and thickness of crystal**

```

1 // Chapter 1 addl_Example 5
2 //


---


3 clc;

```

```

4 clear;
5
6 //input data
7
8 P = 1; // for fundamental mode
9 p = 2660 // density of quartz in
    kg/m^3
10 f = 1300*10^3 // freq of quartz plate
    for sub division ii
11 k = 2.87*10^3
12 //f1 = (k)/t // freq for sub division i
13
14 // Calculations
15
16 //f = (P/(2*t))*sqrt(E/p);
17 E = p*4*(k)^2; //Youngs modulus in
    N/m^2
18 t = (P/(2*f))*sqrt(E/p);
19
20
21 //Output
22 fprintf('Youngs modulus of quartz plate = %e Nm^-2\n'
    'Thickness of the crystal = %e m',E,t);
23
24 //

```

---

### Scilab code Exa 1.1 Calculating frequency of the Oscillator

```

1 // Chapter 1 Example 1
2 //

```

---

```

3 clc;

```

```

4 clear;
5
6 //input data
7
8 P = 1; // for fundamental mode
9 t = 0.1*10^-2; // thickness of piezo
10 electric crystal
11 E = 80*10^9 // young's modulus
12 p = 2654 // density in kg/m^3
13
14 //Calculations
15 f = (P/(2*t))*sqrt(E/p); // frequency
   of the oscillator circuit
16
17 //Output
18 mprintf('The Frequency of the oscillator circuit =
   %e Hz',f);
19
20 //

```

---

### Scilab code Exa 1.2 Calculating Frequency of the vibrating crystal

```

1 // Chapter 1 Example 2
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7
8 P = 1; // for fundamental mode

```

```

9 t = 0.1*10^-2; // thickness of piezo
10 E = 7.9*10^10 // young's modulus
11 p = 2650 // density in kg/m^3
12
13 // Calculations
14
15 f = (P/(2*t))*sqrt(E/p); // frequency
   of the oscillator circuit
16
17 //Output
18 mprintf('The Frequency of the vibrating crystal = %3
   .3f MHz', f/10^6);
19
20 //

```

---

### Scilab code Exa 1.3 Finding velocity of ultrasonic wave

```

1 // Chapter 1 Example 3
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7
8 f = 1.5*10^6; //frequency of
   ultrasonics in Hz
9 d6 = 2.75*10^-3; // distance between 6
   consecutive nodes
10
11 // Calculations

```

```
12 d      = d6/5;           // distance b/w two
   nodes
13 lamda  = 2*d;           // wavelength in m
14 v      = f*lamda;        // velocity of
   ultrasonics
15
16 //Output
17 mprintf('Velocity of ultrasonics = %3.0f m/sec',v);
18
19 //
```

---

# Chapter 2

## Laser

Scilab code Exa 2.a.1 Finding Energy of first Excited State

```
1 // Chapter 2 addl_Example 1
2 //
=====
3 clc;
4 clear;
5
6 //input data
7 h           = 6.625*10^-34;           // planck 's
    constant
8 c           = 3*10^8;                 // vel. of light
    in m/s
9 lamda       = 5890*10^-10;          // wavelength of
    light in m
10 q          = 1.6*10^-19;           // charge of
    electron
11
12
13 // Calculations
14 Eg         = (h*c)/lamda;          // energy in
    joules
```

```

15 E = Eg/q // energy in eV
16
17 //Output
18
19 mprintf('Energy of the first excited state = %3.3f
eV',E);
20
21 //
=====
```

---

### Scilab code Exa 2.a.2 Finding ratio of stimulated emission to spontaneous emission

```

1 // Chapter 2 addl_Example 2
2 //
=====

3 clc;
4 clear;
5
6 //input data
7 h = 6.625*10^-34; // planck's
constant
8 c = 3*10^8; // vel. of light
in m/s
9 lamda = 5890*10^-10; // wavelength of
light in m
10 k = 1.38*10^-23; // Boltzmann
constant
11 Tc = 280 // Temperature
in centigrades
12
13 // Calculations
14 T = Tc+273; // temperature
in kelvin
```

```

15 R = 1/((exp((h*c)/(k*T*lamda))) - 1);
    // ratio of stimulated emission to spontaneous
    emission
16
17 //Output
18
19 mprintf('The ratio between the stimulated emission
    and apontaneoue emission = %3.3e',R);
20
21 //

```

---

### Scilab code Exa 2.a.3 Finding Number of photons emitted

```

1 // Chapter 2 addl_Example 3
2 //


```

---

```

3 clc;
4 clear;
5
6 //input data
7 h = 6.625*10^-34;           // planck 's
    constant
8 c = 3*10^8;                // vel. of light
    in m/s
9 lamda = 6328*10^-10;        // wavelength of
    He-Ne laser source in m
10 q = 1.6*10^-19;             // charge of
    electron
11 P = 3*10^-3                // output power
    of the He-Ne source in watts or J/sec
12
13

```

```

14 // Calculations
15 v = c/lamda // frequency of
   the photon emitted by the laser beam
16 E = h*v; // energy of a
   photon in joules
17 Po = P*60; // conversion
   fro J/sec to J/min
18 N = Po/E; // No of photons
   emitted per minute
19
20 //Output
21
22 mprintf('The No. of Photons emitted per minute = %3
   .3e photons/minute',N);
23
24 //

```

---

#### Scilab code Exa 2.a.4 Finding no of photons emitted

```

1 // Chapter 2 addl_Example 4
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 h = 6.625*10^-34; // planck 's
   constant
8 c = 3*10^8; // vel. of light
   in m/s
9 lamda = 9.6*10^-6; // wavelength of
   CO2 laser source in m

```

```

10 q           = 1.6*10^-19;           // charge of
   electron
11 P           = 10*10^3                // output power
   of the CO2 laser source in watts or J/sec
12
13
14 //Calculations
15 v           = c/lamda              // frequency of
   the photon emitted by the laser beam
16 E           = h*v;                 // energy of a
   photon in joules
17 Po          = P*60*60;             // conversion
   fro J/sec to J/hour
18 N           = Po/E;                // No of photons
   emitted per hour
19
20 //Output
21
22 mprintf( 'The No. of Photons emitted per hour = %3.3e
   photons/hour ',N);
23
24 //

```

---

### Scilab code Exa 2.a.5 Examining possibility of MASER action

```

1 // Chapter 2 addl_Example 5
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data

```

```

7 h           = 6.625*10^-34;           // planck 's
8 c           = 3*10^8;                 // vel. of light
9 lamda       = 10*10^-2;             // wavelength
10 T          = 300                   // Temperature
11 Kb         = 1.38*10^-23        // Boltzmann
12
13 // Calculations
14 // let R    = Rsp/Rst
15 R          = exp((h*c)/(lamda*Kb*T)) - 1;      // ratio of spontaneous to stimulated emission
16 if R<1 then
17   mprintf('Since the spontaneous emission is
              lesser than stimulated emission \n hence
              MASER action is possible at thermal
              equilibrium ')
18 end
19 //

```

---

### Scilab code Exa 2.a.6 Possibility of Laser action in optical frequencies

```

1 // Chapter 2 addl_Example 6
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data

```

```

7 h           = 6.625*10^-34;           // planck 's
8 c           = 3*10^8;                 // vel. of light
9 lamda       = 5000*10^-10;          // wavelength
10 T          = 300                   // Temperature
11 Kb         = 1.38*10^-23          // Boltzmann
12
13 // Calculations
14 // let R    = Rsp/Rst
15 R          = exp((h*c)/(lamda*Kb*T)) - 1;      //
16 if R<1 then
17     mprintf('Since the spontaneous emission is
18     lesser than stimulated emission \n hence
19     LASER action is possible at thermal
20     equilibrium' )
21 else
22
23 //
```

---

### Scilab code Exa 2.a.7 Finding Energy

```
1 // Chapter 2 Additional Example 7
```

```

2 // _____
3 clc;
4 clear;
5
6 //input data
7 h           = 6.625*10^-34;           // plank 's
    constant
8 c           = 3*10^8;                 // vel. of light
    in m/s
9 lamda       = 5511.11*10^-10;       // wavelength of
    green LED light in m
10 q          = 1.6*10^-19;            // charge of
    electron
11
12 // Calculations
13 Eg          = (h*c)/lamda;         // band gap
    energy in joules
14 E           = Eg/q                // bang gap
    energy in eV
15
16 //Output
17
18 mprintf( 'Energy bandgap Eg     = %3.2f eV' ,E);
19
20 // _____

