

Scilab Textbook Companion for  
Transport Phenomena  
by R. S. Brodkey And H. C. Hershey<sup>1</sup>

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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 transport phenomena

Scilab code Exa 1.1 fundamental variables and units

```
1 clc;
2 warning('off');
3 printf("\\n\\n example1.1 - pg6");
4 // given
5 v=0.01283; // [m^3] - volume of tank in m^3
6 v=0.4531; // [ft ^3] - volume of tank in ft ^3
7 p=2; // [atm] - pressure
8 T=1.8*300; // [degR] - temperature
9 R=0.73; // [(atm*ft ^3)/(lbmol*degR)] - gas constant
10 // using the equation of state for an ideal gas pv=
    nRT
11 n=(p*v)/(R*T);
12 disp(n,"no. of moles ,n=");
13 xN2=0.5; // fraction of N2 in tank
14 nN2=xN2*n;
15 Ca=nN2/v;
16 printf("\\n\\n Ca=%elb*mol/ft ^3" ,Ca);
```

---

### Scilab code Exa 1.2 The role of intermolecular forces

```
1 clc;
2 warning(" off");
3 printf("\n\n example1.2 - pg9");
4 // given
5 // the three unknowns are x,y,z
6 // the three equations are-
7 // x+y+z=1500
8 // (1) 0.05*x+0.15*y+0.40*z=1500*0.25
9 // (2) 0.95*x+0.00*y+0.452*z=1500*0.50
10 a=[1 1 1;0.05 0.15 0.40;0.95 0 0.452];
11 d=[1500;1500*0.25;1500*0.50];
12 ainv=inv(a);
13 sol=ainv*d;
14 printf("\n\n the amount of concentrated HNO3 is %fkg
    \n the amount of concentrated H2SO4 is %fkg\n the
    amount of waste acids is %fkg",sol(2),sol(1),sol
    (3));
```

---

# Chapter 2

## molecular transport mechanisms

**Scilab code Exa 2.1** the analogous form

```
1 clc;
2 warning('off');
3 printf("\n\n example2.1 - pg28");
4 // given
5 deltax=0.1; // [m] - thickness of copper block
6 T2=100; // [degC] - temp on one side of copper block
7 T1=0; // [degC] - temp on other side of the copper
        block
8 k=380; // [W/mK] - thermal conductivity
9 // using the formula (q/A)*deltax=-k*(T2-T1)
10 g=-k*(T2-T1)/deltax;
11 g1=(g/(4.184*10000));
12 printf("\n\n The steady state heat flux across the
        copper block is \n q/A=%fW/m^2 \n or in alternate
        units is \n q/A=%fcal/cm*sec",g,g1);
```

---

### Scilab code Exa 2.2 the analogous form

```
1 clc;
2 warning('off');
3 printf("\n\n example2.2 - pg29");
4 // given
5 dely=0.1; // [m] - distance between two parallel
    plates
6 delUx=0.3; // [m/sec] - velocity of a plate
7 mu=0.001; // [kg/m*sec] - viscosity
8 // using the formula tauyx=F/A=mu*(delUx/dely)
9 tauyx=-mu*(delUx/dely);
10 printf("\n\n the momentum flux and the force per
    unit area ,(which are the same thing) is \n tauyx=
    F/A=%fN/m^2",tauyx);
```

---

### Scilab code Exa 2.3 heat transfer

```
1 clc;
2 warning('off');
3 printf("\n\n example2.3 - pg30");
4 // given
5 tauyx=-0.003; // [N/m^2] - momentum flux
6 dely=0.1; // [m] - distance between two parallel
    plates
7 mu=0.01; // [kg/m*sec] - viscosity
8 // using the formula tauyx=F/A=mu*(delUx/dely)
9 delUx=-((tauyx*dely)/mu)*100;
10 printf("\n\n Velocity of the top plate is \n deltaUx
    =%fcm/sec",delUx);
```

---

### Scilab code Exa 2.5 heat transfer

```

1 clc;
2 warning('off');
3 printf("\n\n example2.5 - pg31");
4 // given
5 d=0.0013; // [m] - diameter of the tube
6 delx=1; // [m] - length of the glass tube
7 T2=110.6; // [degC] - temperature on one end of the
    rod
8 T1=0; // [degC] - temperature on other side of the
    rod
9 k=0.86; // [W/m*K] - thermal conductivity
10 Hf=333.5; // [J/g] - heat of fusion of ice
11 // (a) using the equation (q/A)=-k*(delt/delx)
12 A=(%pi*d^2)/4;
13 q=A*(-k*(T2-T1)/delx);
14 printf("\n\n (a) the heat flow is \n q=%fJ/sec",q);
15 // (b) dividing the total heat transfer in 30 minutes
    by the amount of heat required to melt 1g of ice
16 a=abs((q*30*60)/333.5);
17 printf("\n\n (b) the amount or grams of ice melted in
    30 minutes is %fg",a);

```

---

### Scilab code Exa 2.6 mass transfer

```

1 clc;
2 warning('off');
3 printf("\n\n example2.6 - pg36");
4 // given
5 d=1.2*10^-2; // [m] - diameter of the hole
6 Ca1=0.083; // [kmol/m^3]
7 Ca2=0; // [kmol/m^3]
8 L=0.04; // [m] - thickness of the iron piece
9 Dab=1.56*10^-3; // [m^2/sec] - diffusion coefficient
    of CO2
10 A=(%pi*d^2)/4; // area

```

```

11 // (a) using the formula  $(Na/A)A = (Ja/A) = -Dab \frac{\partial Ca}{\partial x}$ 
12 intdCa=integrate('1','Ca',Ca2,Ca1);
13 intdx=integrate('1','x',0,0.04);
14 g=(intdCa/intdx)*Dab;
15 printf("\n\n (a) The molar flux with respect to
           stationary coordinates is \n  $(Na/A) = %f \text{ kmol/m}^2 \text{ sec}$ 
           ",g);
16 // using the formula  $na/A = (Na/A) * Ma$ 
17 Ma=44.01; // [kg/mol] – molecular weight of co2
18 na=(intdCa/intdx)*Dab*Ma*A*(3600/0.4539);
19 printf("\n\n The mass flow rate is %fb/hr",na);

```

---

### Scilab code Exa 2.7 mass transfer

```

1 clc;
2 warning('off');
3 printf("\n\n example2.7 – pg38");
4 // given
5 T=30+273.15; // [K] temperature
6 pA=3; // [atm] partial pressure of the component A
7 R=0.082057; // [atm*m^3*kmol*K] gas constant
8 // (a) using the equation  $Ca = n/V = pA / (R*T)$ 
9 Cco2=pA/(R*T);
10 Cco2=Cco2*(44.01);
11 printf("\n\n (a) The concentration of Co2 entering is
           %f kg/m^3",Cco2);
12 // (b) using the same equation as above
13 pN2=(0.79)*3; // [atm] partial pressure of nitrogen(
                  as nitrogen is 79% in air)
14 R=0.7302; // [atm*ft^3*lb/mol*R] – gas constant
15 T=T*(1.8); // [R] temperature
16 CN2=pN2/(R*T);
17 printf("\n\n (b) The concentration of N2 entering is
           %f lbmol/ft^3",CN2);

```

```

18 // (c) using the same equation as above
19 nt=6;
20 nCo2=4;
21 nO2=2*(0.21);
22 nN2=2*(0.79);
23 yCo2=nCo2/nt;
24 yO2=nO2/nt;
25 yN2=nN2/nt;
26 R=82.057; // [atm*cm^3/mol*K] - gas constant
27 T=30+273.15; // [K] - temperature
28 pCo2=3*yCo2;
29 Cco2=pCo2/(R*T);
30 printf("\n\n (c) The concentrartion of Co2 in the
exit is %fmol/cm^3",Cco2);
31 // (d) using the same equation as above
32 R=8.3143; // [kPa*m^3/kmol*K] - gas constant
33 pO2=3*(yO2)*(101.325); // [kPa] - partial pressure
34 CO2=pO2/(R*T);
35 printf("\n\n (d) The concentration of O2 in the exit
stream is %fkmol/m^3",CO2);

```

---

### Scilab code Exa 2.8 mass transfer

```

1 clc;
2 warning('off');
3 printf("\n\n example2.8 - pg39");
4 // given
5 delx=0.3-0; // [m] - length
6 d=0.05-0; // [m] - diameter
7 A=(%pi*d^2)/4; // [m^2] - area;
8 R=8.314*10^3; // [N*m/kmol*K] - gas constant
9 xco1=0.15; // mole prcent of co in one tank
10 xco2=0; // mole percent of co in other tank
11 p2=1; // [atm] - pressure in one tank
12 p1=p2; // [atm] - pressure in other tank

```

```

13 D=0.164*10^-4; // [m^2/sec] - diffusion coefficient
14 T=298.15; // [K] - temperature
15 // using the formula (Na/A)=(Ja/A)=D*(delca/delx)
16 // =-(D/R*T)*(delpa/delx);
16 delpa=(p2*xco2-p1*xco1)*10^5; // [N/m^2] - pressure
   difference
17 Na=-((D*A)/(R*T))*(delpa/delx);
18 disp(Na)
19 printf("\n\n The initial rate of mass transfer of
   co2 is %ekmol/sec",Na);
20 printf("\n\n In order for the pressure to remain at
   1atm, a diffusion of air must occur which is in
   the opposite direction and equal to %ekmol/sce",
   Na);

```

---

### Scilab code Exa 2.9 momentum transfer

```

1 clc;
2 warning("off");
3 printf("\n\n example2.9 - pg44");
4 // given
5 A=5; // [m^2] - area of the plates
6 Ft=0.083 // [N] - force on the top plate
7 Fb=-0.027; // [N] - force on the bottom plate
8 ut=-0.3; // [m/sec] - velocity of the top plate
9 ub=0.1; // [m/sec] - velocity of the bottom plate
10 dely=0.01; // [m]
11 delux=ut-ub; // [m/sec]
12 // using the formula tauyx=F/A=-mu(delux/dely)
13 tauyx=(Ft-Fb)/A;
14 mu=tauyx/(-delux/dely); // [Ns/m^2]
15 mu=mu*10^3; // [cP]
16 printf("\n\n The viscosity of toulene in centipose
   is %fcP",mu);

