

Scilab Textbook Companion for  
Semiconductor Physics And Devices  
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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

## Introduction to Quantum Mechanics

**Scilab code Exa 1.1** Photon energy

```
1 // Chapter 1_Principles of Quantum Mechanics
2 //Caption_Photon Energy
3 //Ex_1//page 3
4 disp("X-Rays wavelength lambda=0.708*10^-8 cm");
5 lambda=0.708*(10^-10); //Wavelength
6 h=6.625*(10^-34); // Plank's constant
7 c=3*10^8 //speed of light
8
9 E=(h*c)/lambda;
10 printf('The photon energy corresponding to given
wavelength is %fd J\n',E)
11 Evv=E/(1.6*(10^-19));
12 printf('Energy in the units of ev is %fd eV \n',Evv)
```

---

**Scilab code Exa 1.2** Broglie wavelength

```

1 // Chapter 1_Principles of Quantum Mechanics
2 //Caption_Broglie wavelength
3 //Ex_2//page 5
4 disp("Velocity of electron is  $10^7$  cm/s");
5 h=6.625*(10^-34); // Plank's constant
6 m=9.11*(10^-31); //Mass of electron
7 v=10^5; //Velocity of electron
8
9 p=m*v; //Momentum
10 printf('Momentum is %fd \n',p)
11 lambda=(h/p)*10^10; //De Broglie's wavelength in
    angstrom
12 printf('The De Broglie wavelength is %fd Angstrom\n',
    lambda);

```

---

### Scilab code Exa 1.3 Electron in infinite potential well

```

1 // Chapter 1_Principles of Quantum Mechanics
2 //Caption_Electron in an infinite potential well
3 //Ex_3//page 14
4 disp("Width of electrons infinite potential well is
    5 Angstrom");
5 h=1.054*(10^-34); // Plank's constant
6 m=9.11*(10^-31); //Mass of electron
7 v=10^5; //Velocity of electron
8 a=5*(10^-10);
9 c=1.6*(10^-19);
10 n1=1; // assume
11 En1=((h^2)*(n1^2)*(%pi^2)/(2*m*a^2))/c
12 printf('Energy in first energy level is %fd eV\n',
    En1)
13 n2=2;..... //assume
14 En2=((h^2)*(n2^2)*(%pi^2)/(2*m*a^2))/c
15 printf('Energy in first energy level is %fd eV\n',
    En2)

```

```

16 n3=3;      //assume
17 En3=((h^2)*(n3^2)*(%pi^2)/(2*m*a^2))/c
18 printf('Energy in first energy level is %fd eV\n',
En3)

```

---

**Scilab code Exa 1.4** Penetration depth of a particle impinging on a potential barrier

```

1 // Chapter 1_Principles of Quantum Mechanics
2 //Caption_Penetration depth of a particle impinging
   on a potential barrier
3 //Ex_4//page 18
4 v=10^5;           // Given velocity of electron
5 m=9.11*(10^-31); // mass of electron
6 c=1.6*(10^-19)
7 E=((1/2)*m*(v^2))/c;
8 V0=2*E;          //Assume that the potential barrier at x
   =0 is twice as large as total energy of the
   incident particle
9 printf('Energy of the particle is %fd eV\n',E)
10 h=1.054*(10^-34)
11 d=((h^2)/(2*m*E*c))^(1/2)*10^10;
12 printf('The distance at which the wave function
   magnitude has decayed to e^-1 of its value at x=0
   is %2.1f Angstrom \n',d )

```

---

**Scilab code Exa 1.5** Probability of an electron tunneling through a potential barrier

```

1 // Chapter 1_Principles of Quantum Mechanics
2 //Caption_Probability of an electron tunneling
   through a potential barrier
3 //Ex_5//page 21

```

```
4 E=2;      //energy of electron in eV
5 Vo=20;    //potential barrier in eV
6 w=3;      //width of potential barrier in angstrom
7 m=9*(10^-31)
8 h=1.054*(10^-34)
9 c=1.6*(10^-19)
10 K=((2*m)*(Vo-E)*c/(h^2))^(1/2)
11 printf('Factor K is %fd m^-1 \n',K)
12 l=(-2*K*w*(10^-10))
13 a=%e^l
14 x=E/Vo;
15 T=16*x*(1-x)*a;
16 printf('Transmission coefficient i.e. the
           probability of electron to tunnel through the
           potential barrier is %fd \n',T)
```

---

# Chapter 2

## Introduction to the quantum theory of solids

**Scilab code Exa 2.1** Change in kinetic energy

```
1 // Chapter 2_Introduction to the quantum theory of  
solids  
2 //Caption_Change in kinetic energy of electron  
3 //Ex_1//page 37  
4 v=10^5;  
5 delV=0.01;  
6 m=9.1*10^-31;  
7 c=1.6 *(10^-19)  
8 delE=m*v*delV/c;  
9 printf('Change in kinetic energy for a very small  
change in velocity of electron is %1.10fd eV \n',  
delE)
```

---

**Scilab code Exa 2.2** Lowest allowed energy bandwidth

```
1 // Chapter 2_Introduction to the quantum theory of  
solids
```

```

2 //Caption_Lowest allowed energy bandwidth
3 //Ex_2//page 46
4
5 m=9.11*(10^-31)           //mass of electron
6 h=1.054*(10^-34)
7 a=5*(10^-10)
8 c=1.6*(10^-19)           //electron charge
9 E=((%pi^2)*(h^2))/(2*m*(a)^2)
10 E2=E/c;
11 E1= 1.053      // For alpha a=2.628, energy is given
                  in eV
12 delE=E2-E1
13 printf('The allowed energy bandwidth is %fd eV\n',
          delE)

```

---

### Scilab code Exa 2.3 Density of States

```

1 // Chapter 2_Introduction to the quantum theory of
  solids
2 //Caption_Density of States
3 //Ex_3//page 63
4 m=9.11*(10^-31)           //mass of electron
5 h=6.625*(10^-34)
6 E=1.6*(10^-19)
7 N=((4*%pi*(2*m)^(1.5))/(h^3))*(E^(1.5))*(2/3)
8 printf('Density of states per unit volume with
          energies between 0 and 1 eV is %f states per unit
          vol \n',N)

```

---

### Scilab code Exa 2.4 The Fermi Dirac Probability Function

```

1 // Chapter 2_Introduction to the quantum theory of
  solids

```

```

2 //Caption_The Fermi Dirac Probability Function
3 //Ex_4//page 68
4 Ni=10      //given no. of particles
5 gi=10      //given no. of quantum states
6 P=factorial(gi)/(factorial(Ni)*(factorial(gi-Ni)))
    //Possible no of ways of relizing this
    distribution
7 printf('This distribution can be realized in %i ways
\n',P)

```

---

### Scilab code Exa 2.5 The Fermi Dirac Probability Function

```

1 // Chapter 2_Introduction to the quantum theory of
solids
2 //Caption_The Fermi Dirac Probability Function
3 //Ex_5//page 68
4 Ni=9      //given no. of particles
5 gi=10      //given no. of quantum states
6 P=factorial(gi)/(factorial(Ni)*(factorial(gi-Ni)))
    //Possible no of ways of relizing this
    distribution
7 printf('This distribution can be realized in %i ways
\n',P)

```

---

### Scilab code Exa 2.6 The Distribution function and the Fermi Energy

```

1 // Chapter 2_Introduction to the quantum theory of
solids
2 //Caption_The Distribution function and the Fermi
Energy
3 //Ex_6//page 71
4 T=300      // Temperature in kelvin
5 x=3       //x= (E-Ef)/kT

```

```
6 fE=100/(1+e^x);  
7 printf('Probability that an energy level 3kT above  
the Fermi energy is occupied by an electron is %1  
.2fd%%\n',fE)
```

---

### Scilab code Exa 2.7 The Distribution function and the Fermi Energy

```
1 // Chapter 2_Introduction to the quantum theory of  
solids  
2 //Caption_The Distribution function and the Fermi  
Energy  
3 //Ex_7//page 72  
4 fE=0.01  
5 Ef=6.25 //Fermi energy in eV  
6 a=1/(1-fE)  
7 x=log(a-1)  
8 k=1.38*(10^-23)  
9 T=(-0.30)*1.6*(10^-19)/(k*x)  
10 printf('Temperature at which there is a 1 percent  
probability that an energy state is empty is %3.1  
fd K\n',T)
```

---

# Chapter 3

## The Semiconductor in Equilibrium

**Scilab code Exa 3.1** Equilibrium Distribution of Electrons and holes

```
1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Equilibrium Distribution of Electrons and
holes
3 //Ex_1//page 85
4 T=300          // Temperatire in kelvin
5 Nc=2.8*(10^19)      //Effective density of states
function in the conduction band in per cm cube
6 de1E=0.25      //Fermi energy is 0.25eV below the
conduction band
7 k=1.389*(10^-23)      //Boltzmann constant
8 kT=0.0259
9 fF=%e^(-de1E/(kT))
10 no=Nc*fF
11 printf('The thermal equilibrium electron
concentration in siliconn is %1.2 fd per cm^3 ',no)
```

---

### Scilab code Exa 3.2 Equilibrium Distribution of Electrons and holes

```
1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Equilibrium Distribution of Electrons and
holes
3 //Ex_2//page 87
4 T=400;
5 N=1.04*(10^19)
6 kT=0.0259*(T/300);
7 Nv=N*(T/300)^(1.5)
8 po=Nv*(%e^(-0.27/kT))
9 printf('The thermal equilibrium hole concentration
in silicon at T=400K ==%fd per cm^3 \n',po)
```

---

### Scilab code Exa 3.3 Intrinsic carrier concentration

```
1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Intrinsic carrier concentration
3 //Ex_3//page 90
4 T1=300;      //Given temperature in kelvin
5 T2=450;
6 Nc1=4.7*(10^17) //effective density of state
function in cm^-3
7 Nv1=7*(10^18)
8 Eg=1.42      //bandgap energy in eV
9 kT=0.0259*(T2/T1);
10 ni1=(Nc1*Nv1*exp((-Eg)/0.0259))^.5
11 ni2=(Nc1*Nv1*(T2/T1)^3*exp(-Eg/kT))^.5
12 printf('The intrinsic carrier concentration in
gallium arsenide at T=300k is %fd per cm cube and
at 450k is %fd //cm^3 ', ni1, ni2)
```