```

---

### Scilab code Exa 1.1 Calculating Number of electron hole pairs

```

1 // Chapter 2 Example 1
2 //

```

---

```

3  clc;
4  clear;
5
6 //input data
7
8 A           = 4*10^-6;           // Receiving
9 I           = 200;             // Intensity in
10 W/m^2
11 h           = 6.625*10^-34;    // plank 's
12 constant
13 c           = 3*10^8;          // vel. of light
14 in m/s
15 lamda      = 0.4*10^-6;      // wavelength of
16 light in m
17
18 //Calculations
19 v           = c/lamda;        // frequency
20 NOP         = I*A/(h*v)       // number of
21 photons
22
23 //since each photon generates an electron hole pair ,
24 the number of photons is equal to number of
25 electron hole pairs
26
27 //Output
28
29 mprintf('Number of electron hole pairs = %e ', NOP);
30
31 //
```

---



---

### Scilab code Exa 1.2 Calculating Wavelength

```

1 // Chapter 2 Example 1
2 //


---


3 clc;
4 clear;
5
6 //input data
7 Eg = 2.8; // bandgap
8 h = 6.625*10^-34; // plank's
9 c = 3*10^8; // vel. of light
10 q = 1.602*10^-19; // charge of
    electron
11
12 //Calculations
13 E = Eg*q // eV to joules
    conversion
14 lamda = h*c/E; // wavelength
15
16 //Output
17
18 mprintf('wavelength = %3.1f (Blue Colour)',lamda
    *10^10);
19
20 //


---



```

### Scilab code Exa 1.3 Finding Energy Band gap

```

1 // Chapter 2 Example 3
2 //

```

---

```

3  clc;
4  clear;
5
6 //input data
7 h           = 6.625*10^-34;           // plank 's
8 c           = 3*10^8;                 // vel. of light
9 lamda       = 1.55*10^-6;          // wavelength of
10 light in m
11 q           = 1.6*10^-19;          // charge of
12 electron
13
14 //Calculations
15 Eg          = (h*c)/lamda;        // band gap
16 energy in joules
17 E           = Eg/q;               // bang gap
18 energy in eV
19
20 //
```

---

### Scilab code Exa 2.4 Finding number of photons

```

1 // Chapter 2 Example 4
2 //
```

---

```

3 clc;
4 clear;
5
6 //input data
7 h           = 6.625*10^-34;           // plank 's
   constant
8 c           = 3*10^8;                 // vel. of light
   in m/s
9 lamda       = 4961*10^-10;          // wavelength of
   light in m
10
11 //Calculations
12 E           = (h*c)/lamda;          // energy in
   joules
13 N           = 1/E
14 //Output
15
16 mprintf('Number of photons required to do one Joule
   of work = %3.4e /m^3',N);
17
18 //
```

---

### Scilab code Exa 2.5 Calculating Long Wavelength

```

1 // Chapter 2 Example 5
2 //


---


3 clc;
4 clear;
5
6 //input data
7 E           = 0.02;                  // ionisation
```

```

        energy in eV
8 h           = 6.625*10^-34;          // plank 's
        constant
9 c           = 3*10^8;              // vel. of light
        in m/s
10 q          = 1.6*10^-19;         // charge of
        electron

11
12 // Calculations
13
14 lamda       = h*c/(E*q)          // long
        wavelength limit in m
15
16 //Output
17
18 mprintf('long wavelength limit = %3.3e m',lamda);
19
20 //

```

---

### Scilab code Exa 2.6 Finding Wavelength

```

1 // Chapter 2 Example 6
2 //

3 clc;
4 clear;
5
6 //input data
7 E           = 1.44;                  // Bandgap
        energy in eV
8 h           = 6.625*10^-34;         // plank 's
        constant

```

```

9   c           = 3*10^8;           // vel. of light
10  q           = 1.6*10^-19;       // charge of
11                                electron
12 // Calculations
13
14 lamda        = h*c/(E*q)       // Wavelength of
15                                GaAs laser
16 //Output
17
18 mprintf( 'Wavelength of GaAs laser = %3.1f      ', lamda
19                                *10^10 );
20 //
```

---

# Chapter 3

## Fibre Optics And Applications

Scilab code Exa 3.a.1 Finding Refractive index acceptance angle and max no of modes

```
1 // Chapter 3 Additional Example 1
2 //
3 clc;
4 clear;
5
6 //input data
7 n1      = 1.5;          // Refractive index of core
8 NA      = 0.26;         // Numerical aperture
9 d       = 100*10^-6;    // core diameter
10 lamda  = 10^-6;        // wavelength in m
11
12 // Calculations
13 n2      = sqrt( n1^2 - NA^2);           // Refractive
   index of cladding
14 im      = asin(NA);                    // Acceptance
   angle
15 im_d   = im*180/%pi;                  // radian to
   degree conversion
16 N      = 4.9*(d*NA/lamda)^2;          // maximum no of
```

```

modes
17
18 // Output
19 mprintf('Refractive index of cladding n2 = %3.4f\n'
           Acceptance angle = %3.2f degrees\n Maximum number
           of modes that fibre allows = %d ',n2,im_d,N);
20 //


---



```

### Scilab code Exa 3.a.2 Finding Critical angle

```

1 // Chapter 3 Additional Example 2
2 //


---


3 clc;
4 clear;
5
6 //input data
7 delta      = 0.02;          // relative refractive
     index
8 n1         = 1.48;          // refractive index of
     core
9
10 // Calculations
11 NA         = n1*(2*delta)^0.5; // Numerical
     aperture
12 n2         = sqrt( n1^2 - NA^2); // Refractive
     index of cladding
13 cri_ang    = asin(n2/n1);      // critical
     angle
14 cri_ang_d = cri_ang*180/%pi; // critical
     angle in degrees
15

```

```
16 // output
17 mprintf('Numerical Aperture = %3.3f\n The Critical
    angle = %3.2f degrees',NA,cri_ang_d);
18 //
```

---

### Scilab code Exa 3.a.3 Calculating Refractive indices of core and cladding

```
1 // Chapter 3 Additional Example 3
2 //
3 clc;
4 clear;
5
6 //input data
7 delta      = 0.015;           // relative refractive
    index
8 NA         = 0.27;            // Numerical aperture
9
10 // Calculations
11 //we know that NA = n1*sqrt(2* )
12 n1         = NA/sqrt(2*delta) // refractive
    index of core
13 n2         = sqrt( n1^2 - NA^2); // Refractive
    index of cladding
14 // Output
15 mprintf('Refractive index of the core = %3.3f\n
    Refractive index of the cladding = %3.3f\n',n1,n2
    );
16 //
```

---

**Scilab code Exa 3.a.4 Calculating no of modes in a fibre**

```
1 // Chapter 3 Additional Example 4
2 //


---


3 clc;
4 clear;
5
6 //input data
7 NA      = 0.25;          // Numerical aperture
8 d       = 60*10^-6;      // core diameter
9 lamda   = 2.7*10^-6;    // wavelength in m
10
11 // calculations
12 N      = 4.9*(d*NA/lamda)^2;           // no of modes
     for step index fibre
13
14 // Output
15 mprintf('No. of total modes propagating in a
     multimode step index fibre = %d',N);
16 //
```

---

**Scilab code Exa 3.a.5 Calculating No of Modes**

```
1 // Chapter 3 Additional Example 5
2 //


---


3 clc;
```

```

4 clear;
5
6 //input data
7 NA      = 0.25;           // Numerical aperture
8 d       = 6*10^-6;        // core diameter
9 lamda   = 1.5*10^-6;     // wavelength of laser source
10 n1     = 1.47;           // refractive index of core
11 n2     = 1.43;           // refractive index of cladding
12
13 // calculations
14 NA      = sqrt( n1^2 - n2^2);           // Numerical
Aperture
15 N       = 4.9*(d*NA/lamda)^2;           // no of modes
for step index fibre
16
17 // Output
18 mprintf('No. of total modes propagating in the fibre
= %d',N);
19 //