```

---

### Scilab code Exa 2.11 diffusion coefficient

```
1 clc;
2 warning('off');
3 printf("\n\n example2.11 - pg51");
4 // given
5 po=1; // [atm] - pressure
6 p=2; // [atm] - pressure
7 To=0+273.15; // [K] - temperature
8 T=75+273.15; // [K] - temperature
9 Do=0.219*10^-4; // [m^2/sec];
10 n=1.75;
11 // using the formula D=Do*(po/p)*(T/To)^n
12 D=Do*(po/p)*(T/To)^n;
13 printf("\n\n The diffusion coefficient of water
vapour in air at %f atm and %f degC is \n D=%em^2/
sec",p,T-273.15,D);
```

---

### Scilab code Exa 2.12 viscosity

```
1 clc;
2 warning('off');
3 printf("\n\n example2.12 - pg52");
4 // given
5 T=53+273.15; // [K] - temperature
6 mu1=1.91*10^-5;
7 mu2=2.10*10^-5;
8 T1=313.15; // [K] - temperature
9 T2=347.15; // [K] - temperature
10 // for air
11 // using linear interpolation of the values in table
2.2
```

```

12 function b=f(a)
13     b=log(mu1/a)/log(T1);
14 endfunction
15 function y=g(a)
16     y=log(mu2)-log(a)-f(a)*log(T2);
17 endfunction
18 a1=10^-7;
19 A=fsolve(a1,g);
20 B=f(A);
21 // using the formula ln(mu)=lnA+Bln(t)
22 mu=%e^(log(A)+B*log(T))*10^3; // [cP]
23 printf("\n\n the viscosity of air at %fdegC is %fcP"
    ,T-273.15,mu);
24 // similarly for water
25 BdivR=1646;
26 A=3.336*10^-8;
27 mu=A*%e^(BdivR/T)*10^5 // [cP]
28 printf("\n\n the viscosity of water at %fdegC is
    %fcP",T-273.15,mu);

```

---

# Chapter 3

## molecular transport and the general property balance

Scilab code Exa 3.1 balance or conservation concept

```
1 clc;
2 warning(" off");
3 printf("\n\n example3.1 - pg65");
4 // given
5 a=0.0006; // [m^2] - area
6 l=0.1; // [m] - length
7 // (a) using the fourier law
8 deltax=0.1; // [m] - thickness of copper block
9 T2=100; // [degC] - temp on one side of copper block
10 T1=0; // [degC] - temp on other side of the copper
    block
11 k=380; // [W/mK] - thermal conductivity
12 // using the formula (q/A)*deltax=-k*(T2-T1)
13 g=-k*(T2-T1)/deltax;
14 printf("\n\n (a) The steady state heat flux across
    the copper block is \n q/A=%5eJ*m^-2*sec^-1 ",g);
15 // (b)
16 V=a*l; // [m^3] - volume
17 // using the overall balance equation with the
```

```

        accumulation and generation term
18 Qgen=1.5*10^6; // [j*m^-3*sec^-1]
19 SIx=(g*a-Qgen*v)/a;
20 printf("\n\n (b) the flux at face 1 is %5ej*m^-2*sec
    ^-1;\nthe negative sign indicates taht the heat
    flux is from right to left (negative x direction",
SIx);

```

---

**Scilab code Exa 3.2** the balance equation in differential form

```

1 clc;
2 warning('off');
3 printf("\n\n example3.2 - pg68");
4 // given
5 sym x;
6 SIx2=-3.8*10^5; // [j*m^-2*sec^-1] - flux at x=0.1, i
    .e through face2
7 Qgen=1.5*10^6; // [j*m^-3*sec^-1] - uniform
    generation in the volume
8 T2=100+273.15; // [K] temperature at face 2
9 x2=0.1; // [m]
10 k=380; // [W/mK] - thermal conductivity
11 // using the equation der(SIx)*x=SIx+c1; where c1 is
    tyhe constant of integration
12 c1=(Qgen*x2)-SIx2;
13 disp(c1)
14 SIx=Qgen*x-c1;
15 disp(SIx,"SIx=");
16 printf("\n where SIx is in units of j m^-2 sec^-1
    and x is in units of m");
17 // using the equation -k*T=der(SIx)*x^2-c1*x+c2;
    where c2 is the constant of integration
18 c2=-k*T2-(Qgen*(x2)^2)/2+c1*x2;
19 T=-(Qgen/k)*x^2+(c1/k)*x-(c2/k);
20 disp(T,"T=");

```

```
21 printf("\n where T is in units of kelvin(K));
```

---

**Scilab code Exa 3.3** the balance equation in differential form

```
1 clc;
2 warning("off");
3 printf("\n\n example3.3 - pg69");
4 // given
5 syms t x;
6 hf1=-270; // [J/sec] - heat flow at face 1
7 hf2=-228; // [J/sec] - heat flow at face2
8 Qgen=1.5*10^6; // [J*m^-3*sec^-1] generation per
      unit volume per unit time
9 v=6*10^-5; // [m^3] volume
10 Cp=0.093; // [cal*g^-1*K^-1] heat capacity of copper
11 sp=8.91; // specific gravity of copper
12 a=0.0006; // [m^2] - area
13 // (a) using the overall balance
14 acc=hf1-hf2+Qgen*v;
15 printf("\n\n (a) the rate of accumulation is %fJ/sec
      \n\n ",acc);
16 // (b)
17 SIx1=hf1/a;
18 SIx2=hf2/a;
19 x1=0;
20 // solving for the constant of integration c1 in the
      equation [del(p*cp*T)/delt-der(SIx)]*x=-SIx+c1;
21 c1=0+SIx1;
22 x2=0.1;
23 g=(-(SIx2)+c1)/x2+Qgen;
24 SIx=c1-(g-Qgen)*x;
25 disp(SIx,"SI(x)=",(b));
26 // solving for constant of integration c3 in the
      equation p*cp*T=g*t+c3
27 T2=100+273.15;
```

```

28 t2=0;
29 p=sp*10^3; // [kg/m^3] - density
30 cp=Cp*4.1840; // [J*kg^-1*K^-1]
31 c3=p*cp*T2-g*t2;
32 T=(g*(10^-3)/(p*cp))*t+c3/(p*cp);
33 disp(T,"T=");
34 // solving for constant of integration c2 in the
   equation -k*T=der(SIx)*x^2-c1*x+c2
35 k=380; // [w/m^1*K^1]
36 x2=0.1;
37 c2=k*T+(3.5*10^5)*x2^2-(4.5*10^5)*x2;
38 function y=T(t,x)
39     y=(-(3.5*10^5)*x^2+(4.5*10^5)*x+87.7*t
         +1.00297*10^5)/k;
40 endfunction
41 // at face 1;
42 x1=0;
43 t1=60; // [sec]
44 T1=T(t1,x1);
45 disp(T1,"T=","at face 1");
46 // at face 2
47 x2=0.1;
48 t2=60; // [sec]
49 T2=T(t2,x2);
50 disp(T2,"T=","at face 2");

```

---

# Chapter 4

## molecular transport and the general property balance

Scilab code Exa 4.1 variable area transport

```
1 clc;
2 warning('off');
3 printf("\n\n example4.1 - pg99");
4 // given
5 id=2.067; // [in] - inside diameter
6 t=0.154; // [in] - wall thickness
7 od=id+2*t; // [in] - outer diameter
8 a=1.075; // [in^2] - wall sectional area of metal
9 A=a*(1/144); // [ft^2] - wall sectional area of
    metal in ft^2
10 deltaz=5/12; // [ft] - length of transfer in z
    direction
11 T2=10+273.15; // [K] - temperature at the top
12 T1=0+273.15; // [K] - temperature at the bottom
13 q=-3.2; // [Btu/hr] - heat transferred
14 deltaT=(T2-T1)+8; // [degF]
15 k=-(q/A)/(deltaT/deltaz);
16 printf("\n\n korrect=%fBtu h^-1 ft^-1 degF^-1=17.17
    W m^-1 K^-1",k);
```

```

17 Alm=(2*pi*deltaz*((od-id)/(2*12)))/log(od/id); //[
    ft^2] log mean area
18 disp(Alm)
19 kincorrect=k*(A/Alm);
20 printf("\n\n kincorrect=%fBtu h^-1 ft^-1 degF
    ^-1=0.529 W m^-1 K^-1",kincorrect);
21 errorf=(k-kincorrect)/kincorrect;
22 disp(errorf," error factor is-");

```

---

### Scilab code Exa 4.2 variable area transport

```

1 clc;
2 warning('off');
3 printf("\n\n example4.2 - pg100");
4 // given
5 T1=0; // [degC]
6 T2=10; // [degC]
7 km=17.17; // [W/m*K]
8 l=1; // [m]
9 r2=1.1875;
10 r1=1.0335;
11 deltaT=T1-T2;
12 // using the formula Qr=km*((2*pi*l)/ln(r2/r1))*deltaT;
13 Qr=-km*((2*pi*l)/log(r2/r1))*deltaT;
14 printf("\n\n qr=%fW\n the plus sign indicates that
    the heat flow is radially out from the center",Qr
);

```

---

### Scilab code Exa 4.3 variable area transport

```

1 clc;
2 warning('off');

```

```

3 printf("\n\n example4.3 - pg100");
4 // given
5 km=9.92; // [Btu/h*ft*degF]
6 Alm=0.242*(12/5); // [ft^2]
7 T1=0; // [degC]
8 T2=10; // [degC]
9 deltaT=(T1-T2)*1.8; // [degF]
10 r2=1.1875;
11 r1=1.0335;
12 deltar=(r2-r1)/12; // [ft]
13 // using the formula Qr/Alm==km*(deltaT/deltar)
14 Qr=(-km*(deltaT/deltar))*Alm;
15 printf("\n\n qr=%fBtu/h",Qr);
16 // in SI units
17 Alm=0.177; // [m^2]
18 T1=0; // [degC]
19 T2=10; // [degC]
20 km=17.17; // [W/m*K]
21 r2=1.1875;
22 r1=1.0335;
23 deltaT=T1-T2;
24 deltar=(r2-r1)*0.0254; // [m]
25 // using the same formula
26 Qr=(-km*(deltaT/deltar))*Alm;
27 printf("\n\n qr=%fW",Qr);

```

---

### Scilab code Exa 4.4 variable area transport

```

1 clc;
2 warning("off");
3 printf("\n\n example4.4 - pg101");
4 // given
5 x1=0; // [cm]
6 x2=30; // [cm]
7 p1=0.3; // [atm]