---

### Scilab code Exa 3.4 Intrinsic fermi level position

```

1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Intrinsic fermi level position
3 //Ex_4//page 92
4 T=300;      //temperature in kelvin
5 mnr =1.08    //relative effective mass of negative
               charge carrier
6 mpr=0.56     //relative effective mass of positive
               charge carrier
7 kT=0.0259
8 Efm=(3/4)*kT*log(mpr/mnr)      //The intrinsic fermi
               level with respect to the center of bandgap
9 EfmF=-(Efm)*1000
10 printf('The intrinsic feremi level in silicon is %1
               .1 fd meV below the midgap energy ',EfmF)

```

---

### Scilab code Exa 3.5 Extrinsic Semiconductor

```

1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Extrinsic Semiconductor
3 //Ex_5//page 101
4 T=300      //temperature in kelvin
5 Nc=2.8*(10^19);
6 Nv=1.04*(10^19);      //
7 Fe=0.25      //Fermi energy is FeeV below the
               conduction band
8 Eg=1.12      // Bandgap energy of silicon is Eg in eV
9 no=Nc*exp(-Fe/0.0259);
10 po=Nv*exp(-(Eg-Fe)/0.0259);
11 printf('Thermal equilibrium concentration of
               electrons is %1.2 fd cm ^-3 and of holes is %1.2 fd
               cm^-3 ',no,po)

```

---

### Scilab code Exa 3.6 Extrinsic Semiconductor

```

1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Extrinsic Semiconductor
3 //Ex_6//page 104
4 nf=2      // nf=(Ef-Ec)/kT
5 Fe=52     //Fermi energy is above the conduction band
           by Fe meV
6 T=300;
7 Nc=2.8*(10^19);
8 F(nf)=2.3 // Value of fermi dirac integral from the
           graph
9 no=(2/((%pi)^0.5))*Nc*F(nf)
10 printf('Electron concentration using fermi dirac
           integral is %fd per cm cube ',no)

```

---

### Scilab code Exa 3.7 Statistics of acceptors and donors

```

1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Statistics of acceptors and donors
3 //Ex_7//page 108
4 T=300;
5 Nd=10^16    // donor concentration per cm cube
6 kT=0.0259
7 Ecd=0.045   //Ec-Ed
8 Nc=2.8*(10^19);
9 x=1/(1+(Nc/(2*Nd))*exp(-(Ecd)/kT))
10 printf('Fraction of total electrons still in the
           donor state is %fd ',x)

```

---

### Scilab code Exa 3.8 Statistics of acceptors and donors

```

1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Statistics of acceptors and donors
3 //Ex_8//page 110

```

```
4 Na=10^16    //Acceptor concentration
5 kT=0.0259
6 Nv=1.04*(10^19);
7 Eav=0.045
8 x=0.1    //90%of acceptor atoms are ionized
9 y=((1/x)-1)*4*Na/Nv;
10 //((T/300)^1.5*exp(-Eav/kT*(T/300))=y
11 //By trial and error
12 printf('Required temperature is 193 K')
```

---

### Scilab code Exa 3.9 Charge Neutrality

```
1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Charge Neutrality
3 //Ex_9//page 112
4 T=300    //temperature in kelvin
5 Nd=10^16    //donor concentration per cm cube
6 Na=0
7 ni=1.5*(10^10)    //intrinsic carrier concentration
8 no=((Nd-Na)/2)+(((Nd-Na)/2)^2+ni^2)^0.5
9 po=ni^2/no;
10 printf('The majority carrier electron concentration
           is %fd per cm cube while the minority carrier
           hole concentration is %fd per cm cube ',no,po)
```

---

### Scilab code Exa 3.10 Charge Neutrality

```
1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Charge Neutrality
3 //Ex_10//page 114
4 T=300    //temperature in kelvin
5 Nd=5*(10^13)
6 Na=0
```

---

```

7 ni=2.4*(10^13)
8 no=((Nd-Na)/2)+(((Nd-Na)/2)^2+ni^2)^0.5
9 po=ni^2/no;
10 printf('The majority carrier electron concentration
           is %fd per cm cube while the minority carrier
           hole concentration is %fd per cm cube',no,po)

```

---

### Scilab code Exa 3.11 Charge Neutrality

---

```

1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Charge Neutrality
3 //Ex_11//page 115
4 T=300      //temperature in kelvin
5 Na=10^16    //donor concentration per cm cube
6 Nd=3*(10^15)
7 ni=1.5*(10^10)   //intrinsic carrier concentration
8 po=((Na-Nd)/2)+(((Na-Nd)/2)^2+ni^2)^0.5
9 no=ni^2/po;
10 printf('The minority carrier electron concentration
           is %fd per cm cube while the majority carrier
           hole concentration is %fd per cm cube',no,po)

```

---

### Scilab code Exa 3.12 Charge Neutrality

---

```

1 // Chapter 3_The Semiconductor in Equilibrium
2 //Caption_Charge Neutrality
3 //Ex_12//page 116
4 T=550      //temperature in kelvin
5 Nc=2.8*(10^19)
6 Nv=1.04*(10^19)
7 Eg=1.12    // band gap energy in eV
8 ni=(Nc*Nv*(T/300)^3*exp(-Eg/0.0259 *(300/T)))^0.5

```

```
9 //no=1.05*Nd since the intrinsic carrier  
concentration to contribute no more than 5  
percent of the total electron concentration .  
10 Nd=(ni^2/(1.05-1))^0.5  
11 printf('The required impurity doping concentration  
is %fd per cm cube',Nd)
```

---

### Scilab code Exa 3.13 Position of Fermi Energy level

```
1 // Chapter 3_The Semiconductor in Equilibrium  
2 //Caption_Position of Fermi Energy level  
3 //Ex_13//page 116  
4 T=300 //temperature in kelvin  
5 Na=10^16 // acceptor carrier impurity in per cm  
cube  
6 Ef=0.20 //Fermi energy is Ef eV below the  
conduction band edge  
7 Nc=2.8*(10^19)  
8 Nd=Na+(Nc*exp(-Ef/0.0259))  
9 printf('The require donor impurity concentration is  
%fd per cm cube',Nd)
```

---

### Scilab code Exa 3.14 Position of Fermi Energy level

```
1 // Chapter 3_The Semiconductor in Equilibrium  
2 //Caption_Position of Fermi Energy level  
3 //Ex_14//page 121  
4 T=300 //temperature in kelvin  
5 kT=0.0259  
6 ni=1.5*(10^10) //intrinsic carrier concentration  
7 Efa=3*kT //Ef-Ea=3kT  
8 Eav=0.045
```

```
9 Efif=Eg/2-(Eav)-(Efa)      //The position of fermi  
    level at the maximum doping  
10 Na=exp(Efif/kT)*ni  
11 printf('Maximum doping is %fd per cm cube',Na)
```

---

# Chapter 4

## Carrier Transport Phenomenon

Scilab code Exa 4.1 Carrier drift

```
1 // Chapter 4_Carrier Transport Phenomenon
2 //Caption_Carrier drift
3 //Ex_1//page 134
4 T=300      //temperature in kelvin
5 Na=0
6 e=1.6*(10^-19)
7 Nd=10^16    //donor concentration in per cm cube
8 E=10        //Applied electric field in V/cm
9 ni=1.8*(10^6)
10 n=(Nd-Na)/2+(((Nd-Na)/2)^2+ni^2)^0.5
11 p=ni^2/n
12 muN=8500   //mobility of electron in gallium
               arsenide in cm^2/V-s
13 muP=400
14 J=e*(muN*n+muP*p)*E
15 printf('The drift current density for this electric
           field is %1.2f A/cm^2 ',J )
```

---

Scilab code Exa 4.2 Carrier drift

```

1 // Chapter 4_Carrier Transport Phenomenon
2 //Caption_Carrier drift
3 //Ex_2//page 143
4 T=300
5 sig=16    //CONDUCTIVITY IN (OHM-CM)^-1
6 Na=10^16   //acceptor doping concentration
7 e=1.6*(10^-19)
8 // sig=e*muN*(Nd-Na)
9 //By trial and error
10 printf('Doping concentration is 3.5*10^17 cm^-3 and
           mobility is 400 cm^2/V-S')

```

---

### Scilab code Exa 4.3 Conductivity

```

1 // Chapter 4_Carrier Transport Phenomenon
2 //Caption_Conductivity
3 //Ex_2//page 144
4 T=300
5 Nd=5*(10^15)    //donor concentration
6 R=10      //resistance in kohm
7 J=50       //current density in A/cm^2
8 V=5        //voltage in volts
9 i=V/R      //current
10 A=i/J     //cross sectional area
11 E=100
12 L=V/E     //length of the resistor
13 pho=L/(V*A)
14 // The conductivity of a compensated p-type
   semiconductor is
15 //pho=e*muP*(Na-Nd)
16 //where the mobility is a function of the total
   ionized impurity concentration Na+Nd
17 //Using trial and error , if
18 Na=1.25*(10^16)
19 muP=410

```

```
20 e=1.6*(10^-19)
21 sig=e*mup*(Na-Nd)
22 printf('Conductivity obtained is %1.2f which is
           very close to the value we need',sig)
```

---

#### Scilab code Exa 4.4 Carrier diffusion

```
1 // Chapter 4_Carrier Transport Phenomenon
2 //Caption_Carrier diffusion
3 //Ex_4//page 150
4 T=300
5 d=0.10    //distance in cm over which concentration
             varies
6 Dn=225      //diffusion coefficient
7 e=1.6*(10^-19)
8 delN=1*(10^18)-7*(10^17)
9 J=e*Dn*(delN/d)
10 printf('The diffusion current density is %1.2f A/cm
           ^3',J)
```