```

---

### Scilab code Exa 3.1 Finding Numerical Aperture

```

1 // Chapter 3 Example 1
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 n1      = 1.6;           // Refractive index of core
8 n2      = 1.5;           // Refractive index of cladding
9

```

```

10 // Calculations
11 NA      = sqrt(n1^2 - n2^2);           // Numerical
    Aperture of optical fiber
12
13 // Output
14 mprintf('Numerical Aperture of the optical fiber =
    %3.4f',NA);
15 //

```

---

### Scilab code Exa 3.2 Finding Numerical Aperture and Acceptance angle

```

1 // Chapter 3 Example 2
2 //


```

---

```

3 clc;
4 clear;
5
6 //input data
7 n1      = 1.55;           // Refractive index of core
8 n2      = 1.5;            // Refractive index of cladding
9
10 // Calculations
11 NA      = sqrt(n1^2 - n2^2);           // Numerical
    Aperture of optical fiber
12 im      = asin(NA);                  // Acceptance
    angle
13 im_d   = im*180/%pi                // radian to
    degree conversion
14
15 // Output
16 mprintf('Numerical Aperture of the optical fiber =
    %3.4f\n Acceptance angle = %3.2f degrees ',NA,

```

```
    im_d);  
17 //
```

---

### Scilab code Exa 3.3 Calculating Refractive Index of Cladding

```
1 // Chapter 3 Example 3  
2 //  
  
3 clc;  
4 clear;  
5  
6 //input data  
7 NA      = 0.26;           // Numerical aperture  
8 n1      = 1.5 ;           // Refractive index of core  
9 d       = 100*10^-6;      // diameter of the core in m  
10  
11 // Calculations  
12 n2      = sqrt( n1^2 - NA^2);          // Refractive  
    index of cladding  
13  
14 // Output  
15 mprintf('Refractive index of cladding = %3.4f',n2);  
16 //
```

---

### Scilab code Exa 3.4 Finding Numerical Aperture

```
1 // Chapter 3 Example 4
```

```
2 //

---

  
3 clc;  
4 clear;  
5  
6 //input data  
7 n1      = 1.54;      // Refractive index of core  
8 n2      = 1.5;       // Refractive index of cladding  
9  
10 // Calculations  
11 NA      = sqrt(n1^2 - n2^2);           // Numerical  
     Aperture of optical fiber  
12  
13 // Output  
14 mprintf('Numerical Aperture of the optical fiber =  
          %3.4f',NA);  
15 //
```

---

# Chapter 4

## Quantum Physics

Scilab code Exa 4.a.1 Finding no of photons emitted

```
1 // Chapter 4 Addtional Example 1
2 //
3 clc;
4 clear;
5
6 // input data
7 h      = 6.625*10^-34          // plancks constant
8 c      = 3*10^8;                // vel. of light
9 lamda = 5893*10^-10;          // wavelength in m
10 P     = 100                   // power of sodium
    vapour lamp
11
12 // Calculations
13 E     = (h*c)/lamda;          // Energy in joules
14 N     = P/E                   // Number of photons
    emitted
15
16 // Output
17 mprintf('Number of Photons emitted = %3.4e per
```

```
18 // second ',N);
```

---

### Scilab code Exa 4.a.2 Finding wavelength of incident beam

```
1 // Chapter 4 AdditionalExample 2
2 //

3 clc;
4 clear;
5
6 // input data
7
8 lamda1 = 0.022*10^-10;           // wavelength of
         scattered X-rays in m
9 theta    = 45;                  // scattering angle in
         degrees
10 h        = 6.625*10^-34;       // plancks constant
11 mo       = 9.11*10^-31;        // mass in Kg
12 c        = 3*10^8;            // vel. of light
13
14 // Calculations
15 // from Compton theory ,Compton shift is given by
16 // lamda' - lamda = (h/(mo*c))*(1- cos )
17
18 theta_r = theta*%pi/180;       // degree to radian
         conversion
19 lamda   = lamda1-( (h/(mo*c))*(1- cos(theta_r))) // 
         incident Wavelength
20
21 // Output
22 mprintf('Wavelength of incident beam = %3.4f ',
```

```
    lamda*10^10);  
23 //
```

---

### Scilab code Exa 4.a.3 Finding Energy of scattered photon

```
1 // Chapter 4 Additional Example 3  
2 //  
  
3 clc;  
4 clear;  
5  
6 // input data  
7 Ei      = 1.02*10^6           // photon energy in eV  
8 theta   = 90;                // scattered angle in  
     degrees  
9 h       = 6.625*10^-34        // plancks constant  
10 mo     = 9.1*10^-31          // mass of electron in Kg  
11 e      = 1.6*10^-19          // charge of electron  
12 c      = 3*10^8;             // vel. of light in m/s  
13  
14 // Calculations  
15 // from Compton theory ,Compton shift is given by  
16 // lamda' - lamda = (h/(mo*c))*(1- cos )  
17 theta_r = theta*%pi/180;     // degree to radian  
     conversion  
18 c_lamda = ( (h/(mo*c))*(1- cos(theta_r))) // Change  
     in wavelength in m  
19 dv      = c/c_lamda;         // change in frequency  
     of the scattered photon  
20 dE      = (h*dv)/e          // change in energy of  
     scattered photon in eV  
21 // This change in energy is transferred as the KE of
```

```

          the recoil electron
22 Er      = dE;                      // Energy of recoil
          electron
23 Es      = Ei - Er                 // Energy of scattered
          photon
24
25
26 // Output
27 mprintf('Energy of the recoil electron = %3.4f MeV\n'
           'Energy of the Scattered photon = %3.4f MeV',Er
           *10^-6,Es*10^-6);
28 //

```

---

#### Scilab code Exa 4.a.4 Finding wavelength of Scattered X rays

```

1 // Chapter 4 Additional Example 4
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7
8 lamda    = 0.124*10^-10;           // wavelength of X-rays
     in   m
9 theta    = 180;                   // Scattering angle in
     degrees
10 h        = 6.625*10^-34          // plancks constant
11 mo       = 9.11*10^-31          // mass in Kg
12 c        = 3*10^8;              // vel. of light
13
14 // Calculatioms

```

```

15 // from Compton theory ,Compton shift is given by
16 // lamda' - lamda = (h/(mo*c))*(1- cos )
17
18 theta_r = theta*%pi/180;           // degree to radian
   conversion
19 lamda1 = lamda+( (h/(mo*c))*(1-cos(theta_r))) // 
   wavelength of scattered X-rays
20
21 // Output
22 mprintf('Wavelength of Scattered X-rays = %3.4f ', 
   lamda1*10^10);
23 //

```

---

### Scilab code Exa 4.a.5 Finding de Broglie wavelength

```

1 // Chapter 4 Additional Example 5
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7 h      = 6.625*10^-34          // plancks constant
8 m      = 9.11*10^-31          // mass of electron in
   Kg
9 e      = 1.6*10^-19           // charge of electron
10 V     = 2000;                 // potential in volts
11
12 // Calculations
13
14 lamda = h/(sqrt(2*m*e*V)) // de Broglie
   wavelength

```

```

15
16 // Output
17 mprintf('The de-Broglie wavelength of electron = %3
18 .4 f      ,lamda*10^10);
19 //

```

---

### Scilab code Exa 4.a.6 Finding de Broglie Wavelength

```

1 // Chapter 4 Additional Example 6
2 //

3 clc;
4 clear;
5
6 // input data
7 h          = 6.625*10^-34           // plancks constant
8 m          = 1.678*10^-27          // mass of proton in Kg
9 e          = 1.6*10^-19            // charge of electron
10 Kb        = 1.38*10^-23;         // boltzmann constant
11 T          = 300                  // Temperature in
12          kelvin
13
14 lamda     = h/(sqrt(3*m*Kb*T)) // de Broglie
15          wavelength
16 // Output
17 mprintf('The de-Broglie wavelength = %3.4 f      ',lamda
18 *10^10);
19 //