```

```

8 p2=0.03; // [ atm ]
9 D=0.164; // [ am^2/sec ]
10 R=82.057; // [ cm^3*atm/mol*K ]
11 T=298.15; // [ K ]
12 // using the formula Nax*int (dx/Ax)=-(D/RT)*int (1*
   dpa)
13 a=integrate("1/((%pi/4)*(10-(x/6))^2)" , "x" , x1 , x2);
14 b=integrate("1" , "p" , p1 , p2);
15 Nax=-( (D/(R*T))*b )/a;
16 printf("\n\n Nax=%6emol/sec=%3emol/h \n the plus
      sign indicates diffusion to the right" , Nax , Nax
      *3600);

```

---

### Scilab code Exa 4.5 heat or mass transport with constant generation

```

1 clc;
2 warning(" off");
3 printf("\n\n example4.5 - pg105");
4 // given
5 syms r;
6 ro=0.5; // [ inch ] - outside radius
7 ro=0.0127; // [m] - outside radius in m
8 Tg=2*10^7; // [ J/m^3*sec ] - heat generated by
   electric current
9 Tw=30; // [degC] - outside surface temperature
10 km=17.3; // [W/m*K] - mean conductivity
11 // using the formula T=Tw+(Tg/4*km)*(ro^2-r^2)
12 T=Tw+(Tg/(4*km))*(ro^2-r^2);
13 disp(T , "T=");
14 printf("\n where r is in meters and T is in degC");
15 function y=t(r)
16     y=Tw+(Tg/(4*km))*(ro^2-r^2);
17 endfunction
18 printf("\n\n at the centre line (r=0) ,the maximum
      temperature is %fdegC .At the outside ,the

```

temperature reduces to the boundary condition value of %fdegC. The distribution i parabolic between these 2 limits”, t(0),t(0.0127));

---

### Scilab code Exa 4.7 laminar flow in a tube

```
1 clc;
2 warning(" off");
3 printf("\n\n example4.7 - pg119");
4 // given
5 r=10^-3; // [m] - radius
6 l=1; // [m] - length
7 Q=10^-7; // [m^3/s] - flow rate
8 deltap=-10^6; // [N/m^2=Pa] - pressure difference
9 spg=1.1;
10 pwater=1000; // [kg/m^3] - density of water at 4degC
11 pfluid=spg*pwater;
12 mu=(r*(deltap)*(%pi*r^3))/((4*Q)*(2*l));
13 mupoise=mu*10;
14 mucentipoise=mupoise*100;
15 printf("\n\n mu=%fNsM^-2=%fpoise=%fcP",mu,mupoise,
    mucentipoise);
```

---

# Chapter 5

## transport with a net convective flux

Scilab code Exa 5.9 mass fluxes in stationary and convected coordinates

```
1 clc;
2 warning("off");
3 printf("\n\n example5.9 - pg166");
4 // given
5 v=1; // [cm/sec] - volume velocity or bulk velocity
6 vol=1; // [cm^3] - volume
7 na=2; // moles of a
8 nb=3; // moles of b
9 nc=4; // moles of c
10 mma=2; // molecular weight of a
11 mmb=3; // molecular weight of b
12 mmc=4; // molecular weight of c
13 ma=na*mma; // [g] weight of a
14 mb=nb*mmb; // [g] weight of b
15 mc=nc*mmc; // [g] weight of c
16 NabbyA=2+2; // [mol/cm^2*s] - molar flux = diffusing
    flux + convected flux
17 NbbbyA=-1+3; // [mol/cm^2*s] - molar flux = diffusing
    flux + convected flux
```

```

18 NcbyA=0+4; // [mol/cm^2*s] - molar flux = diffusing
   flux + convected flux
19 NtbyA=NabyA+NbbyA+NcbyA; // [mol/cm^2*s] - total
   molar flux
20 // on a mass basis, these corresponds to
21 nabyA=4+4; // [g/cm^2*s]; - mass flux = diffusing
   flux + convected flux
22 nbbyA=-3+9; // [g/cm^2*s]; - mass flux = diffusing
   flux + convected flux
23 ncbyA=0+16; // [g/cm^2*s]; - mass flux = diffusing
   flux + convected flux
24 ntbyA=nabyA+nbbyA+ncbyA; // [g/cm^2*s] - total mass
   flux
25 // concentrations are expressed in molar basis
26 CA=na/vol; // [mol/cm^3]
27 CB=nb/vol; // [mol/cm^3]
28 CC=nc/vol; // [mol/cm^3]
29 CT=CA+CB+CC; // [mol/cm^3] - total concentration
30 // densities are on a mass basis
31 pa=ma/vol; // [g/cm^3]
32 pb=mb/vol; // [g/cm^3]
33 pc=mc/vol; // [g/cm^3]
34 pt=pa+pb+pc; // [g/cm^3]
35 Ua=NabyA/CA; // [cm/sec];
36 Ub=NbbyA/CB; // [cm/sec];
37 Uc=NcbyA/CC; // [cm/sec];
38 // the same result will be obtained from dividing
   mass flux by density
39 Uz=(pa*Ua+pb*Ub+pc*Uc)/(pa+pb+pc);
40 printf("\n\n Uz=%fcm/sec",Uz);
41 Uzstar=(NtbyA/CT);
42 printf("\n\n Uz*=%fcm/sec",Uzstar);
43 printf("\n\n for this example both Uz and Uz* are
   slightly greater than the volume velocity of 1cm/
   sec, because there is a net molar and mass
   diffusion in the positive direction.");

```

---

### Scilab code Exa 5.10 total flux and ficks law

```
1 clc;
2 warning("off");
3 printf("\n\n example5.10 - pg171");
4 // given (from example 5.9)
5 na=2; // moles of a
6 nb=3; // moles of b
7 nc=4; // moles of c
8 mma=2; //molecular weight of a
9 mmb=3; //molecular weight of b
10 mmc=4; //molecular weight of c
11 ma=na*mma; // [g] weight of a
12 mb=nb*mmb; // [g] weight of b
13 mc=nc*mmc; // [g] weight of c
14 NabbyA=2+2; // [mol/cm^2*s] - molar flux = diffusing
    flux + convected flux
15 NbbyA=-1+3; // [mol/cm^2*s] - molar flux = diffusing
    flux + convected flux
16 NcbyA=0+4; // [mol/cm^2*s] - molar flux = diffusing
    flux + convected flux
17 NtbyA=NabyA+NbbyA+NcbyA; // [mol/cm^2*s] - total
    molar flux
18 // on a mass basis, these corresponds to
19 nabbyA=4+4; // [g/cm^2*s]; - mass flux = diffusing
    flux + convected flux
20 nbbyA=-3+9; // [g/cm^2*s]; - mass flux = diffusing
    flux + convected flux
21 ncbyA=0+16; // [g/cm^2*s]; - mass flux = diffusing
    flux + convected flux
22 // concentrations are expressed in molar basis
23 CA=na/vol; // [mol/cm^3]
24 CB=nb/vol; // [mol/cm^3]
25 CC=nc/vol; // [mol/cm^3]
```

```

26 CT=CA+CB+CC; // [ mol/cm^3 ] – total concentration
27 // densities are on a mass basis
28 pa=ma/vol; // [ g/cm^3 ]
29 pb=mb/vol; // [ g/cm^3 ]
30 pc=mc/vol; // [ g/cm^3 ]
31 Ua=NabyA/CA; // [ cm/sec ];
32 Ub=NbbyA/CB; // [ cm/sec ];
33 Uc=NcbyA/CC; // [ cm/sec ];
34 U=(pa*Ua+pb*Ub+pc*Uc)/(pa+pb+pc);
35 Ustar=(NtbyA/CT);
36 // the fluxes relative to mass average velocities
   are found as follows
37 JabyA=CA*(Ua-U); // [ mol/cm^2*sec ]
38 JbbyA=CB*(Ub-U); // [ mol/cm^2*sec ]
39 JcbyA=CC*(Uc-U); // [ mol/cm^2*sec ]
40 printf("\n\n fluxes relative to mass average
   velocities are–");
41 printf("\n\n Ja/A=%fmol/cm^2*sec",JabyA);
42 printf("\n Jb/A=%fmol/cm^2*sec",JbbyA);
43 printf("\n Jc/A=%fmol/cm^2*sec",JcbyA);
44 jabyA=pa*(Ua-U); // [ g/cm^2*sec ]
45 jbbyA=pb*(Ub-U); // [ g/cm^2*sec ]
46 jcbyA=pc*(Uc-U); // [ g/cm^2*sec ]
47 printf("\n\n ja/A=%fg/cm^2*sec",jabyA);
48 printf("\n jb/A=%fg/cm^2*sec",jbbyA);
49 printf("\n jc/A=%fg/cm^2*sec",jcbyA);
50 // the fluxes relative to molar average velocity are
   found as follows
51 JastarbyA=CA*(Ua-Ustar); // [ mol/cm^2*sec ]
52 JbstarbyA=CB*(Ub-Ustar); // [ mol/cm^2*sec ]
53 JcstarbyA=CC*(Uc-Ustar); // [ mol/cm^2*sec ]
54 printf("\n\n fluxes relative to molar average
   velocities are–");
55 printf("\n\n Ja*/A=%fmol/cm^2*sec",JastarbyA);
56 printf("\n Jb*/A=%fmol/cm^2*sec",JbstarbyA);
57 printf("\n Jc*/A=%fmol/cm^2*sec",JcstarbyA);
58 jastarbyA=pa*(Ua-Ustar); // [ g/cm^2*sec ]
59 jbstarbyA=pb*(Ub-Ustar); // [ g/cm^2*sec ]

```

```

60 jcstarbyA=pc*(Uc-Ustar); // [g/cm^2*sec]
61 printf("\n\n ja*/A=%fg/cm^2*sec",jastarbyA);
62 printf("\n jb*/A=%fg/cm^2*sec",jbstarbyA);
63 printf("\n jc*/A=%fg/cm^2*sec",jcstarbyA);

```

---

### Scilab code Exa 5.11 binary mass diffusion in gases

```

1 clc;
2 warning("off");
3 printf("\n\n example5.11 - pg176");
4 // given
5 T=0+273.15; // [K] - temperature in Kelvins
6 pa2=1.5; // [atm] - partial pressure of a at point2
7 pa1=0.5; // [atm] - partial pressure of a at point 1
8 z2=20; // [cm] - position of point 2 from reference
    point
9 z1=0; // [cm] - position of point1 from reference
    point
10 p=2; // [atm] - total pressure
11 d=1; // [cm] - diameter
12 D=0.275; // [cm^2/sec] - diffusion coefficient
13 A=(%pi*((d)^2))/4;
14 R=0.082057; // [atm*m^3*kmol^-1*K^-1] - gas constant
15 // (a) using the formula Na/A=-(D/(R*T))*((pa2-pa1)
    /(z2-z1))
16 Na=(-(D/(R*T))*((pa2-pa1)/(z2-z1)))*(A)/(10^6);
17 printf("\n\n Na=%ekmol/sec\n The negative sign
    indicates diffusion from point 2 to point 1",Na);
18 pb2=p-pa2;
19 pb1=p-pa1;
20 // (b) using the formula Na/A=((D*p)/(R*T*(z2-z1)))*
    ln(pb2/pb1)
21 Na(((D*p)/(R*T*(z2-z1)))*log(pb2/pb1))*(A)/(10^6);
22 printf("\n\n Na=%ekmol/sec",Na);
23 printf("\n The induced velocity increases the net