---

#### Scilab code Exa 4.5 Graded impurity distribution

```
1 // Chapter 4_Carrier Transport Phenomenon
2 //Caption_Graded impurity distribution
3 //Ex_5//page 153
4 T=300
5 x=0      //given 0<x<1 micrometer
6 Nd=10^16-10^19*x
7 //Taking the derivative of donor concentration , we
      have d(ND)/dx=-10^19
8 e=1.6*(10^-19)
9 Ex=-(0.0259)*(-10^19)/Nd
```

```
10 printf('The induced electric field is %1.1f d V/cm',  
Ex)
```

---

### Scilab code Exa 4.6 The Einstein relation

```
1 // Chapter 4_Carrier Transport Phenomenon  
2 //Caption_The Einstein relation  
3 //Ex_6//page 155  
4 T=300  
5 mu=1000 //mobility of a particular carrier  
6 kT=0.0259  
7 e=1.6*(10^-19)  
8 D=(kT)*mu  
9 printf('Diffusion coefficient is %1.2f d cm^2/s ',D)
```

---

### Scilab code Exa 4.7 The hall effect

```
1 // Chapter 4_Carrier Transport Phenomenon  
2 //Caption_The Hall Effect  
3 //Ex_7//page 158  
4 L=10^-3 //LENGTH IN M  
5 W=10^-2 //WIDTH IN CM  
6 d=10^-5  
7 Ix=10^-3 //current in Amp  
8 Vx=12.5  
9 e=1.6*(10^-19)  
10 Bz=500 //magnetic field in gauss  
11 Vh=-6.25*10^-3 //hall voltage  
12  
13 //A negative hall voltage for this geometry implies  
// that we have an n-type semiconductor  
14 BzT=Bz*10^-4 //magnetic field in tesla  
15 n=-(Ix*BzT)/(e*d*Vh*10^6)
```

```
16 mun=(Ix*L)/(e*n*Vx*W*d)
17 printf('Majority carrier concentration is %1.1f d cm
^ -3 and mobility is %1.1f d cm^2/V-s ',n, mun)
```

---

# Chapter 5

## Non Equilibrium excess carriers in semiconductors

Scilab code Exa 5.5 Relaxation time

```
1 // Chapter 5_Non equilibrium excess carriers in
   semiconductors
2 //Caption_Relaxation time
3 //Ex_5//page 190
4 Nd=10^16      //donor concentration
5 e=1.6*(10^-19)    //electronic charge
6 mun=1200      //mobility
7 sig=e*mun*Nd
8 epsR=11.7     //dielectric constant for silicon
9 epso=8.85*(10^-14)
10 eps=epso*epsR      //permittivity of silicon
11 taud=eps/sig      //dielectric relaxation time
   constant
12 tau=taud*10^12
13 printf('The dielectric relaxation time constant for
   this semiconductor is %1.2f ps',tau)
```

---

### Scilab code Exa 5.6 Quasi Energy Fermi levels

```
1 // Chapter 5_Non equilibrium excess carriers in  
    semiconductors  
2 //Caption_Q quasi Energy Fermi Levels  
3 //Ex_6//page 194  
4 T=300      //temperature in kelvin  
5 no=10^15    //carrier concentration  
6 ni=10^10    //intrinsic concentration  
7 po=10^5  
8 deln=10^13   //excess carrier concentration  
9 delp=10^13  
10 EFnEfi=0.0259*log(no/ni)      //fermi level for thermal  
    equilibrium  
11 EfnEfi=0.0259*log((no+deln)/ni)  
12 EfiEfp=0.0259*log((po+delp)/ni)  
13 printf('Quasi fermi level for electrons in non  
    equilibrium is %1.4f eV and for holes is %1.3f  
    eV ',EfnEfi,EfiEfp)
```

---

### Scilab code Exa 5.10 Surface effects

```
1 // Chapter 5_Non equilibrium excess carriers in  
    semiconductors  
2 //Caption_Surface effects  
3 //Ex_10//page 206  
4 gtaupo=10^14  
5 Dp=10  
6 Lp=31.6*(10^-4)  
7 delpo=10^13  
8 s=(Dp/Lp)*((gtaupo/delpo)-1)  
9 printf('Surface recombination velocity is %1.2fd cm/  
    s ',s)
```

---

# Chapter 6

## The pn junction

**Scilab code Exa 6.1** Zero applied bias

```
1 // Chapter 6_The pn junction
2 //Caption_Zero applied bias
3 //Ex_1//page 220
4 Na=10^18      //acceptor ion concentration
5 T=300        //temperature in kelvin
6 Nd=10^15
7 ni=1.5*(10^10)    //intrinsic ion concentration
8 Vbi=(0.0259)*log(Na*Nd/(ni^2))
9 printf('The built in potential barrier is %1.3f V', Vbi)
```

---

**Scilab code Exa 6.2** Space charge width

```
1 // Chapter 6_The pn junction
2 //Caption_Space charge width
3 //Ex_2//page 224
4 T=300
5 Na=10^16      //acceptor ion concentration
```

```

6 Nd=10^15      //donor ion concentration
7 eps=11.7*8.85*(10^-14)
8 e=1.6*(10^-19)
9 Vbi=0.635    //built in potential barrier
10 W=(2*eps*Vbi/e*(Na+Nd)/(Na*Nd))^0.5
11 Emax=-e*Nd*W/eps
12 printf('The space charge width is %f cm and the
          electric field is %f V/cm',W,Emax)

```

---

### Scilab code Exa 6.3 Space charge width

```

1 // Chapter 6_The pn junction
2 //Caption_Space charge width
3 //Ex_3//page 227
4 Na=10^16    //acceptor ion concentration
5 T=300      //temperature in kelvin
6 Nd=10^15
7 ni=1.5*(10^10)  //intrinsic ion concentration
8 Vr=5       //Reverse applied voltage
9 Vbi=0.635
10 V=Vr+Vbi
11 W=(2*eps*V/e*(Na+Nd)/(Na*Nd))^0.5
12 printf('The space charge width is %f cm ',W)

```

---

### Scilab code Exa 6.4 Space charge width

```

1 // Chapter 6_The pn junction
2 //Caption_Space charge width
3 //Ex_4//page 228
4 T=300
5 Na=10^18    //acceptor ion concentration
6 Emax=3*10^5  //Max electric field
7 Vr=25      //Reverse bias voltage

```

```
8 eps=11.7*8.85*(10^-14)
9 e=1.6*(10^-19)
10 x=eps*(Emax^2)/(2*e*Vr);
11 Nd=Na*x/(Na-x)
12 printf('The ntype doping concentration such that the
           maximum electric field is obtained is %f /cm^3',
           Nd)
```

---

### Scilab code Exa 6.5 Junction capacitance

```
1 // Chapter 6_The pn junction
2 //Caption_Junction capacitance
3 //Ex_5//page 230
4 Na=10^16 //acceptor ion concentration
5 T=300 //temperature in kelvin
6 Nd=10^15
7 ni=1.5*(10^10) //intrinsic ion concentration
8 Vr=5 //Reverse applied voltage
9 Vbi=0.635
10 V=Vr+Vbi
11 C=(e*eps*Na*Nd/(2*(V)*(Na+Nd)))^0.5
12 A=10^-4 //Area of the pn junction
13 Ca=A*C*10^12
14 printf('The junction capacitance for the given
           semiconductor is %1.3 f pF',Ca)
```

---

### Scilab code Exa 6.6 Junction capacitance

```
1 // Chapter 6_The pn junction
2 //Caption_Junction capacitance
3 //Ex_6//page 232
4 T=300 //temperature in kelvin
```

```
5 S=1.32*10^15      //slope of the curve between Vr and  
6 (1/c)^2  
7 ni=1.5*(10^10)  
8 Vbi=0.855      //Built in potential barrier  
9 eps=11.7*8.85*(10^-14)  
10 e=1.6*(10^-19)  
11 Nd=2/(e*eps*S)  
12 Na=((ni^2)/Nd)*exp(Vbi/0.0259)  
12 printf('The impurity doping concentration in p+n  
junction is -acceptor %f cm^-3 and donor %f cm^-3  
' ,Na ,Nd)
```

---

# Chapter 7

## The pn junction diode

Scilab code Exa 7.1 pn junction current

```
1 // Chapter 7_The pn junction Diode
2 //Caption_pn Junction current
3 //Ex_1//page 252
4 T=300      //temperature in kelvin
5 ni=1.5*(10^10)    //intrinsic ion concentration
6 Nd=10^16
7 Vf=0.60        //forward bias voltage
8 pno=(ni^2)/Nd
9 e=1.6*10^-19
10 pn=pno*exp(Vf/0.0259)
11 printf('Minority carrier hole concentration is %f cm
           ^-3',pn)
```

---

Scilab code Exa 7.2 pn junction current

```
1 // Chapter 7_The pn junction Diode
2 //Caption_pn Junction current
3 //Ex_2//page 258
```

```

4 Na=10^16      // acceptor impurity
5 Nd=10^16      // donor impurity
6 ni=1.5*10^10   // intrinsic concentration
7 Dn=25
8 Dp=10
9 tau_po=5*10^-7
10 tau_no=5*10^-7
11 epsr=11.7
12 Js=e*ni^2*((1/Na*(Dn/tau_no)^0.5)+(1/Nd)*(Dp/tau_po)
    ^0.5)*10^9
13 printf('Ideal reverse saturation current density is
    %f nA/cm', Js)

```

---

### Scilab code Exa 7.3 pn junction current

```

1 // Chapter 7_The pn junction Diode
2 //Caption_pn Junction current
3 //Ex_3//page 258
4 Jn=20      //electron current density
5 Jp=5       //hole current density
6 T=300
7 Va=0.65
8 ni=1.5*10^10   //intrinsic concentration
9 Dn=25
10 Dp=10
11 e=1.6*10^-19
12 tau_po=5*10^-7
13 tau_no=5*10^-7
14 epsr=11.7
15 Na=1/(Jn/((e*(Dn/tau_no)^0.5)*(ni^2*(exp(Va/0.0259)
    -1))))
16 Nd=1/(Jp/((e*(Dp/tau_po)^0.5)*(ni^2*(exp(Va/0.0259)
    -1))))
17 printf('The design parameters for this semiconductor
    are Na=%f cm^-3 and Nd=%f cm^-3',Na,Nd)