```

---

---

### Scilab code Exa 4.a.7 Finding Energy of electron

```
1 // Chapter 4 Additional Example 7
2 //
3 clc;
4 clear;
5 // input data
6 h      = 6.625*10^-34           // plancks constant
7 m      = 9.11*10^-31           // mass of electron in
8 Kg
9 lamda = 3*10^-2;                // wavelength of
10 electron wave
11 e      = 1.6*10^-19;            // charge of electron
12 // Calculations
13
14 E      = (h^2)/(2*m*lamda^2);  // Energy in Joules
15 E1     = E/e;
16 // Output
17 mprintf('Energy of the electron E = %3.4e eV\n',E1);
18 mprintf(' Note: Calculation mistake in textbook')
19 //
```

---

### Scilab code Exa 4.a.8 Proving de Broglie is equal to compton wavelength

```
1 // Chapter 4 Additional Example 8
2 //
```

---

```

3 clc;
4 clear;
5 // input data
6 h = 6.625*10^-34           // plancks constant
7 m = 9.11*10^-31           // mass of electron in
     Kg
8 c = 3*10^8;                // velocity of light in
     m/s
9
10 // Calculations
11 ve = 0.7071*c              // velocity of electron
12 lamda = h/(m*ve*sqrt(1-(ve/c)^2)) // de Broglie
     wavelength
13
14 // we know Compton wavelength ,lamda' - lamda = (h
     /(m*c))*(1- cos )
15 // maximum shift = 180
16 theta = 180
17 theta1 = theta*%pi/180;
18 d_lamda = (h/(m*c))*(1- cos(theta1))
19 mprintf('de Broglie wavelength = %e m\n',lamda);
20 mprintf(' compton wavelength = %e m\n',d_lamda)
21 mprintf(' The de-Broglie wacelength is equal to the
     compton wavelength');
22 //

```

---

### Scilab code Exa 4.a.9 Finding Eigen Values

```

1 // Chapter 4 Additional Example 9
2 //

```

---

```

3 clc;

```

```

4 clear;
5
6 // input data
7 l      = 10^-10;           // side of one
   dimensional box
8 h      = 6.625*10^-34    // plancks constant in
   Jsec
9 m      = 9.11*10^-31    // mass of electron in
   Kg
10 n1     = 1;              // for 1st eigen value
11 n2     = 2;              // for 2nd eigen value
12 n3     = 3;              // for 3rd eigen value
13 n4     = 4;              // for 4th eigen value
14 e      = 1.6*10^-19    // charge of electron
   in columbs
15
16 // Calculations
17 E1     = (h^2 * n1^2)/(8*m*l^2 *e) // first Eigen
   value
18 E2     = (h^2 * n2^2)/(8*m*l^2 *e) // second Eigen
   value
19 E3     = (h^2 * n3^2)/(8*m*l^2 *e) // third Eigen
   value
20 E4     = (h^2 * n4^2)/(8*m*l^2 *e) // fourth Eigen
   value
21
22 // Output
23 fprintf('1st Eigen value = %3.1f eV\n 2nd Eigen
   value = %3.1f eV\n 3rd Eigen value = %3.1f eV\n
   4th Eigen value = %3.1f eV',E1,E2,E3,E4);
24 //

```

---



---

**Scilab code Exa 4.a.10 Finding Energy of system having two electrons**

```

1 // Chapter 4 Additional Example 10
2 //


---


3 clc;
4 clear;
5
6 // input data
7 l      = 10^-10 ;           // length of one
8 h      = 6.625*10^-34     // plancks constant in
9 m      = 9.11*10^-31     // mass of electron in
10 Kg
11 n      = 1;               // for ground state
12 e      = 1.6*10^-19     // charge of electron
13 in columbs
14 // Calculations
15 E      = 2*(h^2 * n^2)/(8*m*l^2 *e) // Energy of
16 system having two electrons
17 // Output
18 mprintf('Energy of the system having two electrons =
19 %3.4f eV',E);
20 //


---



```

### Scilab code Exa 4.a.11 Finding Magnifying power

```

1 // Chapter 4 Additional Example 10
2 //


---


3 clc;

```

```

4 clear;
5
6 // input data
7 b = 40;           // angle subtended by final
                     images at eye in degrees
8 a = 10;           // angle subtended by the object
                     at the eye kept at near point in degrees
9
10 // Calculations
11 b_r = b*%pi/180; // degree to radian
                     conversion
12 a_r = a*%pi/180; // degree to radian
                     conversion
13 M = tan(b_r)/tan(a_r); // magnifying power
14
15 // Output
16 mprintf('Magnifying power = %3.3f',M);
17 //

```

---

### Scilab code Exa 4.1 Finding Wavelength of the Scattered photons

```

1 // Chapter 4 Example 1
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7
8 lamda = 3*10^-10;           // wavelength of
                             incident photons in m
9 theta = 60;                 // viewing angle in

```

```

    degrees
10 h      = 6.625*10^-34      // plancks constant
11 mo     = 9.11*10^-31       // mass in Kg
12 c      = 3*10^8;           // vel. of light
13
14 // Calculations
15 // from Compton theory ,Compton shift is given by
16 // lamda' - lamda = (h/(mo*c))*(1- cos )
17
18 theta_r = theta*pi/180;    // degree to radian
   conversion
19 lamda1 = lamda+( (h/(mo*c))*(1- cos(theta_r))) // 
   wavelength of scattered photons
20
21 // Output
22 mprintf( 'Wavelength of Scattered photons = %3.4f      '
   ,lamda1*10^10);
23 //

```

---

### Scilab code Exa 4.2 Finding Change in Wavelength

```

1 // Chapter 4 Example 2
2 //


```

---

```

3 clc;
4 clear;
5
6 // input data
7 theta    = 135;           // angle in degrees
8 h        = 6.625*10^-34   // plancks constant
9 mo       = 9.1*10^-31     // mass in Kg
10 c       = 3*10^8;         // vel. of light in m/s

```

```

11
12 // Calculations
13 // from Compton theory ,Compton shift is given by
14 // lamda' - lamda = (h/(mo*c))*(1- cos )
15 theta_r = theta*pi/180;           // degree to radian
        conversion
16 c_lamda = ( (h/(mo*c))*(1- cos(theta_r))) // Change
        in wavelength in m
17
18 // Output
19 mprintf('Change in Wavelength = %3.5f ',c_lamda
        *10^10);
20 //


---



```

### Scilab code Exa 4.3 Finding wavelength of Scattered beam

```

1 // Chapter 4 Example 3
2 //


---


3 clc;
4 clear;
5
6 // input data
7
8 lamda    = 0.1*10^-9;           // wavelength of X-rays
        in m
9 theta    = 90;                 // angle with incident
        beam in degrees
10 h        = 6.625*10^-34;      // plancks constant
11 mo       = 9.11*10^-31;      // mass in Kg
12 c        = 3*10^8;            // vel. of light
13

```

```

14 // Calculations
15 // from Compton theory ,Compton shift is given by
16 // lamda' - lamda = (h/(mo*c))*(1- cos )
17 theta_r = theta*%pi/180;           // degree to radian
    conversion
18 lamda1 = lamda+( (h/(mo*c))*(1-cos(theta_r))) // 
    wavelength of scattered beam
19
20 // Output
21 mprintf('Wavelength of Scattered beam = %3.4f ', 
    lamda1*10^10);
22 //


---



```

#### Scilab code Exa 4.4 Finding De Broglie wavelength

```

1 // Chapter 4 Example 4
2 //


---


3 clc;
4 clear;
5
6 // input data
7 h      = 6.625*10^-34          // plancks constant
8 m      = 9.11*10^-31          // mass of electron in
    Kg
9 e      = 1.6*10^-19           // charge of electron
10 V     = 150;                  // potential difference
    in volts
11
12 // Calculations
13
14 lamda = h/(sqrt(2*m*e*V)) // de Broglie

```

```

        wavelength
15
16 // Output
17 mprintf('The de-Broglie wavelength = %d      ', lamda
           *10^10);
18 //

```

---

### Scilab code Exa 4.5 Finding de Broglie wavelength

```

1 // Chapter 4 Example 5
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7 h      = 6.625*10^-34          // plancks constant
8 m      = 9.11*10^-31          // mass of electron in
                                Kg
9 e      = 1.6*10^-19           // charge of electron
10 V     = 5000;                 // potential in volts
11
12 // Calculations
13
14 lamda = h/(sqrt(2*m*e*V))   // de Broglie
                                wavelength
15
16 // Output
17 mprintf('The de-Broglie wavelength of electron = %3
           .5 f      ', lamda*10^10);
18 //