```

transport of A by the ratio of  $10.6 \times 10^{-10}$  to  $4.82 \times 10^{-10}$  or 2.2 times. This increase is equivalent to 120 percent");

---

### Scilab code Exa 5.12 binary mass diffusion in gases

```
1 clc;
2 warning("off");
3 printf("\n\n example5.12 - pg178");
4 // given
5 T=0+273.15; // [K] - temperature in Kelvins
6 pa2=1.5; // [atm] - partial pressure of a at point2
7 pa1=0.5; // [atm] - partial pressure of a at point 1
8 z2=20; // [cm] - position of point 2 from reference
    point
9 z1=0; // [cm] - position of point1 from reference
    point
10 p=2; // [atm] - total pressure
11 d=1; // [cm] - diameter
12 D=0.275; // [cm^2/sec] - diffusion coefficient
13 A=(%pi*((d)^2))/4;
14 R=0.082057; // [atm*m^3*kmol^-1*K^-1] - gas constant
15 k=0.75;
16 // using the formula (Na/A)=-(D/(R*T*(z2-z1)))*ln
    ((1-(pa2/p)*(1-k))/(1-(pa1/p)*(1-k)))
17 NabA=-(D/(R*T*(z2-z1)))*(2*0.7854)*log((1-(pa2/p)
    *(1-k))/(1-(pa1/p)*(1-k)))/(10^6);
18 printf("\n\n (Na/A)=%ekmol/sec",NabA);
19 printf("\n Note that this answer is larger than the
    rate for equimolar counter diffusion but smaller
    than the rate for diffusion through a stagnant
    film. Sometimes the rate for diffusion through a
    stagnant film can be considered as an upper bound
    , if k lies between zero and one");
```

---

### Scilab code Exa 5.13 diffusion due to pressure gradient

```
1 clc;
2 warning(" off");
3 printf("\n\n example5.13 - pg184");
4 // given
5 l=4; // [m] - length of the tube
6 id=1.6*10^-3; // [m] - inside diameter
7 Nkn=10; // - knudsen no.
8 Ma=92; // - molecular weight of gas
9 mu=6.5*10^-4; // [kg/m*sec] - viscosity
10 T=300; // [K] - temperature
11 R=8314; // [kPa*m^3*kmol^-1*K^-1] - gas constant
12 lambdaA=Nkn*id; // [m] mean free path
13 // for calculating pressure using the formula lamdaA
   =32*(mu/p)*((R*T)/(2*pi*Ma))^(1/2)
14 p=32*(mu/lambdaA)*((R*T)/(2*pi*Ma))^(1/2);
15 patm=p/(1.01325*10^5);
16 printf("\n\n p=%f kg/m*sec^2=%f Pa=%e atm" ,p,p,patm);
17 printf("\n The value of 10 for the knudsen number is
   on the border between Knudsen diffusion and
   transition flow");
```

---

# Chapter 6

## flow turbulence

**Scilab code Exa 6.1** the reynolds experiment

```
1 clc;
2 warning(" off");
3 printf("\\n\\n example6.1 - pg200");
4 // given
5 q=50; // [gal/min] - volumetric flow rate
6 d=2.067/12; // [ft] - diameter
7 A=0.02330; // [ft ^2] - flow area
8 p=0.99568*62.43; // [lb/ft ^3] - density of water at
86degF
9 mu=0.8007*6.72*10^-4; // [lb/ft *sec] - viscosity of
water at 86degF
10 u=q/(60*7.48*A);
11 // using the formula Nre=d*u*p/mu;
12 Nre=(d*u*p)/mu;
13 disp(Nre," Nre=");
14 printf("\\n Hence the flow is turbulent. Note also
that Nre is dimensionless");
```

---

**Scilab code Exa 6.2** transitional flow

```

1 clc;
2 warning(" off");
3 printf("\n\n example6.2 - pg202");
4 // given
5 p=0.99568*62.43; // [lb/ft ^3] - density of water at
86degF
6 mu=0.8007*6.72*10^-4; // [lb/ft*sec] - viscosity of
water at 86degF
7 u=4.78; // [ft/sec] - free stream velocity
8 Nre=5*10^5; // the lower limit for the transition
reynolds number range is substituted
9 x=(Nre*mu)/(p*u);
10 disp(x,"x");
11 printf("\\nThus the transition could start at about
%fft. The reynolds number at the upper end of the
transition range is %e. The value of x at this
location is ten times then the value obtained
above i.e %fft",x,Nre*10,x*10);

```

---

**Scilab code Exa 6.3** the equations for transport under turbulent conditions

```

1 clc;
2 warning(" off");
3 printf("\n\n example6.3 - pg212");
4 // given
5 t=[0 0.01 0.02 0.03 0.04 0.05 0.06 0.07 0.08 0.09
0.10 0.11 0.12];
6 Ux=[3.84 3.50 3.80 3.60 4.20 4.00 3.00 3.20 3.40
3.00 3.50 4.30 3.80];
7 Uy=[0.43 0.21 0.18 0.30 0.36 0.28 0.35 0.27 0.21
0.22 0.23 0.36 0.35];
8 Uz=[0.19 0.16 0.17 0.13 0.09 0.10 0.16 0.15 0.13
0.18 0.17 0.18 0.17];
9 // using the formula AREA=(deltat /2)*(U1+U13+2*(U2+

```

```

        U3+U4+U5+U6+U7+U8+U9+U10+U11+U12)) )
10 // for Uxmean
11 deltat=0.01;
12 T=t(13)-t(1);
13 AREA=(deltat/2)*(Ux(1)+Ux(13)+2*(Ux(2)+Ux(3)+Ux(4) +
    Ux(5)+Ux(6)+Ux(7)+Ux(8)+Ux(9)+Ux(10)+Ux(11)+Ux
    (12)));
14 Uxmean=AREA/T;
15 disp(Uxmean,"Uxmean=");
16 // for Uymean
17 deltat=0.01;
18 T=t(13)-t(1);
19 AREA=(deltat/2)*(Uy(1)+Uy(13)+2*(Uy(2)+Uy(3)+Uy(4) +
    Uy(5)+Uy(6)+Uy(7)+Uy(8)+Uy(9)+Uy(10)+Uy(11)+Uy
    (12)));
20 Uymean=AREA/T;
21 disp(Uymean,"Uymean=");
22 // for Uzmean
23 deltat=0.01;
24 T=t(13)-t(1);
25 AREA=(deltat/2)*(Uz(1)+Uz(13)+2*(Uz(2)+Uz(3)+Uz(4) +
    Uz(5)+Uz(6)+Uz(7)+Uz(8)+Uz(9)+Uz(10)+Uz(11)+Uz
    (12)));
26 Uzmean=AREA/T;
27 disp(Uzmean,"Uzmean=");
28 U=(Uxmean^2+Uymean^2+Uzmean^2)^(1/2);
29 disp(U,"U=");

```

---

### Scilab code Exa 6.5 the prandtl mixing theory

```

1 clc;
2 warning('off');
3 printf("\n\n example6.5 - pg232");
4 // given
5 UzmaxbyU=24.83;

```

```

6 roUbyv=2312;
7 Re=100000;
8 // using the formula Et/v=95.5*((r/ro)/slope)-1
9 // from fig 6.6 at Re=100000
10 rbyro=[0 0.040 0.100 0.200 0.300 0.4 0.5 0.6 0.7 0.8
11 0.9 0.960 1];
12 slope=[0 0.105 0.112 0.126 0.144 0.168 0.201 0.252
13 0.336 0.503 1.007 2.517 94.59];
14 for i=2:13
15 Etbyv(i)=95.5*((rbyro(i))/slope(i))-1;
16 end
17 clf;
18 xtitle("eddy viscosity ratio versus dimensionless
19 radius","r/ro","Et/v");
20 plot(rbyro,Etbyv);

```

---

### Scilab code Exa 6.9 friction factor

```

1 clc;
2 warning("off");
3 printf("\n\n example6.9 - pg258");
4 // given
5 spg=0.84;
6 p=0.84*62.4; // [lbf/ft^3] - density
7 dP=80*144; // [lbf/ft^2] - pressure
8 dz=2000; // [ft] - length of pipe
9 gc=32.174; // [(lbm*ft)/(lbf*sec^2)] - gravitational
10 // conversion constant
11 dpbydz=-dP/dz;
12 do=2.067/12; // [ft]
13 U=2000*(1/24)*(1/3600)*(42)*(1/7.48)*(1/0.02330);
14 // using the formula f=((do/2)*(-dp/dz)*gc)/(p*(U)
15 // ^2)
16 f=((do/2)*(-dpbydz)*gc)/(p*(U)^2)
17 disp(f,"f=");

```



# Chapter 7

## integral methods of analysis

**Scilab code Exa 7.2** the integral mass balance

```
1 clc;
2 warning(" off");
3 printf("\\n\\n example7.2 - pg273");
4 // given
5 id=4; // [m] - inside diameter
6 h=2; // [m] - water level
7 ro=0.03; // [m] - radius of exit hole
8 rt=id/2; // [m] - inside radius
9 g=9.80665; // [m/sec^2] - gravitational acceleration
10 // using the equation dh/h^(1/2)=-((ro^2)/(rt^2))
    *(2*g)^(1/2)dt and integrating between h=2 and h
    =1
11 t1=integrate('(1/h^(1/2))*(1/(-((ro^2)/(rt^2))*(2*g)
    ^(1/2))))','h',2,1);
12 printf("\\n\\n Time required to remove half of the
    contents of the tank is \\n t=%fsec=%fmin",t1,t1
    /60);
13 // integrating between h=2 and h=0
14 t2=integrate('(1/h^(1/2))*(1/(-((ro^2)/(rt^2))*(2*g)
    ^(1/2))))','h',2,0);
15 printf("\\n\\n Time required to empty the tank fully
```

```
    is \n t=%fsec=%fmin" ,t2 ,t2/60) ;
```