```

---

### Scilab code Exa 7.4 pn junction current

```
1 // Chapter 7_The pn junction Diode
2 //Caption_pn Junction current
3 //Ex_4//page 261
4 T=300
5 Va=0.65
6 Js=4.15*10^-11
7 e=1.6*10^-19
8 J=Js*(exp(Va/0.0259)-1)
9 mun=1350
10 Nd=10^16
11
12 E=J/(e*mun*Nd)
13 printf('The electric field required to produce a
given majority carrier drift is %f V/cm',E)
```

---

### Scilab code Exa 7.5 Temperature effects on pn junction

```
1 // Chapter 7_The pn junction Diode
2 //Caption_Temperature effects on pn Junction current
3 //Ex_5//page 262
4 T2=310
5 T1=300
6 Eg=1.12      //energy band gap
7 Va1=0.60      //forward bias voltage at 300k
8 Va2=Eg-((T2/T1)*(Eg-Va1))
9 delV=Va2-Va1
10 printf('Change in fprward bias voltage on a pn
junction with the change in temperature is %f V',
delV)
```

---

### Scilab code Exa 7.6 small signal admittance

```
1 // Chapter 7_The pn junction Diode
2 //Caption_Small signal admittance
3 //Ex_6//page 272
4 T=300
5 tau_po=10^-7
6 Ipo=0.001
7 Idq=0.001
8 Cd=10^9*(1/(2*0.0259))*(Ipo*tau_po)
9 rd=0.0259/(Idq)      //diffusion resistance
10 printf('Diffusion capacitance is %1.2f nF and
           diffusion resistance is %1.2f ohm',Cd,rd)
```

---

### Scilab code Exa 7.7 Generation recombination currents

```
1 // Chapter 7_The pn junction Diode
2 //Caption_Generation Recombination currents
3 //Ex_7//page 277
4 T=300
5 tau_o=5*10^-7
6 tau_po=5*10^-7
7 tau_no=5*10^-7
8 Na=10^16      //acceptor impurity
9 Nd=10^16      // donor impurity
10 ni=1.5*10^10    //intrinsic concentration
11 epsr=11.7
12 eps=epsr*8.85*10^-14
13 V=5          //V=Vbi+VR
14 e=1.6*10^-19
15 W=((2*eps/e)*((Na+Nd)/(Na*Nd))*(V))^0.5
16 Jgen=e*ni*10^9*W/(2*tau_o)
```

```
17 printf('The ideal reverse saturation current density  
was calculated in example 2 and it was  
4.15810^-11 A/cm^2 and the generation current  
density calculated here is %f nA/cm^2 ', Jgen)
```

---

# Chapter 8

## Metal semiconductors and semiconductor heterojunctions

Scilab code Exa 8.1 Shottky barrier diode

```
1 // Chapter 8_Metal Semiconductor and Semiconductor  
    heterojunctions  
2 //Caption_Shottky barrier diode  
3 //Ex_1//page 308  
4 T=300      //temperature in kelvin  
5 Nd=10^16    //donor impurity  
6 phi_m=4.55  //metal work function for tungsten  
7 xi=4.01     //electron affinity for silicon  
8 phi_bo=phi_m-xi  
9 phi_n=0.0259*log(2.8*10^19/Nd)  
10 Vbi=phi_bo-phi_n  
11 xn=(2*eps*Vbi/(e*Nd))^(1/2)      //space charge width  
    at zero bias  
12 Emax=e*Nd*xn/eps      //maximum electric field  
13 printf('Theoretical barrier height is %f V, built-in  
        potential barrier is %f V and maximum electric  
        field is %f V/cm', phi_bo,phi_n,Emax)
```

---

### Scilab code Exa 8.2 Non ideal effects on barrier height

```
1 // Chapter 8_Metal Semiconductor and Semiconductor  
heterojunctions  
2 //Caption_Non ideal effects on the barrier height  
3 //Ex_3//page 312  
4 E=6.8*10^4  
5 T=300  
6 e=1.6*10^-19  
7 eps=13.1*8.85*10^-14  
8 delphi=(e*E/(4*pi*eps))^.5  
9 xm=(e/(16*pi*eps*E))^.5*10^8  
10 printf('Position of the maximum barrier height is %1  
.0 f Angstrom ',xm)
```

---

### Scilab code Exa 8.3 Non ideal effects on barrier height

```
1 // Chapter 8_Metal Semiconductor and Semiconductor  
heterojunctions  
2 //Caption_Non ideal effects on the barrier height  
3 //Ex_3//page 312  
4 E=6.8*10^4  
5 T=300  
6 e=1.6*10^-19  
7 eps=13.1*8.85*10^-14  
8 delphi=(e*E/(4*pi*eps))^.5  
9 xm=(e/(16*pi*eps*E))^.5*(10^8)  
10 printf('Position of maximum barrier height is %fA ',  
xm)
```

---

### Scilab code Exa 8.4 Current voltage relationship

```
1 // Chapter 8_Metal Semiconductor and Semiconductor  
heterojunctions  
2 //Caption_Current voltage relationship  
3 //Ex_4/page 318  
4 phi_bn=0.67      //barrier height  
5 Jst=6*10^-5      //reverse saturation current density  
6 T=300  
7 e=1.6*10^-19  
8 A=Jst/(T^2)*exp(phi_bn/0.0259)  
9 printf('The effective Richardson constant is %1.0f A  
/K^2-cm^2',A)
```

---

### Scilab code Exa 8.5 Comparison of the schottky barrier diode and the pn junction diode

```
1 // Chapter 8_Metal Semiconductor and Semiconductor  
heterojunctions  
2 //Caption_Comparison of the schottky barrier diode  
and the pn junction diode  
3 //Ex_5/page 319  
4 e_phi_bn=0.67  
5 A=114      //effective richardson constant  
6 T=300  
7 Jst=A*T^2*exp(-e_phi_bn/0.0259)  
8 //if we neglect the barrier lowering effect , we have  
// for the schottky barrier diode  
9 //for a pn junction  
10 Na=10^18  
11 Nd=10^16  
12 Dp=10  
13 Dn=25  
14 tau_po=10^-7  
15 tau_no=10^-7
```

```

16 Lp=(Dp*tau_po)^0.5
17 Ln=(Dn*tau_no)^0.5
18 pno=2.25*10^4
19 npo=2.25*10^2
20 //the ideal reverse saturation current density of
   the pn junction diode can be determined as
21 Js=e*Dn*npo/Ln+(e*Dp*pno/Lp)
22 J=10^9*(Js+5.7*10^-13)
23 printf('Reverse saturation current density for
   schottky barrier diode is %f A/cm^2 and for pn
   junction is %f nA/cm^2 ',Jst,J)

```

---

### Scilab code Exa 8.6 Schottky barrier diode and pn junction

```

1 // Chapter 8_Metal Semiconductor and Semiconductor
   heterojunctions
2 //Caption_Schottky barrier diode and pn junction
3 //Ex_6//page 308
4 Jf=10 //forward biased current density
5 Jst=5.98*10^-5
6 Va=(0.0259*log(Jf/Jst))
7 //for pn junction diode
8 Js=3.66*10^-11 //reverse saturation current
   density
9 Va_pn=0.0259*log(Jf/Js)
10 printf('Forward bised voltage required for schottky
   is %1.3f V and for pn junction is %1.3fV ',Va,
   Va_pn)

```

---

### Scilab code Exa 8.7 Tunnelling barrier

```

1 // Chapter 8_Metal Semiconductor and Semiconductor
   heterojunctions

```

```

2 //Caption_Tunnelling barrier
3 //Ex_7//page 325
4 T=300 //temperature in kelvin
5 Nd=7*10^18 //donor impurity
6 e=1.6*10^-19
7 phi_bn=0.67 //schottky barrier
8 Vbi=phi_bn //assumption with neglection of
    barrier lowering effect
9 eps=11.7*8.85*10^-14
10 xn=(2*eps*Vbi/(e*Nd))^0.5
11 x=10^8*xn
12 printf('The space charge width is %1.0f Angstrom',x)

```

---

### Scilab code Exa 8.8 Equilibrium electrostatics

```

1 // Chapter 8_Metal Semiconductor and Semiconductor
    heterojunctions
2 //Caption_Equilibrium electrostatics
3 //Ex_8//page 333
4 Nd=10^16 //donor impurity
5 Na=10^16 //acceptor impurity
6 ni=2.4*10^13 //intrinsic ion concentration
7 T=300
8 e=1.6*10^-19
9 xi_n=4.13 //electron affinity
10 xi_p=4.07
11 del_Ec=(xi_n-xi_p) //difference between two
    conduction band energies
12 del_Eg=1.43-0.67
13
14 del_Ev=del_Eg-del_Ec //difference between two
    valence band energies
15 pno=ni^2/Nd
16 Ncp=6*10^18
17 Ncn=7*10^18

```

```
18 Vbi=del_Ev+(0.0259*log(Na*Ncp/(pno*Ncn)))
19 printf('Difference between two conduction band
    energies is %1.2f eV , difference between two
    valence band energies is %f eV and Vbi=%fV',
    del_Ec,del_Ev,Vbi)
```

---

# Chapter 9

## The Bipolar transistor

**Scilab code Exa 9.1** Gain factors

```
1 // Chapter 9_The bipolar transistor
2 //Caption_Gain factors
3 //Ex_1//page 371
4 gamma=0.9967      //emitter injection efficiency
                      factor
5 Nbe=(1/gamma)-1    //base to emitter doping
                      concentration ratio
6 NEB=1/Nbe
7 printf('The emitter doping concentration must be
          much larger i.e.%1.0ftimes that of base',NEB)
```