```

---

---

### Scilab code Exa 4.6 Finding de Broglie wavelength

```
1 // Chapter 4 Example 6
2 //
3 clc;
4 clear;
5
6 // input data
7 E      = 100           // Energy of electron
8 in eV
9 h      = 6.625*10^-34 // plancks constant
10 m     = 9.11*10^-31  // mass of electron in
11 Kg
12 e      = 1.6*10^-19   // Charge of electron
13 in Columbs
14
15 // Calculations
16
17 E1      = E*e          // Energy conversion
18 from eV to Joule
19 lamda   = h/(sqrt(2*m*E1)) // de Broglie wavelength
20
21 // Output
22 mprintf('The de-Broglie wavelength = %3.3f ,'
23 lamda*10^10);
24 //
```

---

### Scilab code Exa 4.7 Finding de Broglie Wavelength

```
1 // Chapter 4 Example 7
2 //


---


3 clc;
4 clear;
5
6 // input data
7 m      = 1.675*10^-27;           // Mass of proton in kg
8 c      = 3*10^8;                 // velocity of light in
9      m/s
10 h      = 6.625*10^-34;          // plancks constant
11
12 // Calculations
13 vp     = c/20;                  // velocity of proton in
14      m/s
15 lamda  = h/(m*vp);             // de-Broglie wavelength
16      in m
17
18 // Output
19 mprintf('de-Broglie wavelength = %e m',lamda);
20 //
```

---

### Scilab code Exa 4.8 Finding de Broglie Wavelength of neutron

```
1 // Chapter 4 Example 8
2 //


---


3 clc;
```

```

4 clear;
5
6 // input data
7 E      = 10000           // Energy of neutron in
   eV
8 h      = 6.625*10^-34    // plancks constant
9 m      = 1.675*10^-27    // mass of neutron in
   Kg
10 e     = 1.6*10^-19
11 // Calculations
12
13 E1     = E*e           // Energy conversion
   from eV to Joule
14 lamda  = h/(sqrt(2*m*E1)) // de Broglie wavelength
15
16 // Output
17 mprintf('The de-Broglie wavelength of neutron = %3.3
   e m', lamda);
18 //

```

---

### Scilab code Exa 4.10 Finding Energy level and Temperature of molecules

```

1 // Chapter 4 Example 10
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7 l      = 0.1*10^-9;        // side of cubical box
8 h      = 6.625*10^-34     // plancks constant in
   Jsec

```

```

9 m      = 9.11*10^-31           // mass of electron in
10 Kg
11 Kb      = 1.38*10^-23          // Boltzmann constant
12 // Calculations
13 // for cubical box the energy eigen value is Enx ny
14 nz = (h^2/(8*m*l^2))*(nx^2 + ny^2 +nz^2)
15 // For the next energy level to the lowest energy
16 level nx = 1 , ny = 1 and nz = 2
17 nx      = 1
18 ny      = 1
19 nz      = 2
20 E112    = (h^2/(8*m*l^2))*( nx^2 + ny^2 + nz^2);
21
22 // we know the average energy of molecules of
23 aperfect gas = (3/2)*(Kb*T)
24 T      = (2*E112)/(3*Kb);       // Temperature in
25 kelvin
26 // Output
27 mprintf('E112 = %3.4e Joules\n Temperature of the
28 molecules T = %3.4e K',E112,T);
29 //

```

---

### Scilab code Exa 4.11 Finding Minimum energy of an electron

```

1 // Chapter 4 Example 11
2 //

```

---

```

3 clc;
4 clear;
5

```

```

6 // input data
7 l = 4*10^-9; // width of infinitely
    deep potential
8 h = 6.625*10^-34 // plancks constant in
    Jsec
9 m = 9.11*10^-31 // mass of electron in
    Kg
10 n = 1; // minimum energy
11 e = 1.6*10^-19 // charge of electron
    in columbs
12
13 // Calculations
14 E = (h^2 * n^2)/(8*m*l^2) // Energy of
    electron in an infinitely deep potential well
15 E1 = E/e // energy
    conversion from joules to eV
16
17 // Output
18 mprintf('Minimum energy of an electron = %3.4f eV', E1);
19 //

```

---

### Scilab code Exa 4.12 Finding Energy for Exciting a electron

```

1 // Chapter 4 Example 12
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7 l = 0.1*10^-9; // length of one

```

```

        dimensional box
8 h      = 6.625*10^-34           // plancks constant in
     Jsec
9 m      = 9.11*10^-31           // mass of electron in
     Kg
10 n     = 1;                   // for ground state
11 n5    = 6;                   // n value for fifth
     excited state
12 e      = 1.6*10^-19          // charge of electron
     in columbs
13
14 // Calculations
15 Eg     = (h^2 * n^2)/(8*m*l^2 *e) // Energy in
     ground state in eV
16 Ee     = (h^2 * n5^2)/(8*m*l^2 * e) // Energy in
     excited state in eV
17 E      = Ee - Eg;             // energy req
     to excite electrons from ground state to fift
     excited state
18
19 // Output
20 mprintf('Energy required to excite an electron from
     ground state to fifth excited state = %3.2f eV',E
     );
21 //

```

---

### Scilab code Exa 4.13 Finding Energy of electron

```

1 // Chapter 4 Example 13
2 //

```

---

```

3 clc;

```

```

4 clear;
5
6 // input data
7 l      = 0.1*10^-9;           // length of one
     dimensional box
8 h      = 6.625*10^-34        // plancks constant in
     Jsec
9 m      = 9.11*10^-31         // mass of electron in
     Kg
10 n     = 1;                  // for ground state
11 e     = 1.6*10^-19          // charge of electron
     in columbs
12
13 // Calculations
14 E      = (h^2 * n^2)/(8*m*l^2 *e) // Energy of
     electron in eV
15 // Output
16 mprintf('Energy of an electron = %3.3f eV',E);
17 //

```

---

### Scilab code Exa 4.14 Finding Least energy of electron

```

1 // Chapter 4 Example 14
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7 l      = 0.5*10^-9;           // width of one
     dimensional box in m
8 h      = 6.625*10^-34          // plancks constant in

```

```

Jsec
9 m      = 9.11*10^-31           // mass of electron in
Kg
10 n      = 1;                  // for ground state
11 e      = 1.6*10^-19          // charge of electron
     in columbs
12
13 // Calculations
14 E      = (h^2 * n^2)/(8*m*l^2 *e) // Energy of
     electron in eV
15 // Output
16 mprintf('Least Energy of an electron = %3.4f eV',E);
17 //

```

---

# Chapter 5

## Crystal Physics

Scilab code Exa 5.a.1 Finding atomic radius

```
1 // Chapter 5 additional Example 1
2 //
=====
3 clc;
4 clear;
5
6 // input data
7 // Copper has FCC structure
8 a = 3.6;           // lattice parameter of copper in
9
10 // Calculations
11
12 r = a*sqrt(2)/4;      // atomic radius of copper
13
14 // Output
15 mprintf('Atomic Radius of copper = %3.3f ',r);
16 //
```

---

### Scilab code Exa 5.a.2 Finding density of copper

```
1 // Chapter 5 additional Example 2
2 //


---


3 clc;
4 clear;
5
6 // input data
7 // Copper has FCC structure
8
9 r      = 1.278;           // Atomic radius in
   angstrom
10 N     = 6.023*10^26;    // Avagadros number in
   atoms/kilomole
11 A     = 63.54;          // Atomic weight of
   copper
12 n     = 4;              // No. of atoms per unit
   cell for FCC
13
14 // Calculations
15 r1    = r*10^-10;       // Radius conversion
   from angstrom to m
16 a    = (4*r1)/sqrt(2); // lattice parameter
   for FCC
17 p    = (n*A)/(N*a^3); // Density of copper
18
19 // Output
20
21 mprintf(' Density of copper = %3.2f kg/m^3 ',p);
22 //
```