---

### Scilab code Exa 7.3 integral balance on an individual species

```
1 clc;
2 warning(" off");
3 printf("\n\n example7.3 - pg274");
4 // given
5 // composition of fuel gas
6 nH2=24;
7 nN2=0.5;
8 nCO=5.9;
9 nH2S=1.5;
10 nC2H4=0.1;
11 nC2H6=1;
12 nCH4=64;
13 nCO2=3.0;
14 // calculating the theoritical amount of O2 required
15 nO2theoreq=12+2.95+2.25+0.30+3.50+128;
16 // since fuel gas is burned with 40% excess O2, then
   O2 required is
17 nO2req=1.4*nO2theoreq;
18 nair=nO2req/0.21; // as amount of O2 in air is 21%
19 nN2air=nair*(0.79); // as amount of N2 in air is 79%
20 nN2=nN2+nN2air;
21 nO2=nO2req-nO2theoreq;
22 nH2O=24+0+0.2+3.0+128;
23 nCO2formed=72.1;
24 nCO2=nCO2+nCO2formed;
25 nSO2=1.5;
26 ntotal=nSO2+nCO2+nO2+nN2+nH2O;
27 mPSO2=(nSO2/ntotal)*100;
28 mPCO2=(nCO2/ntotal)*100;
29 mPO2=(nO2/ntotal)*100;
```

```

30 mpN2=(nN2/ntotal)*100;
31 mpH2O=(nH2O/ntotal)*100;
32 printf("\n\n gas          N2
           O2          H2O          CO2
           SO2");
33 printf("\n\n moles          %f      %f
           %f      %f", nN2, nO2, nH2O, nCO2, nSO2);
34 printf("\n\n mole percent    %f      %f
           %f      %f", mpN2, mpO2, mpH2O, mpCO2,
           mpSO2);

```

---

### Scilab code Exa 7.4 integral momentum balance

```

1 clc;
2 warning("off");
3 printf("\n\n example7.4 - pg280");
4 // given
5 id=6; // [inch] - inlet diameter
6 od=4; // [inch] - outlet diameter
7 Q=10; // [ft^3/sec] - water flow rate
8 alpha2=%pi/3; // [radians] - angle of reduction of
                 elbow
9 alpha1=0;
10 p1=100; // [psi] - absolute inlet pressure
11 p2=29; // [psi] - absolute outlet pressure
12 S1=(%pi*((id/12)^2))/4;
13 S2=(%pi*((od/12)^2))/4;
14 U1=Q/S1;
15 U2=Q/S2;
16 mu=6.72*10^-4; // [lb*ft^-1*sec^-1]
17 p=62.4; // [lb/ft^3]
18 Nrei=((id/12)*U1*p)/(mu);
19 disp(Nrei,"Nre(inlet)=");
20 Nreo=((od/12)*U2*p)/(mu);
21 disp(Nreo,"Nre(outlet)=");

```

```

22 // thus
23 b=1;
24 w1=p*Q; // [lb/sec] - mass flow rate
25 w2=w1;
26 gc=32.174;
27 // using the equation (w/gc)*((U1)*(cos(alpha1))-(U2)
28 // *(cos(alpha2)))+p1*S1*cos(alpha1)-p2*S2*cos(
29 // alpha2)+Fextx=0;
30 Fextx=-(w1/gc)*((U1)*(cos(alpha1))-(U2)*(cos(alpha2))
31 // )-p1*144*S1*cos(alpha1)+p2*144*S2*cos(alpha2);
32 disp(Fextx,"Fext,x=");
33 Fexty=-(w1/gc)*((U1)*(sin(alpha1))-(U2)*(sin(alpha2))
34 // )-p1*144*S1*sin(alpha1)+p2*144*S2*sin(alpha2);
35 disp(Fexty,"Fext,y=");
36 printf("\n\n the forces Fext,x and Fext,y are the
37 forces exerted on the fluid by the elbow.Fext,x
38 acts to the left and Fext,y acts in the positive
39 y direction. Note that the elbow is horizontal ,and
40 gravity acts in the z direction");

```

---

### Scilab code Exa 7.5 integral momentum balance

```

1 clc;
2 warning("off");
3 printf("\n\n example7.5 - pg 282");
4 // given
5 Fextx=-2522; // [lb] - force in x direction
6 Fexty=2240; // [lb] - force in y direction
7 // the force exerted by the elbow on the fluid is
7 // the resolution of Fext,x and Fext,y , therefore
8 Fext=((Fextx)^2+(Fexty)^2)^(1/2);
9 alpha=180+atan(Fexty/Fextx)*(180/%pi);
10 printf("\n\n the force has a magnitude of %flb and a
11 direction of %f from the positive x direction (in
12 the second quadrant",Fext,alpha);

```

---

### Scilab code Exa 7.6 integral momentum balance

```
1 clc;
2 warning("off");
3 printf("\n\n example7.6 - pg283");
4 // given
5 id=6; // [inch] - inlet diameter
6 od=4; // [inch] - outlet diameter
7 Q=10; // [ft^3/sec] - water flow rate
8 alpha2=%pi/3; // [radians] - angle of reduction of
elbow
9 alpha1=0;
10 p1=100; // [psi] - absolute inlet pressure
11 p2=29; // [psi] - absolute outlet pressure
12 patm=14.7; // [psi] - atmospheric pressure
13 p1gauge=p1-patm;
14 p2gauge=p2-patm;
15 S1=(%pi*((id/12)^2))/4;
16 S2=(%pi*((od/12)^2))/4;
17 U1=Q/S1;
18 U2=Q/S2;
19 p=62.4; // [lb/ft^3]
20 b=1;
21 w1=p*Q; // [lb/sec] - mass flow rate
22 w2=w1;
23 gc=32.174;
24 // using the equation Fpress=p1gauge*S1-p2gauge*S2*
cos(alpha2);
25 Fpressx=p1gauge*144*S1-p2gauge*144*S2*cos(alpha2);
26 Fpressy=p1gauge*144*S1*sin(alpha1)-p2gauge*144*S2*
sin(alpha2);
27 wdeltaUx=(w1/gc)*((U2)*(cos(alpha2))-(U1)*(cos(
alpha1)));
28 wdeltaUy=(w1/gc)*((U2)*(sin(alpha2))-(U1)*(sin(
```

```

        alpha));
29 Fextx=wdeltaUx-Fpressx;
30 Fexty=wdeltaUy-Fpressy;
31 Fext=((Fextx)^2+(Fexty)^2)^(1/2);
32 alpha=180+(atan(Fexty/Fextx))*(180/%pi);
33 printf("\n\n The force has a magnitude of %flb and a
         direction of %f from the positive x direction(in
         the second quadrant",Fext,alpha);
34 printf("\n\n Also there is a force on the elbow in
         the z direction owing to the weight of the elbow
         plus the weight of the fluid inside");

```

---

### Scilab code Exa 7.7 integral energy balance

```

1 clc;
2 warning("off");
3 printf("\n\n example7.7 - pg293");
4 // given
5 Uo=1; // [m/sec]
6 // using Ux/Uo=y/yo
7 // assuming any particular value of yo will not
   change the answer, therefore
8 yo=1;
9 Uxavg=integrate('(Uo*y)/yo','y',0,yo);
10 Ux3avg=integrate('((Uo*y)/yo)^3','y',0,yo);
11 // using the formula alpha=(Uxavg)^3/Ux3avg
12 alpha=(Uxavg)^3/Ux3avg;
13 disp(alpha,"alpha=");
14 printf("\n\n Note that the kinetic correction factor
         alpha has the same final value for laminar pipe
         flow as it has for laminar flow between parallel
         plates.");

```

---

### Scilab code Exa 7.8 integral energy balance

```
1 clc;
2 warning("off");
3 printf("\n\n example7.8 - pg293");
4 // given
5 Q=0.03; // [m^3/sec] - volumetric flow rate
6 id=7; // [cm] - inside diameter
7 deltaz=-7; // [m] - length of pipe
8 T1=25; // [degC] - lowere side temperature
9 T2=45; // [degC] - higher side temperature
10 g=9.81; // [m/sec^2] - acceleration due to gravity
11 deltaP=4*10^4; // [N/m^2] - pressure loss due to
    friction
12 p=1000; // [kg/m^3] - density of water
13 w=Q*p;
14 C=4184; // [J/kg*K] - heat capacity of water
15 deltaH=w*C*(T2-T1);
16 // using the formula Qh=deltaH+w*g*deltaz
17 Qh=deltaH+w*g*deltaz;
18 printf("\n\n the duty on heat exchanger is \n Q=%6eJ
    /sec",Qh);
```

---

### Scilab code Exa 7.10 the energy equation and the engineering bernoulli equation

```
1 clc;
2 warning("off");
3 printf("\n\n example7.10 - pg298");
4 // given
5 d=0.03; // [m] - diameter
6 g=9.784; // [m/sec] - acceleration due to gravity
7 deltaz=-1;
8 // using the equation (1/2)*(U3^2/alpha3-U1^2/alpha1
    )+g*deltaz=0
```

```

9 // assuming
10 alpha1=1;
11 alpha3=1;
12 // also since the diameter of the tank far exceeds
    the diameter of the hose , the velocity at point
    1 must be negligible when compared to the
    velocity at point 3
13 U1=0;
14 U3=(-2*g*deltaz+(U1^2)/alpha1)^(1/2);
15 p=1000; // [kg/m^3] - density of water
16 S3=(%pi/4)*(d)^2
17 w=p*U3*S3;
18 printf("\n\n the mass flow rate is \n w=%fkg/sec",w)
;
19 // the minimum pressure in the siphon tube is at the
    point 2. Before the result of 3.13 kg/sec is
    accepted as the final value , the pressure at
    point 2 must be calculated in order to see if the
    water might boil at this point
20 // using deltap=p*((U3^2)/2+g*deltaz)
21 deltap=p*((U3^2)/2+g*deltaz);
22 p1=1.01325*10^5; // [N/m^2] - is equal to
    atmospheric pressure
23 p2=p1+deltap;
24 vp=0.02336*10^5;
25 if p2>vp then
26     printf("\n\n the siphon can operate since the
        pressure at point 2 is greater than the value
        at which the liquid boils");
27 else
28     printf("\n\n the siphon cant operate since the
        pressuer at point 2 is less than the value at
        which the liquid boils");
29
30 end