---

**Scilab code Exa 9.2** Gain factors

```
1 // Chapter 9_The bipolar transistor
2 //Caption_Gain factors
3 //Ex_2//page 372
4 alpha_T=0.9967
5 Db=10
```

```
6 tau_bo=10^-7
7 xbLb=abs(acosh(alpha_T)) //xB/Lb where LB is the
    length
8 Lb=(Db*tau_bo)^0.5
9 xb=xbLb*Lb*10^4
10 printf('Base width required to achieve the given
        base transport factor is %1.3f micrometer',xb)
```

---

### Scilab code Exa 9.3 Gain factors

```
1 // Chapter 9_The bipolar transistor
2 //Caption_Gain factors
3 //Ex_3//page 373
4 delta=0.9967      // recombination favtor
5 T=300
6 Jro=10^-8
7 Jso=10^-11
8 del=1/delta-1
9 x=del*Jso/Jro
10 Vbe=-2*0.0259*log(x)
11 printf('Forward biased BE voltage required to
        achieve the given delta is %1.3f V',Vbe)
```

---

### Scilab code Exa 9.4 Gain factors

```
1 // Chapter 9_The bipolar transistor
2 //Caption_Gain factors
3 //Ex_4//page 373
4 DE=10
5 DB=25
6 XB=0.70*10^-4 //width of base
7 XE=0.50*10^-4 //width of emitter
8 NE=10^18 //doping concentration in emitter
```

```

9 NB=10^16      //doping concentration in base
10 VBE=0.65
11 e=1.6*10^-19
12 tau_eo=10^-7    //minority carrier lifetime in
13          emitter
14 tau_bo=5*10^-7    //minority carrier lifetime in base
15 Jro=5*10^-8
16 T=300
17 peo=(1.5*10^10)^2/NE
18 nbo=(1.5*10^10)^2/NB
19 Le=(DE*tau_eo)^0.5
20 Lb=(DB*tau_bo)^0.5
21 gamma_i=1/(1+((peo*DE*Lb*tanh(0.0198))/(nbo*DB*Le*
22          tanh(0.050))))
23 alpha_T=1/(cosh(XB/Lb))
24 Jso=e*DB*nbo/(Lb*tanh(XB/Lb))
25 delta=1/1+(Jro*exp(-VBE/(2*0.0259)/Jso))
26 delta=0.99986
27 alpha=gamma_i*alpha_T*delta
28 beta_i=alpha/(1-alpha)
29 printf('Common emitter current gain is %1.0f',beta_i
)

```

---

### Scilab code Exa 9.5 Non ideal effects

```

1 // Chapter 9_The bipolar transistor
2 //Caption_Non ideal effects
3 //Ex_5//page 376
4 T=300
5 e=1.6*10^-19
6 NB=5*10^16      //doping concentration in base
7 NC=2*10^15      //doping concentration in collector
8 XB=0.70*10^-4    //metallurgical base width
9 ni=1.5*10^10     //intrinsic ion concentration
10 Vbi=0.718       //built-in potential

```

```

11 Vcb1=2
12 eps=11.7*8.85*10^-14
13 xdb1=(2*eps*(Vbi+Vcb1)*NC/(e*NB*(NC+NB)))^0.5
14 Vcb2=10
15 xdb2=(2*eps*(Vbi+Vcb2)*NC/(e*NB*(NC+NB)))^0.5
16 xb1=XB-xdb1      //neutral base width
17 xb2=XB-xdb2
18 del_xb=((xb1-xb2)/xb1)*100
19 printf('The neutral base width changes by %1.0f
           percent as the C-B voltage changes from 2 to 10 V
           ',del_xb)

```

---

### Scilab code Exa 9.6 Non ideal effects

```

1 // Chapter 9_The bipolar transistor
2 //Caption_Non ideal effects
3 //Ex_6//page 377
4 DB=25
5 VBE=0.60
6 T=300
7 e=1.6*10^-19
8 NB=5*10^16      //doping concentration in base
9 NC=2*10^15      //doping concentration in collector
10 XB=0.70*10^-4   //mettulurgical base width
11 ni=1.5*10^10    //intrinsic ion concentration
12 nB0=ni^2/NB
13 xb1=0.648*10^-4
14 Jc1=e*DB*nB0*exp(VBE/0.0259)/xb1
15 xb2=0.597*10^-4
16 Jc2=e*DB*nB0*exp(VBE/0.0259)/xb2
17 VCE1=2.6
18 VCE2=10.6
19 del_JC_VCE=(Jc2-Jc1)/(VCE2-VCE1)
20 Va=3.20/del_JC_VCE-2.6
21 printf('The early voltage is %1.0f V',Va)

```

---

### Scilab code Exa 9.7 Non ideal effects

```
1 // Chapter 9_The bipolar transistor
2 //Caption_Non ideal effects
3 //Ex_7//page 382
4 T=300
5 NE1=10^18    //emitter doping
6 NE2=10^19
7 ni=1.5*10^10      //intrinsic ion concentration
8 pE01=ni^2/NE1
9 pE02=ni^2/NE2
10 //This we did by neglecting bandgap narrowing , if
    we consider it , we get
11 pE011=pE01*exp(0.030/0.0259)
12 pE021=pE02*exp(0.1/0.0259)
13 printf('The thermal equilibrium minority carrier
    concentration increases by a factor of 1.5
    instead of decreasing by a factor of 9.This
    effect is due to bandgap narrowing')
```

---

### Scilab code Exa 9.8 Breakdown voltage

```
1 // Chapter 9_The bipolar transistor
2 //Caption_Breakdown voltage
3 //Ex_8//page 387
4 Wb=0.5*10^-4    //metallurgical base width
5 NB=10^16
6 eps=11.7*8.85*10^-14
7 e=1.6*10^-19
8 Vpt=25      //punch through voltage
9 x=Vpt*2*eps/(e*Wb^2*Nb)
```

```

10 y=x-1
11 NC=NB/y
12 xn=(2*eps*(Vpt)*NB/(e*NC*(NB+NC)))^0.5*10000
13 printf('The collector doping is %1.2f per cm^3 and
           collector widt is %1.2f micrometer',NC,xn)

```

---

### Scilab code Exa 9.9 Breakdown voltage

```

1 // Chapter 9_The bipolar transistor
2 //Caption_Breakdown voltage
3 //Ex_9//page 390
4 bet=100    //common emitter current gain
5 NB=10^17   //base doping concentration
6 vmin=15    //minimum open base breakdown voltage
7 BVcbo=(bet)^(1/3)*vmin
8 printf('To achieve this breakdown voltage , the
           maximum collector doping concentration should be
           7*10^15 cm^-3 from the figure')

```

---

### Scilab code Exa 9.10 Ebers moll model

```

1 // Chapter 9_The bipolar transistor
2 //Caption_Ebers Moll model
3 //Ex_10//page 394
4 T=300
5 alpha_f=0.99
6 alpha_r=0.20
7 Ic=.001
8 Ib=50*10^-6
9 Vt=0.0259
10 x=Ic*(1-alpha_r)+Ib
11 y=alpha_f*Ib-((1-alpha_f)*Ic)
12 z=alpha_f/alpha_r

```

```
13 VCEsat=Vt*log(x*z/y)
14 printf('The collector emitter saturation voltage is
    %1.3f V',VCEsat)
```

---

### Scilab code Exa 9.12 Transistor cut off frequency

```
1 // Chapter 9_The bipolar transistor
2 //Caption_Transistor cut off frequency
3 //Ex_12//page 403
4 Ie=0.001      //emitter current
5 Cje=10^-12
6 xb=0.5*10^-4
7 vs=10^7
8 Dn=25
9 xdc=2.4*10^-4
10 rc=20
11 Cu=0.1*10^-12      //B-C junction capacitance
12 Cs=0.1*10^-12      //collector to substrate
    capacitance
13 re=0.0259/Ie
14 tau_e=re*Cje*10^12 //emitter base junction
    charging time
15 tau_b=(xb^2)/(2*Dn) *10^12      //base transit time
16 tau=xdc/vs*10^12
17 tau_c=rc*(Cu+Cs)*10^12
18 tau_ec=(tau_e+tau_b+tau+tau_c) //total emitter to
    collector time delay
19 fT=(10^3)/(2*pi*tau_ec)
20 bet=100
21 fB=fT/bet      //beta cutoff frequency
22 printf('Emitter to collector transit time is %1.1f
    psec and cut off frequency is %1.2f GHz',tau_ec,
    fT)
```

---

# Chapter 10

## Fundamentals of the Metal Oxide semiconductor Field Effect Transistor

Scilab code Exa 10.1 The two terminal MOS structure

```
1 // Chapter 10_Fundamentals of the Metal Oxide
   Semiconductor Field Effect Transistor
2 //Caption_The two terminal MOS structure
3 //Ex_1//page 434
4 Na=10^16
5 T=300
6 eps=11.7*8.85*10^-14
7 e=1.6*10^-19
8 ni=1.5*10^10      //intrinsic carrier concentration
9 phi_fp=0.0259*log(Na/ni)
10 xdT=10^4*(4*eps*phi_fp/(e*Na))^.5
11 printf('The maximum space charge width is %1.2 f
           micrometer', xdT)
```

---

### Scilab code Exa 10.2 Work function

```
1 // Chapter 10_Fundamentals of the Metal Oxide  
    Semiconductor Field Effect Transistor  
2 //Caption_Work function  
3 //Ex_2//page 437  
4 phi_m=3.2      //work function for Al-Si junction  
5 xi=3.25       //oxide electron affinity  
6 Eg=1.11  
7 ni=1.5*10^10    //intrinsic carrier concentration  
8 Na=10^14  
9 phi_fp=0.0259*log(Na/ni)  
10 phi_ms=phi_m-(xi+Eg/(2)+phi_fp)  
11 printf('Metal semiconductor work function difference  
is %1.2 f V',phi_ms)
```