---

### Scilab code Exa 5.a.3 Finding distance between adjacent atoms

```
1 // Chapter 5 additional Example 3
2 //


---


3 clc;
4 clear;
5
6 // input data
7 // NaCl has FCC structure
8
9 ANa      = 23;                      // atomic wt of sodium
10 AC1      = 35.45;                   // atomic wt of chlorine
11 N        = 6.023*10^26;           // Avagadros number in
12          atoms/kilomole
13 n        = 4;                      // No. of atoms per unit
14          cell for FCC
15 p        = 2180;                   // density in kg/m^-3
16
17 // Calculations
18
19 // p      = (n*A)/(N*a^3);       density
20 A        = ANa+AC1;                // atomic wt of NaCl
21 a        = ((n*A)/(N*p))^(1/3);   // lattice constant
22 r        = a/2;                   // Distance b/w two
23          adjacent atoms
24 //Output
25 mprintf('Distance between two adjacent atoms is r =
26 %3.2e m',r);
27 //
```

---

### Scilab code Exa 5.a.4 Calculating atomic radius of Fe

```
1 // Chapter 5 additional Example 4
2 //


---


3 clc;
4 clear;
5
6 // input data
7 // iron has BCC structure
8
9 r      = 1.273;           // Atomic radius in
  angstrom
10 N     = 6.023*10^26;    // Avagadros number in
   atoms/kilomole
11 A     = 55.85;          // Atomic weight of Fe
12 n     = 2;              // No. of atoms per unit
   cell for BCC
13 p     = 7860;           // density in kg/m^-3
14
15 // Calculations
16
17 // p     = (n*A)/(N*a^3);   density
18
19 a     = ((n*A)/(N*p))^(1/3); // lattice constant
20 a1    = a*10^10;          // m to angstrom
   conversion
21 r     = (a1*sqrt(3))/4     // atomic radius
   for BCC
22
23 //Output
24 mprintf('The Radius of the Fe = %3.3f ',r);
25 //
```

---

---

### Scilab code Exa 5.a.5 calculating lattice constant

```
1 // Chapter 5 additional Example 5
2 //
3 clc;
4 clear;
5
6 // input data
7 // KBr has FCC structure
8
9 N      = 6.023*10^26;           // Avagadros number in
10 atoms/kilomole
11 A      = 119;                  // Atomic weight of
12 potassium bromide
13 n      = 4;                   // No. of atoms per unit
14 cell for FCC
15
16 p      = 2700;                // density in kg/m^-3
17
18 // Calculations
19
20 // p      = (n*A)/(N*a^3);    density
21
22 a      = ((n*A)/(N*p))^(1/3); // lattice constant
23 a1     = a*10^10;              // m to angstrom
24 conversion
25
26 // Output
27 mprintf('Lattice constant = %3.1f ',a1);
28 //
```

---

---

### Scilab code Exa 5.a.6 Calculating No of atoms per unit cell

```
1 // Chapter 5 additional Example 6
2 //
3 clc;
4 clear;
5 // input data
6 a      = 4.3*10^-10;           // Lattice constant in
7 p      = 960;                 // Density of crystal in
8 kg/m^3
9 A      = 23;                  // Atomic wt
10 N     = 6.023*10^26;         // avogadros no in atoms
11 /kilomole
12
13 n      = (p*N*(a^3))/A;     // No. of atoms per
14 unit cell
15
16 mprintf('No. of atoms per unit cell = %3.0f (BCC)',n)
17 );
```

---

### Scilab code Exa 5.a.7 Finding Volume of unit cell

```
1 // Chapter 5 additional Example 7
```

```

2 // _____
3 clc;
4 clear;
5 // input data
6 // given crystal has BCC structure
7 r = 1.2*10^-10;           // atomic radius in m
8
9 // Calculations
10
11 a = (4*r)/sqrt(3);      // lattice constant
12 V = a^3;                 // volume of cell
13
14 //Output
15 mprintf('Volume of the cell = %3.3e m^3 ',V);
16 //

```

---

### Scilab code Exa 5.a.8 Finding planar atomic density

```

1 // Chapter 5 additional Example 8
2 // _____
3 clc;
4 clear;
5 // input data
6 a = 4*10^-10;           // lattice constant of the
    crystal
7 h = 1;                  // miller indice
8 k = 0;                  // miller indice
9 l = 0;                  // miller indice
10

```

```

11 // Calculations
12
13 // in fig consider (100) plane. the no of atoms in
14 N = 4*(1/4);           // Number of atoms
15 p = N/(a*a);          // planar atomic density in
16 atoms/m^2
17 p1 = p*10^-6;          // planar atomic density in
18 atoms/mm^2
19 mprintf('planar atomic density = %3.2e atoms/mm^2 ', p1);
20 //

```

---

### Scilab code Exa 5.a.9 Finding miller indices of planes

```

1 // Chapter 5 additional Example 9
2 //

```

---

```

3 clc;
4 clear;
5 // input data
6 // in fig 5(b) the given plane is parallel to X and
   Z axes.Thus,its numerical intercepts on these
   axes is infinity
7 //The numerical intercept on y axis is 1/2. Thus the
   numerical intercepts of plane is (    1/2    )
8 mprintf('Miller indices of plane shown in fig 5.(b)
   = (0 2 0)\n');
9 // in fig 5(c) the given plane is parallel to Z axis
   .Thus its numerical intercept on z axis is

```

```

    infinity
10 // The numerical intercept on x axis is 1 and y axis
   is 1/2. this numerical intercepts on plane is (1
      1/2      )
11 mprintf(' Miller indices of plane shown in fig 5.(c)
   = (1 2 0)\n')
12 // in fig 5(d) the given plane is parallel to Z axis
   .Thus its numerical intercept on z axis is
   infinity
13 // The numerical intercept on x axis is 1/2 and y
   axis is 1/2. this numerical intercepts on plane
   is (1/2 1/2      )
14 mprintf(' Miller indices of plane shown in fig 5.(d)
   = (2 2 0)\n')
15 //

```

---

### Scilab code Exa 5.a.11 Calculating interplanar spacing

```

1 // Chapter 5 additional Example 11
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 // (311) plane in simple cubic lattice
8 h      = 3;           // miller indice
9 k      = 1;           // miller indice
10 l     = 1;           // miller indice
11 a     = 2.109*10^-10 // lattice constant in m
12
13 // Calculations

```

```

14 dhkl      = a/sqrt((h^2)+(k^2)+(l^2)); // interplanar
     distance
15
16 // Output
17 mprintf('d = %3.3e m', dhkl);
18 //

```

---

### Scilab code Exa 5.a.12 Finding lattice constant

```

1 // Chapter 5 additional Example 12
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7
8 h          = 1;           // miller indice
9 k          = 1;           // miller indice
10 l          = 0;          // miller indice
11 d          = 2.86*10^-10 // interplanar distance in m
12
13 // Calculations
14 a          = d*sqrt((h^2)+(k^2)+(l^2)); // interplanar
     distance
15
16 // Output
17 mprintf('Lattice constant a = %3.3e m', a);
18 //

```

---

### Scilab code Exa 5.a.13 Proof

```
1 // Chapter 5 Additional Example 13
2 //


---


3 clc;
4 clear;
5
6 h1      = 1;
7 h0      = 0;
8 k0      = 0;
9 l0      = 0;
10 l1     = 1;
11 // calculations
12
13 // we know that dhkl = a/sqrt( h^2 + k^2 + l^2)
14 // let sqrt( h^2 + k^2 + l^2) = p
15 p101    = sqrt( h1^2 + k0^2 + l1^2);
16 p100    = sqrt( h1^2 + k0^2 + l0^2);
17 p001    = sqrt( h0^2 + k0^2 + l1^2);
18
19 // output
20 mprintf('d101 : d100 : d001 :: a/3.4 f : a/d : a/d
', p101, p100, p001);
21 //
```

---

### Scilab code Exa 5.a.14 Finding ratio of intercepts

```
1 // Chapter 5 additional Example 14
```

```

2 // _____
3 clc;
4 clear;
5
6 // if a plane cut intercepts of lengths l1,l2,l3 the
    on three crystal axes ,then
7 // l1 : l2 : l3 = pa : pq :rc
8 // where a,b and c are primitive vectors of the unit
    cell and p,q and r are numbers related to miller
    indices (hkl) of plane by relation
9 // 1/p : 1/q : 1/r = h : k : l
10 //since , the crystal is simple cubic a = b = c and
    given that h = 1, k = 1 and l = 1
11 // p : q : r = 1/h : 1/k : 1/l = 1/1 : 1/1 : 1/1
12 // p : q : r = 1 : 1 : 1
13 //similarly l1 : l2 : l3 = 1a : 1a : 1a
14 mprintf('ratio of intercepts on the three axes by
    (111) plane is l1 : l2 : l3 = 1 : 1 : 1');