```

---

**Scilab code Exa 7.11** the energy equation and the engineering bernoulli equation

```

1 clc;
2 warning("off");
3 printf("\n\n example7.11 - pg300");
4 // given
5 sp=1.45; // specific gravity of trichloroethylene
6 pwater=62.4; // [lb/ft ^3] - density of water
7 p=sp*pwater;
8 d1=1.049; // [inch] - density of pipe at point 1
9 d2=0.6; // [inch] - density of pipe at point 2
10 d3=1.049; // [inch] - density of pipe at point 3
11 // using the formula U1*S1=U2*S2; we get U1=U2*(d2/
    d1);
12 // then using the bernoulli equation deltap/p=(1/2)
    *(U2^2-U1^2);
13 deltap=4.2*(144); // [lb/ft ^2] - pressure difference
14 U2=((2*(deltap/p)*(1/(1-(d2/d1)^4)))^(1/2))*(32.174)
    ^(1/2);
15 // using the formula w=p*U2*S
16 w=p*U2*((%pi/4)*(0.6/12)^2);
17 w1=w/(2.20462);
18 printf("\n\n the mass flow rate is \n w=%f lb/sec\n
        or in SI units \n w=%f kg/sec",w,w1);

```

---

**Scilab code Exa 7.12** the mechanical energy equation and the engineering bernoulli equation

```

1 clc;
2 warning("off");
3 printf("\n\n example7.12 - pg301");

```

```

4 // given
5 Q=50/(7.48*60); // [ ft/sec ] - volumetric flow rate
       of water
6 d1=1; // [ inch ] - diameter of pipe
7 deltaz=-5; // [ ft ] - distance between end of pipe
       and tank
8 g=32.1; // [ ft/sec ] - acceleration due to gravity
9 Cp=1; // [ Btu/lb*F ] - heat capacity of water
10 p=62.4; // [ lb/ft^3 ] - density of water
11 S1=(%pi/4)*(d1/12)^2;
12 U1=Q/S1;
13 w=p*Q;
14 U2=0;
15 gc=32.174;
16 // using the formula deltaH=(w/2)*((U2)^2-(U1)^2)+w*
       g*deltaz
17 deltaH=-(w/(2*gc))*((U2)^2-(U1)^2)-w*(g/gc)*deltaz;
18 disp(deltaH);
19 deltaH=deltaH/778; // converting from ftlb/sec to
       Btu/sec
20 deltaT=deltaH/(w*Cp);
21 printf("\n\n The rise in temperature is %fdegF",
       deltaT);

```

---

**Scilab code Exa 7.13** the mechanical energy equation and the engineering bernoulli equation

```

1 clc;
2 warning(" off");
3 printf("\n\n example7.13 - pg303");
4 // given
5 deltaz=30; // [ ft ] - distance between process and
       the holding tank
6 Q=100; // [gpm] - volumetric flow rate of water
7 p1=100; // [ psig ]

```

```

8 p2=0; // [psig]
9 g=32.1; // [ft/sec] - acceleration due to gravity
10 sv=0.0161; // [ft^3/lb] - specific volume of water
11 p=1/sv; // [lb/ft^3] - density of water
12 e=0.77; // efficiency of centrifugal pump
13 deltap=(p1-p2)*(144); // [lbf/ft^2]
14 gc=32.174;
15 // using the equation deltap/p+g*(deltaz)+Ws=0;
16 Wst=-deltap/p-(g/gc)*(deltaz);
17 // using the formula for efficiency e=Ws(theoretical)
    // /Ws(actual)
18 // therefore
19 Wsa=Wst/e;
20 // the calculated shaft work is for a unit mass flow
    rate of water, therefore for given flow rate
    multiply it by the flow rate
21 w=(Q*p)/(7.48*60);
22 Wsactual=Wsa*w;
23 power=-Wsactual/(778*0.7070);
24 printf("\n\n the required horsepower is %fhp",power)
;
```

---

**Scilab code Exa 7.14** the mechanical energy equation and the engineering bernoulli equation

```

1 clc;
2 warning("off");
3 printf("\n\n example7.14 - pg304");
4 // given
5 p1=5; // [atm] - initial pressure
6 p2=0.75; // [atm] - final pressure after expansion
    through turbine
7 T=450; // [K] - temperature
8 y=1.4; // cp/cv for nitrogen
9 // using the equation Ws=-(y/(y-1))*(p1/density1)*((
```

```

    p2/p1)^(y-1)/y)-1)
10 R=8314; // gas constant
11 p1bydensity=R*T;
12 Ws=-(y/(y-1))*(p1bydensity)*((p2/p1)^(y-1)/y)-1);
13 printf("\n\n the shaft work of the gas as it expands
        through the turbine and transmits its molecular
        energy to the rotating blades is \n Ws=%eJ/kmol",
        Ws);

```

---

### Scilab code Exa 7.15 manometers

```

1 clc;
2 warning("off");
3 printf("\n\n example 7.15 - pg311");
4 // given
5 T=273.15+25; // [K] - temperature
6 R=8.314; // [kPa*m^3/kmol*K] - gas constant
7 p=101.325; // [kPa] - pressure
8 M=29; // molecular weight of gas
9 pa=(p*M)/(R*T);
10 sg=13.45; // specific gravity
11 pm=sg*1000;
12 g=9.807; // [m/sec^2] - acceleration due to gravity
13 deltaz=15/100; // [m]
14 // using the equation p2-p1=deltap=(pm-pa)*g*deltaz;
15 deltap=-(pm-pa)*g*deltaz;
16 printf("\n\n the pressure drop is %eN/m^2",deltap);
17 printf("\n the minus sign means the upstream
        pressure p1 is greater than p2, i.e ther is a
        pressure drop.");

```

---

### Scilab code Exa 7.16 manometers

```

1 clc;
2 warning(" off");
3 printf("\n\n example7.16 - pg312");
4 // given
5 T=536.67; // [degR] - temperature
6 R=10.73; // [(lbf/in^2*ft^3)*lb*mol^-1*degR] - gas
    constant
7 p=14.696; // [lbf/in^2];
8 g=9.807*3.2808; // [ft/sec^2] - acceleration due to
    gravity
9 M=29; // molecular weight of gas
10 pa=(p*M)/(R*T);
11 sg=13.45; // specific gravity
12 pm=sg*62.4;
13 deltaz=15/(2.54*12); // [ft]
14 gc=32.174;
15 // using the equation p2-p1=deltap=(pm-pa)*g*deltaz
16 deltap=(pm-pa)*(g/gc)*deltaz;
17 printf("\n\n the pressure drop is %flbf/ft^2",deltap
);

```

---

### Scilab code Exa 7.18 manometers

```

1 clc;
2 warning(" off");
3 printf("\n\n example7.18 - pg315");
4 // given
5 at=0.049; // [in^2] - cross sectional area of the
    manometer tubing
6 aw=15.5; // [in^2] - cross sectional area of the
    well
7 g=32.174; // [ft/sec^2] - acceleration due to
    gravity
8 gc=32.174;
9 sg=13.45; // [ specific garvity of mercury

```

```

10 p=62.4; // [lb / ft ^ 3] – density of water;
11 pm=sg*p;
12 deltaz_waterleg=45.2213;
13 // using the equation A(well)*deltaz(well)=A(tube)*
    deltaz(tube)
14 deltazt=70; // [cm]
15 deltazw=deltazt*(at/aw);
16 deltaz=deltazt+deltazw;
17 deltap_Hg=-pm*(g/gc)*(deltaz/(2.54*12));
18 disp(deltap_Hg);
19 deltazw=45.2213; // [cm]
20 deltap_tap=deltap_Hg+p*(g/gc)*(deltazw/(12*2.54));
21 printf("\n\n deltap_tap=%f lbf / ft ^ 2",deltap_tap);
22 printf("\ndeltap is negative and therefore p1 is
        greater than p2");

```

---

### Scilab code Exa 7.19 buoyant forces

```

1 clc;
2 warning("off");
3 printf("\n\n example7_19 – pg317");
4 // given
5 p=749/760; // [atm]
6 T=21+273.15; // [K]
7 R=82.06; // [atm*cm^3/K] – gas constant
8 v=(R*T)/p; // [cm^3/mole] – molar volume
9 M=29; // [g/mole] – molecular weight
10 pair=M/v;
11 m_air=53.32; // [g]
12 m_h2o=50.22; // [g]
13 ph2o=0.998; // [g/cm^3] – density of water
14 V=(m_air-m_h2o)/(ph2o-pair); // [cm^3]
15 density=m_air/V;
16 printf("\n\n The density of coin is \n density=%fg/
        cm^3",density);

```

```
17 printf("\n\n Consulting a handbook it is seen that  
this result is correct density for gold");
```

---

### Scilab code Exa 7.20 buoyant forces

```
1 clc;  
2 warning("off");  
3 printf("\n\n example7.20 - pg318");  
4 // given  
5 P=749/760; // [atm] - pressure  
6 T=21+273.15; // [K] - temperature  
7 poak=38*(1/62.4); // [g/cm^3] - density of oak  
8 pbrass=534/62.4; // [g/cm^3] - density of brass  
9 m_brass=6.7348; // [g]  
10 pair=0.001184; // [g/cm^3] - density of air  
11 // using the formula m_oak=m_brass*((1-(pair/pbrass)  
// (1-(pair/poak)))  
12 m_oak=m_brass*((1-(pair/pbrass))/(1-(pair/poak)));  
13 printf("\n\n m_oak=%fg",m_oak);
```

---

### Scilab code Exa 7.21 variation of pressure with depth

```
1 clc;  
2 warning("off");  
3 printf("\n\n example7.21 - pg320");  
4 // given  
5 T=545.67; // [degR] - temperature  
6 R=1545; // [Torr*ft^3/degR*mole] - gas constant  
7 M=29; // [g/mole] - molecular weight  
8 g=9.807; // [m/sec^2] - acceleration due to gravity  
9 gc=9.807;  
10 po=760; // [Torr] - pressure  
11 deltaz=50; // [ft]
```

```

12 // using the equation p=po*exp(-(g/gc)*M*(deltaz/R*T
    ))
13 p=po*%e^(-(g/gc)*M*(deltaz/(R*T)));
14 printf("\n\n p=%fTorr\n Thus, the pressure decrease
        for an elevation of 50ft is very small",p);

```

---

**Scilab code Exa 7.22** variation of pressure with depth

```

1 clc;
2 warning("off");
3 printf("\n\n example7.22 - pg321");
4 // given
5 To=545.67; // [degR] - air temperature at beach
    level
6 betaa=-0.00357; // [degR/ft] - constant
7 R=1545; // [Torr*ft^3/degR*mole] - gas constant
8 M=29;
9 deltaz=25000; // [ft]
10 // using the equation ln(p/po)=((M)/(R*betaa))*ln(To
    /(To+betaa*deltaz))
11 p=po*exp(((M)/(R*betaa))*log(To/(To+betaa*deltaz)));
12 printf("\n\n p=%fTorr",p);
13 // using the equation T=To+betaa*deltaz
14 T=To+betaa*deltaz;
15 printf("\n\n T=%fdegR",T);