---

### Scilab code Exa 10.3 Flat band voltage

```
1 // Chapter 10_Fundamentals of the Metal Oxide  
    Semiconductor Field Effect Transistor  
2 //Caption_Flat band voltage  
3 //Ex_3//page 442  
4 Na=10^16  
5 tox=500*10^-8      //oxide thickness  
6 Qss=10^11          //trapped charge per unit area  
7 e=1.6*10^-19  
8 eps_ox=3.9*8.85*10^-14  
9 Cox=eps_ox/tox  
10 QSS=Qss*e  
11 phi_ms=-1.1  
12 Vfb=phi_ms-(QSS/Cox)  
13 printf('Flat band voltage for this MOS capacitor is  
%1.2 f V',Vfb)
```

---

### Scilab code Exa 10.4 Flat band voltage

```
1 // Chapter 10_Fundamentals of the Metal Oxide  
    Semiconductor Field Effect Transistor  
2 //Caption_Flat band voltage  
3 //Ex_4//page 445  
4 Na=3*10^16  
5 eps=11.7*8.85*10^-14  
6 eps_ox=3.9*8.85*10^-14  
7 e=1.6*10^-19  
8 Qss=10^11  
9 Vtn=0.65  
10 ni=1.5*10^10      //intrinsic carrier concentration  
11 phi_ms=-1.13  
12 phi_fp=0.0259*log(Na/ni)  
13 xdt=(4*eps*phi_fp/(e*Na))^0.5  
14 QSD=e*Na*xdt  
15 x=Vtn-phi_ms-2*phi_fp  
16 y=(QSD-Qss*e)/eps_ox  
17 z=x/y*10^8  
18 printf('The oxide thickness of this MOS system is %1  
.0 f angstrom ',z)
```

---

### Scilab code Exa 10.5 Threshold voltage

```
1 // Chapter 10_Fundamentals of the Metal Oxide  
    Semiconductor Field Effect Transistor  
2 //Caption_Threshold voltage voltage  
3 //Ex_5//page 446  
4 T=300  
5 eps=11.7*8.85*10^-14  
6 eps_ox=3.9*8.85*10^-14
```

```

7 e=1.6*10^-19
8 ni=1.5*10^10      //intrinsic carrier concentration
9 Na=10^14          //acceptor impurity concentration
10 Qss=10^10
11 tox=500*10^-8     //oxide thickness
12 phi_ms=-0.83
13 phi_fp=0.0259*log(Na/ni)
14 xdt=(4*eps*phi_fp/(e*Na))^0.5
15 QSD_MAX=e*Na*xdt;
16 Vtn=(QSD_MAX-Qss*e)*(tox/eps_ox)+phi_ms+2*phi_fp
17 printf('The threshold voltage of this MOS system is
%1.3f V',Vtn)

```

---

### Scilab code Exa 10.6 Threshold voltage

```

1 // Chapter 10_Fundamentals of the Metal Oxide
   Semiconductor Field Effect Transistor
2 //Caption_Threshold voltage voltage
3 //Ex_6//page 448
4 tox=650*10^-8
5 eps=11.7*8.85*10^-14
6 eps_ox=3.9*8.85*10^-14
7 Qss=10^10
8 Vtp=-1
9 Nd=2.5*10^14
10 ni=1.5*10^10      //intrinsic carrier concentration
11 phi_tn=0.0259*log(Nd/ni)
12 xdt=(4*eps*phi_tn/(e*Nd))^0.5
13 QSD_MAX=e*Nd*xdt;
14 phi_ms=-0.35
15 Vtp2=(-QSD_MAX-Qss*e)*(tox/eps_ox)+phi_ms-2*phi_tn
16 q=abs(Vtp2)==Vtp
17 printf('Since Vtp2=Vtp, it is essentially equal to
the desired result')

```

---

### Scilab code Exa 10.7 Capacitance Voltage characteristics

```
1 // Chapter 10_Fundamentals of the Metal Oxide  
    Semiconductor Field Effect Transistor  
2 //Caption_Capacitance Voltage characteristics  
3 //Ex_7//page 455  
4 Na=10^16  
5 tox=550*10^-8      //oxide thickness  
6 eps=11.7*8.85*10^-14  
7 eps_ox=3.9*8.85*10^-14  
8 Cox=eps_ox/tox*10^9  
9 ni=1.5*10^10      //intrinsic carrier concentration  
10 phi_fp=0.0259*log(Na/ni)  
11 xdt=(4*eps*phi_fp/(e*Na))^0.5  
12 Cmin=eps_ox/(tox+(eps_ox/eps)*xdt)*10^9  
13 r=Cmin/Cox  
14 CFB=eps_ox/(tox+(eps_ox/eps)*(0.0259*eps/(e*Na))  
    ^0.5) *10^9    //flat band capacitance  
15 r2=CFB/Cox  
16 printf('The value of oxide capacitance , minimum  
    capacitance and flat band capacitance are %1.2 f  
    nF, %1.2 f nF and %1.2 f nF respectively ',Cox,Cmin,  
    CFB)
```

---

### Scilab code Exa 10.8 Current voltage relationship

```
1 // Chapter 10_Fundamentals of the Metal Oxide  
    Semiconductor Field Effect Transistor  
2 //Caption_Current voltage relationship  
3 //Ex_8//page 473  
4 L=1.25*10^-4      //channel length  
5 mun=650      //mobility of electrons
```

```

6 Cox=6.9*10^-8 //oxide capacitance
7 Vt=0.65 //thermal voltage
8 Idsat=4*10^-3 //saturated current
9 VGS=5
10 W=2*L*Idsat/(mun*Cox*(VGS-Vt)^2)*10^4
11 printf('The width of MOSFET such that the specified
           current is induced is %1.1f micrometer',W)

```

---

### Scilab code Exa 10.9 Threshold voltage

```

1 // Chapter 10_Fundamentals of the Metal Oxide
   Semiconductor Field Effect Transistor
2 //Caption_Current voltage relationship
3 //Ex_9//page 474
4 W=15*10^-4 //Width of MosFET
5 L=2*10^-4 //length of MOSFET
6 COX=6.9*10^-8 //oxide capacitance
7 VDS=0.10
8 ID1=35*10^-6 //DRAIN CURRENT
9 VGS1=1.5
10 ID2=75*10^-6
11 VGS2=2.5
12 mun=L*(ID2-ID1)/(W*COX*(VGS2-VGS1)*VDS)
13 printf('The inversion carrier mobility is %1.0f cm
           ^2/V-s ',mun)

```

---

### Scilab code Exa 10.10 Substrate bias effects

```

1 // Chapter 10_Fundamentals of the Metal Oxide
   Semiconductor Field Effect Transistor
2 //Caption_Substrate bias effects
3 //Ex_10//page 478
4 T=300

```

```

5 Na=3*10^16
6 tox=500*10^-8
7 VSB=1
8 ni=1.5*10^10      //intrinsic carrier concentration
9 phi_fp=0.0259*log(Na/ni)
10 eps_ox=3.9*8.85*10^-14
11 Cox=eps_ox/tox
12 delVT=(2*e*eps*Na)^0.5*((2*phi_fp+VSB)^0.5-(2*phi_fp
    )^0.5)/Cox
13 printf('The change in threshold voltage is %1.2fV' ,
    delVT)

```

---

### Scilab code Exa 10.11 Cut off frequency

```

1 // Chapter 10_Fundamentals of the Metal Oxide
   Semiconductor Field Effect Transistor
2 //Caption_Cut off frequency
3 //Ex_11//page 484
4 mun=400      //mobility
5 L=4*10^-4
6 VT=1
7 VGS=3
8 fT=mun*(VGS-VT)/(2*pi*L^2)*10^-6
9 printf('The cut off frequency of this MOSFET with
    constant mobility is %1.0f MHz',fT)

```

---

# Chapter 11

## Metal semiconductors and semiconductor heterojunctions Additional concepts

Scilab code Exa 11.1 Mobility variation

```
1 // Chapter 11_ Metal–Oxide–Semiconductor Field
   Effect Transistor: Additional Concepts
2 //Caption_Mobility variation
3 //Ex_1//page 509
4 T=300
5 Na=3*10^16
6 ni=1.5*10^10      //intrinsic carrier concentration
7 phi_fp=0.0259*log(Na/ni)
8 xdt=(4*eps*phi_fp/(e*Na))^.5
9 QSD_MAX=e*Na*xdt;
10 Eeff=1/eps*QSD_MAX
11 printf('Effective electric field at threshold is %1
.2 f V/cm' ,Eeff)
```

---

### Scilab code Exa 11.2 Mobility variation

```
1 // Chapter 11_ Metal–Oxide–Semiconductor Field  
Effect Transistor: Additional Concepts  
2 //Caption_Mobility variation  
3 //Ex_2//page 517  
4 Na=3*10^16  
5 tox=450*10^-8  
6 eps=11.7*8.85*10^-14  
7 e=1.6*10^-19  
8 eps_ox=3.9*8.85*10^-14  
9 ni=1.5*10^10      //intrinsic carrier concentration  
10 L=1.25*10^-4  
11 rj=0.5*10^-4  
12 Cox=eps_ox/tox    //oxide capacitance  
13 phi_fp=0.0259*log(Na/ni)  
14 xdt=(4*eps*phi_fp/(e*Na))^0.5  
15 x=e*Na*xdt/Cox  
16 y=(1+(2*xdt/rj))^0.5-1  
17 delVt=-x*(rj*y/L)      //voltage shift  
18  
19 printf('Threshold voltage shift due to short channel  
effects is %1.3f V',delVt)
```

---

### Scilab code Exa 11.3 narrow channel effects

```
1 // Chapter 11_ Metal–Oxide–Semiconductor Field  
Effect Transistor: Additional Concepts  
2 //Caption_Narrow channel effects  
3 //Ex_3//page 520  
4 Na=3*10^16  
5 tox=450*10^-8      //oxide thickness  
6 fi=%pi/2        //fitting parameter  
7 delVt=0.2  
8 Cox=7.67*10^-8    //oxide capacitance
```

---

```

9 xdt=0.18*10^-4
10 W=10^4*e*Na*(fi*xdt^2)/(Cox*delVt)
11 printf('The channel width that will limit the
           threshold voltage is %1.2f micrometer',W)