```

---

### Scilab code Exa 5.a.15 Finding interplanar distance

```

1 // Chapter 5 additional Example 15
2 //
3 clc;
4 clear;
5
6 //input data
7 r    = 1.246*10^-10;           // atomic radius in m
8 h1   = 1                      // miller indice
9 h2   = 2                      // miller indice
10 k0  = 0                      // miller indice

```

```

11 k1 = 1 // miller indice
12 k2 = 2 // miller indice
13 l0 = 0 // miller indice
14 l1 = 1 // miller indice
15
16 // Calculations
17 a = (4*r)/sqrt(2); // lattice constant
18 d111 = a/sqrt((h1^2)+(k1^2)+(l1^2)); // interplanar
    distance
19 d200 = a/sqrt((h2^2)+(k0^2)+(l0^2)); // interplanar
    distance
20 d220 = a/sqrt((h2^2)+(k2^2)+(l0^2)); // interplanar
    distance
21
22 // Output
23 mprintf('d111 = %3.3e m\n d200 = %3.4e m\n d220 = %3
    .3e m\n', d111, d200, d220);
24 //

```

---

### Scilab code Exa 5.a.16 Finding Miller indices

```

1 // Chapter 5 additional Example 16
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 // the intercept along X-axis be c1 = a
8 // the intercept along Y-axis be c2 = b/2 and
9 // the intercept along Z-axis be c3 = 3c
10 // Therefore , p = c1/a = a/a = 1

```

```

11 // q = c2/b = (b/2)/b = 1/2
12 // r = c3/c = (3c)/c = 3
13 // therefore h = 1/p = 1
14 // k = 1/q = 2
15 // l = 1/r = 1/3
16 // lcm of 1 1 and 3 = 3
17 h = 1
18 k = 2
19 l = 1/3
20 p = [1 1 3]
21 s = lcm(p);
22 h1= s*h
23 k1= s*k
24 l1= s*l;
25 // Output
26 mprintf( '(h k l) = (%d %d %d)',h1,k1,l1);
27 //

```

---



---

### Scilab code Exa 5.a.17 Calculating Wavelength of X ray

```

1 // Chapter 5 Additional Example 17
2 //

```

---



---

```

3 clc;
4 clear;
5
6 //input data
7
8 d = 1.3*10^-10      // interplanar distance
9 n = 1;                // given first order
10 theta = 23;           // Bragg reflection angle in
                           degrees

```

```

11
12 // Calculations
13 theta1 = theta*pi/180; // degree to radian
   conversion
14 // d = (n*lamda)/(2* sin ); by Braggs law
   _____ 1
15 lamda = (2*d*sin(theta1)/n)
16
17 // Output
18 mprintf('Wavelength of X-ray = %3.4f ', lamda
   *10^10);
19 //

```

---

### Scilab code Exa 5.1 Finding lattice parameter and Density of copper

```

1 // Chapter 5 Example 1
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 //Copper has FCC structure
8
9 r = 1.273; // Atomic radius in
   angstrom
10 N = 6.023*10^26; // Avagadros number in
   atoms/kilomole
11 A = 63.5; // Atomic weight of
   copper in grams
12 n = 4; // No. of atoms per unit
   cell for FCC

```

```

13
14 // Calculations
15 r1      = r*10^-10;           // Radius conversion
   from angstrom to m
16 a       = (4*r1)/sqrt(2);    // lattice parameter
   for FCC
17 p       = (n*A)/(N*a^3);    // Density of copper
18
19 // Output
20
21 fprintf('Lattice Constant a = %3.1e m\n Density of
copper = %3.1f kg/m^3',a,p);

```

---

### Scilab code Exa 5.2 Finding Miller indice and interplanar distance

```

1 // Chapter 5 Example 1
2 //

=====

3 clc;
4 clear;
5
6 //input data
7 //given intercepts 3,4 and , the reciprocals of
   intercepts is
8 // (1/3):(1/4):(1/ )
9 // LCM = 12
10 // multiplying by LCM we get miller indices
11 // miller indices of a plane are the smallest
   integers of the reciprocals of its intercepts
12 // therefore miller indices(h k l) is (4 3 0);
13
14 h    = 4;          // miller indice
15 k    = 3;          // miller indice
16 l    = 0;          // miller indice

```

```

17 a      = 2;           // primitive vector of lattice in
                        angstrom
18
19 // Calculations
20
21 dhkl    = a/sqrt((h^2)+(k^2)+(l^2)); // interplanar
                        distance
22
23 // Output
24 mprintf('Miller indices = (4 3 0)\n');
25 mprintf(' The interplanar distance d = %3.1f      ,\n',
            dhkl);
26 //


---



```

### Scilab code Exa 5.3 Finding Radius of an atom

```

1 // Chapter 5 Example 3
2 //


---


3 clc;
4 clear;
5
6 //input data
7 // -Iron solidifies to BCC structure
8
9 r      = 1.273;           // Atomic radius in
                           angstrom
10 N     = 6.023*10^26;     // Avagadros number in
                           atoms/kilomole
11 A     = 55.85;           // Atomic weight of -
                           Iron in kilograms
12 n     = 2;               // No. of atoms per unit

```

```

    cell for BCC
13 p      = 7860;           // density in kg/m^3
14
15 // Calculations
16
17 // p      = (n*A)/(N*a^3);   density
18
19 a      = ((n*A)/(N*p))^(1/3); // lattice constant
20 a1     = a*10^10;           // m to angstrom
21 r      = (a1*sqrt(3))/4      // atomic radius
for BCC
22
23 // Output
24 mprintf('The Radius of the atom = %3.5f \n',r);
25 mprintf(' Note : atomic wt taken as 55.58*10^-3
instead of 55.85 in calculation')
26 //

```

---

### Scilab code Exa 5.4 Calculating interatomic Spacing

```

1 // Chapter 5 Example 4
2 //

3 clc;
4 clear;
5
6 //input data
7 lamda = 1.5418;           // wavelength in
8 h      = 1;                // miller indice
9 k      = 1;                // miller indice
10 l     = 1;                // miller indice

```

```

11 n          = 1;                      // given first order
12 theta      = 30;                    // diffraction angle in
   degrees
13
14 // Calculations
15 theta1    = theta*%pi/180;        // degree to radian
   conversion
16 // d      = (n*lamda)/(2* sin );    by Braggs law
   _____ 1
17 // d      = a/sqrt ((h^2)+(k^2)+(l^2));  interplanar
   distance _____ 2
18 // equating 1 and 2
19
20 a          = (n*lamda*sqrt((h^2)+(k^2)+(l^2)))/(2*sin(
   theta1)))
21
22 // Output
23 mprintf('Interatomic spacing a = %f ',a);
24 //

```

---

**Scilab code Exa 5.5 Finding interplanar distance between planes**

```

1 // Chapter 5 Example 5
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 h1      = 1;                      // miller indice
8 k1      = 1;                      // miller indice
9 l1      = 1;                      // miller indice

```

```

10 h0          = 0;                      // miller indice
11 k0          = 0;                      // miller indice
12 l0          = 0;                      // miller indice
13
14 // calculations
15 // dhkl     = a/sqrt((h^2)+(k^2)+(l^2)); //
   interplanar distance
16 // assume a = 1(constant) for easier calculation in
   scilab
17
18 a            = 1;
19 d100         = a/sqrt((h1^2)+(k0^2)+(l0^2)); //
   interplanar distance
20 d110         = a/sqrt((h1^2)+(k1^2)+(l0^2)); //
   interplanar distance
21 d111         = a/sqrt((h1^2)+(k1^2)+(l1^2)); //
   interplanar distance
22
23 // Output
24 mprintf('d100 : d110 : d111 = %d : %3.2f : %3.2f' ,
   d100,d110,d111);
25
26 //
```