```

---

# Chapter 9

## agitation

**Scilab code Exa 9.3** scale up procedures for turbulent flow with a single test volume

```
1 clc;
2 warning("off");
3 printf("\n\n example9.3 - pg389");
4 Nblades=4; // no. of blades
5 d=9/12; // [ft] - diameter of the impeller
6 dt=30/12; // [ft] - diameter of the tank
7 Nbaffles=4; // no. of baffles
8 h=30; // [inch] - height of unit
9 mu=10; // [cP] - viscosity of fluid
10 sg=1.1; // specific gravity of fluid
11 s=300; // [rpm] - speed of agitator
12 CbyT=0.3;
13 V=(%pi*dt^3)/4; //volume of tank in ft^3
14 V1=V*7.48; // [gal] - volume of tank in gallons
15 mu=mu*(6.72*10^-4); // [lb/ft*sec]
16 p=sg*62.4; // [lb/ft^3] - density of fluid
17 N=s/60; // [rps] - impeller speed in revolutions per
second
18 Nre=((d^2)*N*p)/mu;
19 disp(Nre,"Nre=");
```

```

20 printf("\n\n Therefore the agitator operates in the
           turbulent region");
21 Npo=1.62;
22 gc=32.174;
23 P=(Npo*(p*(N^3)*(d^5)))/(gc*550);
24 Cf=63025;
25 Tq=(P/s)*Cf;
26 PbyV=P/V;
27 PbyV1=P/V1;
28 TqbyV=Tq/V;
29 TqbyV1=Tq/V1;
30 printf("\n\n The power per unit volume and the
           torque per unit volume is \n P/V=%f hp/ ft ^3=%f hp
           /gal\n Tq/V=%f in*lb/ ft ^3=%f in*lb/ gal" ,PbyV ,
           PbyV1 ,TqbyV ,TqbyV1 );

```

---

**Scilab code Exa 9.4** scale up procedures for turbulent flow with a single test volume

```

1 clc;
2 warning(" off");
3 printf("\n\n example9.4 - pg391");
4 // given
5 Tpilot=30;
6 Tlab=10;
7 N1=690;
8 N2=271;
9 D2=3;
10 D1=1;
11 n=(log(N1/N2))/(log(D2/D1));
12 V=12000/7.48; // [ft ^3]
13 T=((4*V)/pi)^(1/3); // [ft ]
14 R=12.69/(30/12);
15 N3=N2*(1/R)^n; // [rpm] - impeller speed in the
                    reactor

```

```

16 disp(N3," impeller speed in rpm=");
17 D3=0.75*R; // [ft] - reactor impeller diameter
18 disp(D3," reactor impeller diameter in ft=");
19 P=0.1374*((N3/N2)^3)*(R^5);
20 disp(P," power in hp=");
21 Cf=63025;
22 Tq=(P/N3)*Cf; // [inch*lb]
23 disp(Tq," torque in inch*lb=");
24 printf("\n\n At this point, the design is complete.
         A standard size impeller would be chosen as well
         as a standard size motor(7.5 hp or 10 hp)");

```

---

**Scilab code Exa 9.5** scale up procedures for turbulent flow with a single test volume

```

1 clc;
2 warning(" off");
3 printf("\n\n example9.5 - pg 393");
4 // given
5 n=[0.5 0.6 0.7 0.8 0.9 1.0];
6 D2=3.806;
7 D1=0.25;
8 R=D2/D1;
9 N1=690;
10 N2=N1*((D1/D2)^n);
11 P1=9.33*10^-3; // [hp]
12 P2=P1*R^(5-3*n);
13 printf("\n\n n          N, rpm
           P, hp");
14 for i=1:6
15 printf("\n %f          %f" ,n(i),
           N2(i),P2(i));
16 end

```

---

# Chapter 10

## transport in ducts

Scilab code Exa 10.1 laminar pipe flow

```
1 clc;
2 warning(" off");
3 printf(" \n\n example10.1 - pg405");
4 T=30; // [degC] - temperature
5 d=8.265*10^-4; // [m] - diameter of the capillary
viscometer
6 deltapbyL=-0.9364; // [psi/ft] - pressure drop per
unit length
7 deltapbyL=deltapbyL*(2.2631*10^4); // [kg/m^2*sec^2]
- pressure drop per unit length
8 Q=28.36*(10^-6)*(1/60);
9 p=(0.88412-(0.92248*10^-3)*T)*10^3; // [kg/m^3] -
density
10 s=(%pi*(d^2))/4;
11 U=Q/s;
12 tauw=(d/4)*(-deltapbyL);
13 shearrate=(8*U)/d;
14 mu=tauw/(shearrate);
15 printf(" \n\n The viscosity is \n mu=%f kg/m*sec=%f
cP" ,mu ,mu*10^3);
16 printf(" \n\n Finally , it is important to check the
```

```

    reynolds number to make sure the above equation
    applies");
17 Nre=(d*U*p)/(mu);
18 disp(Nre,"Nre=");
19 printf("\n\n The flow is well within the laminar
    region and therefore the above equation applies")
;

```

---

### Scilab code Exa 10.2 turbulent pipe flow

```

1 clc;
2 warning("off");
3 printf("\n\n example10.2 - pg407");
4 Nreold=1214;
5 Uold=0.8810;
6 Nre=13700;
7 U=Uold*(Nre/Nreold);
8 Lbyd=744;
9 // using the newton raphson method to calculate the
   value of f from the equation - 1/(f^(1/2))=4*log(
   Nre*(f^(1/2)))-0.4
10 f=0.007119;
11 p=(0.88412-(0.92248*10^-3)*T)*10^3; // [kg/m^3] -
   density
12 tauw=(1/2)*p*(U^2)*f;
13 deltap=tauw*(4)*(Lbyd);
14 d=0.03254/12; // [ft]
15 L=Lbyd*d;
16 printf("\n\n Pressure drop is \n -deltap=%e N/m^2=%f
   kpa=130 psi",deltap,deltap*10^-3);
17 printf("\n\n A pressure drop of 130 psi on a tube of
   length of %f ft is high and shows the
   impracticality of flows at high reynolds number
   in smaller tubes",L);

```

---

### Scilab code Exa 10.3 pressure drop in rough pipes

```
1 clc;
2 warning("off");
3 printf("\n\n example10.3 - pg414");
4 // given
5 u=1/60; // [m/sec] - velocity
6 p=1000; // [kg/m^3] - density
7 mu=1*10^-3; // [kg/m*sec] - viscosity
8 d=6*10^-2; // [m] - inside diameter of tube
9 L=300; // [m] - length of the tube
10 Nre=(d*u*p)/(mu);
11 disp(" therefore the flow is laminar",Nre,"Nre=");
12 f=16/Nre;
13 disp(f);
14 deltap=(4*f)*(L/d)*((p*(u^2))/2);
15 printf("\n\n -deltap=%f N/m^2 = %f kPa = %e psi",
deltap,deltap*10^-3,deltap*1.453*10^-4);
```

---

### Scilab code Exa 10.4 pressure drop in rough pipes

```
1 clc;
2 warning("off");
3 printf("\n\n example10.4 - pg415");
4 // given
5 d=6*10^-2; // [m] - inside diameter of tube
6 p=1000; // [kg/m^3] - density
7 // for smooth pipe
8 Nre=[10^4 10^5];
9 f=[0.0076 0.0045];
10 mu=10^-3; // [kg/m^2*s]
11 U=(Nre*mu)/(d*p);
```

```

12 L=300; // [m] – length of the tube
13 for i=1:2
14 deltap(i)=(4*f(i))*(L/d)*((p*(U(i)^2))/2);
15 end
16 disp(" for smooth pipe");
17 printf(" Nre %f -deltap");
18 printf("\n %f ,Nre(1),deltap(1)
19 );
20 printf("\n %f %f \n",Nre(2),deltap
21 (2));
22 // for commercial steel
23 Nre=[10^4 10^5];
24 f=[0.008 0.0053];
25 U=(Nre*mu)/(d*p);
26 L=300; // [m] – length of the tube
27 for i=1:2
28 deltap(i)=(4*f(i))*(L/d)*((p*(U(i)^2))/2);
29 end
30 disp(" for commercial steel pipe");
31 printf(" Nre %f -deltap");
32 printf("\n %f ,Nre(1),deltap(1)
33 );
34 printf("\n %f %f \n",Nre(2),deltap
35 (2));
36 // for cast iron pipe
37 Nre=[10^4 10^5];
38 f=[0.009 0.0073];
39 U=(Nre*mu)/(d*p);
40 L=300; // [m] – length of the tube
41 for i=1:2
42 deltap(i)=(4*f(i))*(L/d)*((p*(U(i)^2))/2);
43 end
44 disp(" for cast iron pipe");
45 printf(" Nre %f -deltap");
46 printf("\n %f ,Nre(1),deltap(1)
47 );
48 printf("\n %f %f ",Nre(2),deltap(2));

```

---

### Scilab code Exa 10.5 von karman number

```
1 clc;
2 warning(" off");
3 printf("\n\n example10.5 - pg417");
4 // given
5 L=300; // [m] - length of pipe
6 d=0.06; // [m] - inside diameter
7 deltap=147*10^3; // [Pa] - pressure the pump can
    supply
8 ebyd=0.000762; // relative roughness
9 p=1000; // [kg/m^3] - density
10 mu=1*10^-3; // [kg/m*sec] - viscosity
11 tauw=(d*(deltap))/(4*L);
12 // using the hit and trial method for estimation of
    flow velocity
13 // let
14 f=0.005;
15 U=((2*tauw)/(p*f))^(1/2);
16 Nre=(d*U*p)/mu;
17 // from the graph value of f at the above calculated
    reynolds no. and the given relative roughness(e/
        d)
18 f=0.0054;
19 U=((2*tauw)/(p*f))^(1/2);
20 Nre=(d*U*p)/mu;
21 // from the graph value of f at the above calculated
    reynolds no. and the given relative roughness(e/
        d)
22 f=0.0053;
23 U=((2*tauw)/(p*f))^(1/2);
24 Nre=(d*U*p)/mu;
25 // from the graph value of f at the above calculated
    reynolds no. and the given relative roughness(e/
```

```

        d)
26 f=0.0053;
27 // At this point the value of f is deemed unchanged
   from the last iteration .Hence, the values
   obtained after the third iteration are the
   converged values
28 printf("\n\n The maximum flow velocity is \n U=%f m/
   sec",U);