```

---

### Scilab code Exa 11.4 Breakdown voltage

```

1 // Chapter 11_ Metal–Oxide–Semiconductor Field
   Effect Transistor: Additional Concepts
2 //Caption_Breakdown voltage
3 //Ex_4//page 527
4 Nd=10^19      //donor concentration
5 Na=10^16      //acceptor concentration
6 L=1.2*10^-4   //channel length
7 ni=1.5*10^10  //intrinsic carrier concentration
8 Vbi=0.0259*log(Na*Nd/ni^2)
9 xdo=(2*eps*Vbi/(e*Na))^.5    //zero biased source-
   substrate pn junction width
10 //xd=(2*eps*(VbiVDS)/(e*Na))^.5    //reverse biased
    drain substrate pn junction width
11 xd=L-xdo   //at punch through
12 VbiVDS=xd^2*e*Na/(2*eps)      //Vbi+VDS
13 VDS=VbiVDS-Vbi
14 printf('The punch through voltage is %1.1f V',VDS)

```

---

### Scilab code Exa 11.5 Lightly doped drain transistor

```

1 // Chapter 11_ Metal–Oxide–Semiconductor Field
   Effect Transistor: Additional Concepts
2 //Caption_Lightly doped drain transistor
3 //Ex_5//page 531
4 tox=500*10^-8
5 VFB0=-1.25      //initial flat band voltage

```

```

6 e=1.6*10^-19
7 eps_ox=3.9*8.85*10^-14
8 ni=1.5*10^10      //intrinsic carrier concentration
9 VT=0.70
10 Na=5*10^15
11 phi_fpo=0.0259*log(Na/ni)
12 xdto=(4*eps*phi_fpo/(e*Na))^0.5
13 Cox=eps_ox/tox
14 VTO=VFB0+2*phi_fpo+(e*Na*xdto)/Cox
15 x=VT-VTO
16 Dt=Cox*x/e      //implant dose
17 xt=0.15*10^-4    //depth to which uniform implant
                     extends
18 Nsa=Dt/xt
19 Ns=Nsa+Na
20 printf('The required implant dose to achieve the
           desired threshold voltage is %1.2f per cm^2 ',Dt)

```

---

### Scilab code Exa 11.6 Radiation and hot electron effect

```

1 // Chapter 11 Metal–Oxide–Semiconductor Field
   Effect Transistor: Additional Concepts
2 //Caption_Radition and hot electron effect
3 //Ex_6//page 535
4 tox=500*10^-8 //oxide thickness
5 p=0.2 //20% are trapped at oxide semiconductor
         surface
6 N=10^18 //electron hole pair
7 e=1.6*10^-19
8 eps_ox=3.9*8.85*10^-14
9 ni=1.5*10^10 //intrinsic carrier concentration
10 Nh=N*tox //areal density of holes
11 Qss=Nh*p //trapped surface charge density
12 Cox=eps_ox/tox
13 delVt=-Qss*e/Cox

```

```
14 printf('The threshold voltage shift due to radiation  
induced oxide charge trapping is %1.2f V',delVt)
```

---

# Chapter 12

## The junction field effect transistor

Scilab code Exa 12.1 Device characteristics

```
1 // Chapter 12_The junction field effect transistor
2 //Caption_Device characteristics
3 //Ex_1//page 557
4 T=300
5 Na=10^18
6 e=1.6*10^-19
7 eps=8.85*10^-14*11.7
8 ni=1.5*10^10
9 Nd=10^16          //donor concentration
10 a=0.75*10^-4      //metallurgical channel thichness
11 Vpo=e*a^2*Nd/(2*eps)      //internal pinch off
    voltage
12 Vbi=0.0259*log(Na*Nd/ni^2)      //built in potential
    barrier
13 Vp=Vbi-Vpo      //pinch off voltage
14 printf('The pinch off voltage of this n-channel JFET
    is %1.2fV ',Vp)
```

---

### Scilab code Exa 12.2 Device characteristics

```
1 // Chapter 12_The junction field effect transistor
2 //Caption_Device characteristics
3 //Ex_2//page 558
4 T=300
5 Nd=10^18
6 Na=2*10^16
7 e=1.6*10^-19
8 eps=8.85*10^-14*11.7
9 ni=1.5*10^10
10 Vp=2.25      //pinchoff voltage
11 Vbi=0.0259*log(Na*Nd/ni^2)
12 Vpo=Vp+Vbi
13 a=(2*eps*Vpo/(e*Na))^.5*10^4
14 printf('Metallurgical channel thickness is %1.3 f
micrometer',a)
```

---

### Scilab code Exa 12.3 Depletion mode JFET

```
1 // Chapter 12_The junction field effect transistor
2 //Caption_Depletion mode JFET
3 //Ex_3//page 558
4 T=300
5 Na=10^18
6 e=1.6*10^-19
7 eps=8.85*10^-14*11.7
8 Vbi=0.814
9 Vpo=4.35
10 ni=1.5*10^10
11 Nd=10^16
12 a=0.75*10^-4      //metallurgical channel thickness
```

```

13 L=10*10^-4      //channel length
14 W=30*10^-4      //channel width
15 mun=1000
16 Ipi=10^3*mun*(e*Nd)^2*W*a^3/(6*eps*L)
17 IDmax=Ipi*(1-3*(Vbi/Vpo)*(1-(2/3)*(Vbi/Vpo)^0.5))
18 printf('The maximum current is %1.3f mA while pinch
          off current is %1.3fmA',IDmax,Ipi)

```

---

### Scilab code Exa 12.4 Transconductance

```

1 // Chapter 12_The junction field effect transistor
2 //Caption_Transconductance
3 //Ex_4//page 566
4 Ipi=0.522*10^-3
5 Vbi=0.814    //built in potential barrier
6 Vpo=4.35     //pinch off
7 VGS=0
8 gms_max=10^3*3*Ipi*(1-(Vbi/Vpo)^0.5)/Vpo
9 printf('The maximum transconductance is %1.3f mA/V' ,
        gms_max)

```

---

### Scilab code Exa 12.5 The MESFET

```

1 // Chapter 12_The junction field effect transistor
2 //Caption_The MESFET
3 //Ex_5//page 567
4 Nc=4.7*10^17
5 e=1.6*10^-19
6 eps=8.85*10^-14*13.1
7 T=300
8 phi_bn=0.89      //barrier height
9 Nd=2*10^15
10 Vt=0.25

```

```

11 phi_n=0.0259*log(Nc/Nd)
12 Vbi=phi_bn-phi_n //built in potential barrier
13 Vpo=Vbi-Vt
14 a=10^4*(Vpo*2*eps/(e*Nd))^0.5
15 printf('The channel thickness of GaAs is %1.3f
micrometer',a)

```

---

### Scilab code Exa 12.6 The MESFET

```

1 // Chapter 12_The junction field effect transistor
2 //Caption_The MESFET
3 //Ex_6//page 568
4 e=1.6*10^-19
5 eps=8.85*10^-14*13.1
6 T=300
7 ni=1.8*10^6
8 Na=10^18
9 Nd=3*10^15
10 a=0.70*10^-4
11 Vbi=0.0259*log(Na*Nd/ni^2)
12 Vpo=e*a^2*Nd/(2*eps) //internal pinch off
voltage
13 Vt=Vbi-Vpo //threshold voltage
14 h=0.6*10^-4
15 VGS=Vbi-(e*h^2*Nd/(2*eps))
16 printf('The forward bias voltage required in an n
channel GaAs enhancement mode pn jfet to open up
a channel is %1.2f V',VGS)

```

---

### Scilab code Exa 12.7 The MESFET

```

1 // Chapter 12_The junction field effect transistor
2 //Caption_The MESFET

```

```

3 //Ex_7//page 570
4 e=1.6*10^-19
5 eps=8.85*10^-14*13.1
6 T=300
7 ni=1.8*10^6
8 L=1.2*10^-4
9 mun=8000
10 a=0.70*10^-4
11 Idi=75*10^-6
12 VGS=0.5
13 Vt=0.24
14 kn=Idi/(VGS-Vt)^2      //conduction parameter
15 W=10^4*kn*2*a*L/(mun*eps)
16 printf('The required channel width is %1.2f
micrometer',W)

```

---

**Scilab code Exa 12.8** The MESFET channel length modulation

```

1 // Chapter 12_The junction field effect transistor
2 //Caption_The MESFET-Channel length modulation
3 //Ex_8//page 573
4 Nd=3*10^15
5 eps=8.85*10^-14*11.7
6 L=10
7 ID1=4
8 VDSSat=0 //assume
9 VDS1=VDSSat+2
10 VDS2=VDSSat+2.5
11
12 delL2=10^4*(2*eps*(VDS2-VDSSat)/(e*Nd))^0.5    //
   change in length
13 delL1=10^4*(2*eps*(VDS1-VDSSat)/(e*Nd))^0.5    //
   change in length
14 //drain currents are
15 ID22=ID1*(L/(L-0.5*delL2))

```

```

16 ID11=ID1*(L/(L-0.5*dell1))
17 rds=(VDS2-VDS1)/(ID22-ID11)
18 printf('The small signal output resistance at the
      drain terminal due to channel length modulation
      effects is %1.1f kohm',rds)

```

---

### Scilab code Exa 12.9 cut off frequency

```

1 // Chapter 12_The junction field effect transistor
2 //Caption_Cutoff frequency
3 //Ex_9//page 579
4 e=1.6*10^-19
5 mun=1000
6 L=5*10^-4
7 eps=8.85*10^-14*11.7
8 a=0.60*10^-4
9 Nd=10^16
10 fT=(e*mun*Nd*a^2)/(2*pi*eps*L^2)*10^-9
11 printf('The cutoff frequency of silicon JFET with
      given parameters is %1.2f GHz',fT)

```

---

### Scilab code Exa 12.10 High electron mobility transistor

```

1 // Chapter 12_The junction field effect transistor
2 //Caption_High electron mobility transistor
3 //Ex_10//page 585
4 Nd=10^18
5 d=20*10^-8
6 dd=500*10^-8    // thickness
7 phi_B=0.85
8 q=1.6*10^-19
9 VG=0
10 epsn=12.2 // relative dielectric constant