---

### Scilab code Exa 5.6 Finding number of unit cells

```

1 // Chapter 5 Example 6
2 //
```

---

```

3 clc;
4 clear;
5
```

```

6 // input data
7 // Aluminium is FCC
8 a = 0.405*10^-9; // lattice constant
of aluminium
9 t = 0.005*10^-2; // thickness of
aluminium foil in m
10 s = 25*10^-2; // side of square in
m
11
12 // Calculations
13 VUC = a^3; // volume of unit
cell
14 Val = (s^2)*t // volume of
aluminium foil (area*thickness)
15 N = Val/VUC // Number of unit
cells
16
17 // Output
18 mprintf('Number of unit cells = %3.3e',N);
19 //

```

---

### Scilab code Exa 5.7 Finding percentage change in volume

```

1 // Chapter 5 Example 7
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7 // metallic iron changes from BCC to FCC form at
910 degrees

```

```

8 rb      = 0.1258*10^-9;           // atomic radius of BCC
    iron atom
9 rf      = 0.1292*10^-9;           // atomic radius of FCC
    iron atom
10
11 // Calculations
12
13 ab      = (4*rb)/(sqrt(3));     // lattice constant for
    BCC
14 Vbcc    = (ab^3)/2;             // volume occupied by
    one BCC atom
15 af      = (4*rf)/(sqrt(2))     // lattice constant for
    FCC
16 Vfcc    = (af^3)/4;             // volume occupied by
    one FCC atom
17 dv      = ((Vbcc-Vfcc)/Vbcc)*100 // percentage change in volume
18
19 // output
20 mprintf('During the structural change the percentage
    change in volume = %3.4f',dv);
21 //

```

---

### Scilab code Exa 5.8 Finding lattice constant

```

1 // Chapter 5 Example 8
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data

```

```

7 //Copper Crystallines in FCC structure
8
9 p = 8960; // Density of copper in
   kg/m^3
10 N = 6.023*10^26; // Avagadros number in
    atoms/kilomole
11 A = 63.5; // Atomic weight of
    copper in kg/mol
12 n = 4; // No. of atoms per unit
    cell for FCC
13
14 //Calculations
15
16 a = ((n*A)/(N*p))^(1/3);
17
18 //Output
19
20 mprintf('Lattice Constant a = %3.4f \n',a*10^10);
21 mprintf(' atomic wt of copper is taken as 63.5*10^-3
           instead of 63.5 in textbook')
22 //

```

---

### Scilab code Exa 5.9 Calculating d spacing

```

1 // Chapter 5 Example 9
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 // (100) planes in rock crystal

```

```

8 h      = 1;           // miller indice
9 k      = 0;           // miller indice
10 l     = 0;           // miller indice
11 a     = 2.814         // lattice constant in
12
13 // Calculations
14 dhkl   = a/sqrt((h^2)+(k^2)+(l^2)); // interplanar
   distance
15
16 // Output
17 mprintf('d-spacing for (100) plane in rock salt = %3
   .3f', dhkl);
18 //

```

---

### Scilab code Exa 5.10 Calculating lattice constant

```

1 // Chapter 5 Example 10
2 //

```

---

```

3 clc;
4 clear;
5
6 // input data
7 // FCC structured crystal
8
9 p      = 6250;           // Density of crystal in
   kg/m^3
10 N     = 6.023*10^26;    // Avagadros number in
   atoms/kilomole
11 A     = 60.2;           // molecular weight
12 n     = 4;              // No. of atoms per unit
   cell for FCC

```

```

13
14 // Calculations
15
16 a = ((n*A)/(N*p))^(1/3);
17
18 //Output
19
20 mprintf('Lattice Constant a = %3.1e m ',a);
21 //

```

---

### Scilab code Exa 5.11 Calculating interplanar distance

```

1 // Chapter 5 Example 11
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 // (321) plane in simple cubic lattice
8 h = 3; // miller indice
9 k = 2; // miller indice
10 l = 1; // miller indice
11 a = 4.12 // inter atomic space
12
13 // Calculations
14 dhkl = a/sqrt((h^2)+(k^2)+(l^2)); // interplanar
   distance
15
16 // Output
17 mprintf('d = %3.2f ', dhkl);
18 //

```

---

---

### Scilab code Exa 5.12 Calculating lattice constant of Fe

```
1 // Chapter 5 Example 12
2 //
3 clc;
4 clear;
5
6 // input data
7 // BCC structured crystal
8
9 p      = 7860;           // Density of iron in kg
10      /m^3
11 N      = 6.023*10^26;   // Avagadros number in
12      atoms/kilomole
13 A      = 55.85;          // Atomic weight
14 n      = 2;              // No. of atoms per unit
15      cell for BCC
16
17 // Calculations
18
19 a      = ((n*A)/(N*p))^(1/3); // lattice constant
20
21 // Output
22
23 mprintf('Lattice Constant of Fe = %3.3f \n',a
24      *10^10);
25 mprintf(' Note: density of iron is taken as 7.86
26      instead of 7860 in calculation ')
27 //
```

---

---

### Scilab code Exa 5.14 Finding Volume of Unit cell

```
1 // Chapter 5 Example 14
2 //


---


3 clc;
4 clear;
5
6 // input data
7 r           = 0.123*10^-10;           // Radius of the
                                         atom
8
9 // Calculations
10 a          = (4*r)/sqrt(3);        // Lattice constant
                                         in m For a BCC structure
11 V          = a*a*a;              // Volume of BCC
12
13 // Output
14 mprintf('Volume of the unit cell = %3.4e m^3',V);
15 //
```

---

### Scilab code Exa 5.15 Finding Miller indices

```
1 // Chapter 5 Example 15
2 //


---


3 clc;
```

```

4 clear;
5
6 // input data
7 a = 0.05;      // unit cell edge of an orthorhombic
                  crystal in nm
8 b = 0.05;      // unit cell edge of an orthorhombic
                  crystal in nm
9 c = 0.03;      // unit cell edge of an orthorhombic
                  crystal in nm
10 Ia = 0.025    // intercept on 'a' in nm
11 Ib = 0.02     // intercept on 'b' in nm
12 Ic = 0.01     // intercept on 'c' in nm
13
14 // Calculations
15
16 h = a/Ia;     // miller indice h
17 k = b/Ib;     // miller indice k
18 l = c/Ic;     // miller indice l
19
20 // Output
21 mprintf('Miller indices (h k l) = (%d %d %d)',h,k,l)
               ;
22 //
=====
```

---

### Scilab code Exa 5.16 Finding volume of unit cell

```

1 // Chapter 5 Example 16
2 //
=====

3 clc;
4 clear;
5 // Magnesium has HCP structure
```

```

6 // for HCF(Hexagonal closed packed structure)
    consider the relation between 'c' and 'a';
7 // c/a = sqrt(8/3) = 1.6329
8 //input data
9 r = 0.1605*10^-9;           // radius of magnesium atom
    in m
10
11 // Calculations
12
13 a = 2*r                  // lattice constant of HCP
14 c = a*sqrt(8/3);          // relation b/w c and a in
    HCP
15 V = (3*3^0.5)*(a*a*c)/2; //Volume of unit
    cell in m^3
16
17 // Output
18 mprintf('Volume of the unit cell of magnesium = %3.3
    e m^3',V);
19 //

```

---

### Scilab code Exa 5.17 Finding interplanar distance

```

1 // Chapter 5 Example 17
2 //

```

---

```

3 clc;
4 clear;
5
6 //input data
7 // (101),(221) planes in simple cubic lattice
8 h1 = 1;                      // miller indice
9 k0 = 0;                      // miller indice

```

```

10 l1      = 1;           // miller indice
11 h2      = 2;           // miller indice
12 k2      = 2;           // miller indice
13 l1      = 1;           // miller indice
14 a       = 4.2          // inter atomic space
15
16 // Calculations
17 d101    = a/sqrt((h1^2)+(k0^2)+(l1^2)); // 
    interplanar distance
18 d221    = a/sqrt((h2^2)+(k2^2)+(l1^2)); // 
    interplanar distance
19
20
21 // Output
22 mprintf('d(101) = %3.4f\n d(221) = %3.1f', 
    d101,d221);
23 //

```

---