```

```
11 Vp2=q*Nd*dd^2/(2*epsn*8.85*10^-14) //a parameter
12 x=0.22 //x=del Ec/q
13 Voff=phi_B-x-Vp2 //threshold voltage
14 ns=(VG-Voff)*epsn*8.85*10^-14/(q*(dd+d+80*10^-8))
15 printf('The two dimensional electron concentration
is %1.2f cm^-2',ns)
```

---

# Chapter 13

## Optical devices

**Scilab code Exa 13.1** Optical absorption

```
1 // Chapter 13_Optical Devices
2 //Caption_Optical absorption
3 //Ex_1//page 598
4 lambdai1=1*10^-4      //incident wavelength
5 lambdai2=0.5*10^-4
6 alpha1=100    //absorption coefficient
7 d1=1*log(1/0.1)/alpha1    //If 90 percent of the
                             incident flux is to be absorbed in a distance d ,
                             then the flux emerging at x=d will be 10% of the
                             incident flux
8 alpha2=10000
9 d2=1*log(1/0.1)/alpha2*10^4
10 printf('As the incident photon energy increases , the
           absorption coefficient increases rapidly since
           d1=%1.4f cm and d2=%1.2f micrometer ',d1,d2)
```

---

**Scilab code Exa 13.2** Electron hole pair generation rate

```

1 // Chapter 13_Optical Devices
2 //Caption_Electron hole pair generation rate
3 //Ex_2//page 600
4 T=300
5 Ivx=0.05      //photon intensity
6 lambda=0.75   //wavelength
7 alpha=0.7*10^4           //absorption coefficient
8 h=1.24
9 v=1/lambda // v is the frequency
10 E=h*v    //energy in eV,
11 g=alpha*Ivx/(1.6*10^-19*h*v)      //generation rate of
    electron hole pair
12 tau=10^-7      //lifetime of minority carrier
13 deln=g*tau      //excess carrier concentration
14 printf('The generation rate of electron hole pair is
    %1.2f cm^-3 s^-1',g)

```

---

### Scilab code Exa 13.3 Solar cells

```

1 // Chapter 13_Optical Devices
2 //Caption_Solar cells
3 //Ex_3//page 602
4 Na=5*10^18
5 Nd=10^16
6 Dn=25
7 e=1.6*10^-19
8 ni=1.5*10^10
9 Dp=10
10 tau_no=5*10^-7
11 tau_po=10^-7
12 JL=15*10^-3      //photocurrent density
13 Ln=(Dn*tau_no)^0.5
14 Lp=(Dp*tau_po)^0.5
15 Js=e*(ni^2)*((Dn/(Ln*Na))+(Dp/(Lp*Nd)))
16 Voc=0.0259*log(1+JL/Js)

```

```
17 printf('Open circuit voltage of SI pn juncton solar  
cell is %1.3f V',Voc)
```

---

#### Scilab code Exa 13.4 Solar concentration

```
1 // Chapter 13_Optical Devices  
2 //Caption_Solar concentration  
3 //Ex_4//page 605  
4 JL==150*10^-3      //PHOTOCURRENT DENSITY  
5 Js=3.6*10^-11     //reverse saturation current density  
6 Voc=0.0259*log(1+JL/Js)  
7 printf('Open circuit voltage when solar  
concentration is used is %1.3f V',Voc)
```

---

#### Scilab code Exa 14.4 Heat sinks and junction temperature

```
1 // Chapter 14_Semiconductor Power Devices  
2 //Caption_Heat sinks and junction temperature  
3 //Ex_4//page-663  
4 P=20      //Rated power  
5 Tj_max=175    //Junction temperature  
6 TOC=25  
7 Tamb=25      //ambient temperature  
8 Theta_case_snk=1  
9 Theta_snk_amb=5  
10 Theta_dev_case=(Tj_max-TOC)/P  
11 PD_MAX=(Tj_max-Tamb)/(Theta_dev_case+Theta_case_snk+  
Theta_snk_amb)  
12 printf('Maximum power dissipated is %1.1f W',PD_MAX)
```

---

### Scilab code Exa 13.5 Photo conductor

```
1 // Chapter 13_Optical Devices
2 //Caption_Photo conductor
3 //Ex_5//page 611
4 mup=480
5 mun=1350
6 L=100*10^-4 //length of photoconductor
7 A=10^-7 //cross sectional area
8 tau_p=10^-6 //minority carrier lifetime
9 V=10 //applied voltage
10 tn=L^2/(mun*V)
11 //photoconductor gain is
12 G=(tau_p/tn)*(1+(mup/mun))
13 printf('The photoconductor gain is %1.2f',G)
```

---

### Scilab code Exa 13.6 Photo diode

```
1 // Chapter 13_Optical Devices
2 //Caption_Photodiode
3 //Ex_6//page 616
4 Na=10^16
5 eps=8.85*10^-14;
6 Nd=10^16
7 Dn=25
8 Dp=10
9 tau_no=5*10^-7
10 e=1.6*10^-19
11 ni=1.5*10^10
12 tau_po=10^-7
13 VR=5 //reverse bias voltage
14 GL=10^21 //generation rate of excess carriers
15 Ln=(Dn*tau_no)^0.5
16 Lp=(Dp*tau_po)^0.5
17 Vbi=0.0259*log(Na*Nd/ni^2)
```

```
18 W=((2*eps/e)*((Na+Nd)/(Na*Nd))*(Vbi+VR))^0.5
19 JL=e*(W+Ln+Lp)*GL
20 printf('The steady state photocurrent density is %1
.2 f A/cm^2 ', JL)
```

---

### Scilab code Exa 13.7 PIN Photodiode

```
1 // Chapter 13_Optical Devices
2 //Caption_PIN Photodiode
3 //Ex_7//page 618
4 W=20*10^-4 //intrinsic region width
5 phio=10^17 //photon flux
6 alpha=10^3 //absorption coefficient
7 GL1=alpha*phio //generation rate of electron hole
pair at the front region
8 GL2=GL1*exp(-alpha*W)
9 JL=1000*e*phio*(1-exp(-alpha*W)) //photocurrent
density
10 printf('The photocurrent density in PIN photodiode
is %1.1 f mA/cm^2 ', JL)
```

---

### Scilab code Exa 13.8 Materials

```
1 // Chapter 13_Optical Devices
2 //Caption_Materials
3 //Ex_8//page 625
4 Eg=1.42
5 lambda=1.24/Eg //output wavelength of photon
6 lam=0.653 //desired wavelength
7 E=1.24/lam //bandgap energy
8 printf('The band gap energy corresponding to visible
given wavelength is %1.2 f eV and it would
correspond to a mole fraction of x=4 ',E)
```

---

### Scilab code Exa 13.9 Quantum efficiency

```
1 // Chapter 13_Optical Devices
2 //Caption_Quantum efficiency
3 //Ex_9//page 628
4 n2=3.666 //index of refraction in GaAs
5 n1=1 //index of refraction in air
6 T=((n2-n1)/(n2+n1))^2 //reflection coefficient
7 printf('The reflection coefficient at semiconductor-
air interface ius %1.2f',T)
```

---

### Scilab code Exa 13.10 Quantum efficiency

```
1 // Chapter 13_Optical Devices
2 //Caption_Quantum efficiency
3 //Ex_10//page 629
4 n2=3.66 //index of refraction in GaAs
5 n1=1 //index of refraction in air
6 theta=asind(n1/n2)
7 printf('The critical angle at semiconductor-air
interface is %1.1f degree',theta)
```

---

# Chapter 14

## Semiconductor Power Devices

**Scilab code Exa 14.1** Power transistor characteristics

```
1 // Chapter 14_Semiconductor Power Devices
2 //Caption_Power transistor characteristics
3 //Ex_1//page-651
4 RL=10
5 Vcc=35
6 Ic_max=Vcc/RL
7 Ic=Vcc/(2*RL)
8 VCE=Vcc-Ic*RL      //Collector emitter voltage at
                      maximum power point
9 PT=VCE*Ic          //Maximum transistor power dissipation
10 printf('The maximum power dissipation in transistor
           occurs at centre of the load line. The maximum
           power dissipation is therefore %1.1f W',PT)
```

---

**Scilab code Exa 14.2** Power MOSFET characteristics

```
1 // Chapter 14_Semiconductor Power Devices
2 //Caption_Power MOSFET characteristics
```

```

3 //Ex_2//page-658
4 VDD=24
5 PT=30    //Maximum rated power
6 ID1max=5    //Maximum rated current
7 ID2max=4
8 RD1=VDD/ID1max      //Drain resistance
9 RD2=VDD/ID2max
10 ID1=VDD/(2*RD1)    //Current at the maximum power
    point
11 ID2=VDD/(2*RD2)
12 VDS1=VDD-ID1*RD1    //Drain to source voltage
13 VDS2=VDD-ID2*RD2
14 P1=VDS1*ID1    //Maximum power that may be dissipated
    in transistor
15 P2=VDS2*ID2
16 printf('The maximum dissipated power in first case
    is %1.0f W which corresponds to the maximum rated
    power while in second case is %1.0f W which is
    less than the maximum rated power',P1,P2)

```

---

### Scilab code Exa 14.3 Heat sinks and junction temperature

```

1 // Chapter 14_Semiconductor Power Devices
2 //Caption_Heat sinks and junction temperature
3 //Ex_3//page-662
4 Theta_dev_case=1.75
5 Theta_case_snk=1
6 Theta_snk_amb=5
7 Theta_case_amb=50
8 Tamb=30 //Ambient temperature
9 Tdev=150 //maximum junction or device temperature
10 PD_max=(Tdev-Tamb)/(Theta_dev_case+Theta_case_amb)
    //when no heat sink is used
11 PD_MAX2=(Tdev-Tamb)/(Theta_dev_case+Theta_case_snk+
    Theta_snk_amb)

```

```
12 printf('Maximum power dissipated when no sink was  
used is %1.2f W while with the sink is %1.2f W  
which is more than the previous case. Thus use of  
heat sink allows more power to be dissipated in  
the device.',PD_max,PD_MAX2)
```

---