```

---

### Scilab code Exa 10.6 von karman number

```

1 clc;
2 warning("off");
3 printf("\n\n example10.6 - pg419");
4 // given
5 L=300; // [m] - length of pipe
6 d=0.06; // [m] - inside diameter
7 deltap=147*10^3; // [Pa] - pressure the pump can
   supply
8 ebyd=0.000762; // relative roughness
9 p=1000; // [kg/m^3] - density
10 mu=1*10^-3; // [kg/m*sec] - viscosity
11 Nvk=((d*p)/mu)*((d*(deltap))/(2*L*p))^(1/2);
12 disp(Nvk,"von karman no.-");
13 // From the fig at given von karman no and relative
   roughness the value of f is-
14 f=0.0055;
15 Nre=Nvk/(f^(1/2))
16 U=(Nre*mu)/(d*p);
17 printf("\n\n U=%f m/sec",U);

```

---

### Scilab code Exa 10.7 the velocity head concept

```

1 clc;
2 warning(" off");
3 printf("\n\n example10.7 - pg422");
4 // given
5 L=300; // [m] - length of pipe
6 d=0.06; // [m] - inside diameter
7 p=1000; // [kg/m^3] - density
8 mu=1*10^-3; // [kg/m*sec] - viscosity
9 Nre=[10^4 10^5];
10 U=(Nre*mu)/(d*p);
11 velocityhead=(U^2)/2;
12 N=(L/d)/45; // no of velocity heads
13 deltap=p*N*(velocityhead);
14 for i=1:2
15     disp(Nre(i),"Nre=");
16     printf("\n\n velocity head =%f m^2/sec^2",
17            velocityhead(i));
17     printf("\n\n -deltap = %f kPa = %f psi",deltap(i)
18           )*10^-3,deltap(i)*1.453*10^-4);
18 end

```

---

### Scilab code Exa 10.8 pipe fittings and valves

```

1 clc;
2 warning(" off");
3 printf("\n\n example10.8 - pg439");
4 // given
5 mu=6.72*10^-4; // [lb/ft*sec] - viscosity
6 p=62.4; // [lb/ft^3] - density
7 S=0.03322; // [ft^2] - flow area
8 d=0.206; // [ft]
9 e=1.5*10^-4; // absolute roughness for steel pipe
10 ebyd=e/d;
11 Nre=10^5;
12 // friction factor as read from fig in book for the

```

```

        given reynolds no. and relative roughness is-
13 f=0.0053;
14 U=(Nre*mu)/(p*d);
15 Q=U*S;
16 gc=32.174;
17 // (a) equivalent length method
18 deltapbyL=f*(4/d)*(p*(U^2))*(1/(2*gc))*(6.93*10^-3);
19 // using L=Lpipe+Lfittings+Lloss;
20 Lfittings=2342.1*d;
21 kc=0.50; // due to contraction loss
22 ke=1; // due to enlargement loss
23 Lloss=(kc+ke)*(1/(4*f))*d;
24 Lpipe=137;
25 L=Lpipe+Lfittings+Lloss;
26 deltap=deltapbyL*L;
27 patm=14.696; // [psi] - atmospheric pressure
28 p1=patm+deltap;
29 printf("\n\n (a)The inlet pressure is \n p1=%f psi",
       p1);
30 // (b) loss coefficient method
31 // using the equation deltap/p=-(Fpipe+Ffittings+
   Floss)
32 L=137;
33 kfittings=52.39;
34 sigmaF=((4*f*(L/d))+kc+ke+kfittings)*((U^2)/(2*gc));
35 deltap=(p*sigmaF)/(144);
36 p1=patm+deltap;
37 printf("\n\n (b)The inlet pressure is \n p1=%f psi",
       p1);
38 printf("\n\n Computation of the pressure drop by the
           loss coefficient method differs from the
           equivalent length method by less than 1 psi");

```

---

### Scilab code Exa 10.9 gases

```

1 clc;
2 warning(" off");
3 printf("\n\n example10.9 - pg443");
4 // given
5 L1=50; // [m] - length of first pipe
6 L2=150; // [m] - length of second pipe
7 L3=100; // [m] - length of third pipe
8 d1=0.04; // [m] - diameter of first pipe
9 d2=0.06; // [m] - diameter of second pipe
10 d3=0.08; // [m] - diameter of third pipe
11 deltap=-1.47*10^5; // [kg/m*sec] - pressure drop
12 mu=1*10^-3; // [kg/m*sec] - viscosity
13 p=1000; // [kg/m^3] - density
14 // for branch 1
15 S=(%pi*(d1^2))/4;
16 Nvk=((d1*p)/mu)*(-(d1*deltap)/(2*L1*p))^(1/2);
17 f=(1/(4*log10(Nvk)-0.4))^2;
18 U=(((deltap)/p)*(d1/L1)*(2/4)*(1/f))^(1/2);
19 w1=p*U*S;
20 printf("\n\n For first branch w1=%f kg/sec",w1);
21 // for branch 2
22 S=(%pi*(d2^2))/4;
23 Nvk=((d2*p)/mu)*(-(d2*deltap)/(2*L2*p))^(1/2);
24 f=(1/(4*log10(Nvk)-0.4))^2;
25 U=(((deltap)/p)*(d2/L2)*(2/4)*(1/f))^(1/2);
26 w2=p*U*S;
27 printf("\n\n For second branch w2=%f kg/sec",w2);
28 // for branch 3
29 S=(%pi*(d3^2))/4;
30 Nvk=((d3*p)/mu)*(-(d3*deltap)/(2*L3*p))^(1/2);
31 f=(1/(4*log10(Nvk)-0.4))^2;
32 U=(((deltap)/p)*(d3/L3)*(2/4)*(1/f))^(1/2);
33 w3=p*U*S;
34 printf("\n\n For third branch w3=%f kg/sec",w3);
35 // total flow rate w=w1+w2+w3
36 w=w1+w2+w3;
37 printf("\n\n total flow rate is w=%f kg/sec",w);

```

---

### Scilab code Exa 10.11 complex fluid flow systems

```
1 clc;
2 warning(" off");
3 printf("\n\n example10.11 - pg 447");
4 // given
5 sp=1.1;
6 p=sp*62.4; // [lb/ft^3] - density
7 mu=2*6.72*10^-4; // [lb/ft*sec] - viscosity
8 Q=400; // [gpm] - volumetric flow rate
9 e=1.5*10^4; // roughness of steel pipe
10 gc=32.174;
11 kexit=1;
12 kentrance=0.5;
13 // 4 in schedule pipe
14 d=4.026/12; // [ ft ]
15 U4=Q/39.6; // [ ft / sec ]
16 Lgv=13.08;
17 Lg1v=114.1;
18 Le=40.26;
19 Lpipe_4=22;
20 Lfittings_4=Lgv+Lg1v+Le;
21 Lloss=0;
22 L_4=Lpipe_4+Lfittings_4+Lloss;
23 Nre_4=(d*U4*p)/mu;
24 f=0.00475;
25 Fpipe_4=((4*f*L_4)/d)*(U4^2)*(1/(2*gc));
26 Floss_4=((kentrance+0)*(U4^2))/(2*gc);
27 // 5 in schedule pipe
28 d=5.047/12;
29 U5=Q/62.3;
30 Lgv=10.94;
31 Le=75.71;
32 Lpipe_5=100;
```

```

33 Lfittings_5=Lgv+Le;
34 Lloss=0;
35 L_5=Lpipe_5+Lfittings_5+Lloss;
36 Nre=(d*U5*p)/mu;
37 f=0.00470;
38 Fpipe_5=((4*f*L_5)/d)*(U5^2)*(1/(2*gc));
39 Floss_5=((kexit+0)*(U5^2))/(2*gc);
40 // 6 in schedule pipe
41 d=6.065/12;
42 U6=Q/90;
43 Lgv=6.570;
44 Le=30.36;
45 Lpipe_6=4;
46 Lfittings_6=Lgv+Le;
47 Lloss=0;
48 L_6=Lpipe_6+Lfittings_6+Lloss;
49 Nre=(d*U6*p)/mu;
50 f=0.00487;
51 Fpipe_6=((4*f*L_6)/d)*(U6^2)*(1/(2*gc));
52 kc=0.50;
53 Floss_6=kc*((U6^2)/(2*gc));
54 Ffittings=0;
55 deltap_6=p*(Fpipe_6+Ffittings+Floss_6);
56 // 3/4 in 18 gauge tube
57 d=0.652112/12;
58 L_3by4=15;
59 U_3by4=(Q*0.962)/100;
60 Floss_3by4=100*(kexit+kentrance)*((U_3by4^2)/2);
61 Nre=d*U_3by4*p*(1/mu);
62 // clearly the flow is turbulent
63 f=0.08*((Nre)^(-1/4))+0.012*((d)^(1/2));
64 deltap_3by4=((4*f*p*L_3by4)/d)*((U_3by4^2)/(2*gc));
65 Fpipe_3by4=100*((4*f*L_3by4)/d)*((U_3by4^2)/(2*gc));
66 deltap_spraysystem=25; // [psi]
67 Fspraysystem=(deltap_spraysystem/p)*(144);
68 delta_p=[p*(kexit+kentrance)]*[(U_3by4^2)/(2*gc)];
69 Fpipe=Fpipe_4+Fpipe_5+Fpipe_6;
70 Floss=Floss_4+Floss_5+Floss_6+Floss_3by4;

```

```

71 ws=0+([(15^2)-0]/[2*gc])+38.9+382.5;
72 w=(Q*p)/(7.48);
73 Ws=(ws*w)/(33000);
74 efficiency=0.6;
75 Ws_actual=Ws/efficiency
76 printf("\n\n The power supplied to the pump is\n"
    "W_actual = %f",Ws_actual);

```

---

### Scilab code Exa 10.12 complex fluid flow systems

```

1 clc;
2 warning("off");
3 printf("\n\n example10.12 - pg454");
4 // given
5 kexit=1;
6 kentrance=0.5;
7 Q=400; // [gpm] - volumetric flow rate
8 gc=32.174;
9 // for 4 inch pipe
10 d=4.026; // [inch]
11 L=22; // [ft]
12 Lbyd=(L*12)/(d);
13 // adding the contributions due to fittings
14 Lbyd=Lbyd+3*13+340+4*30;
15 N=Lbyd/45;
16 N=N+kentrance+0;
17 U4=Q/39.6; // [ft/sec]
18 Fpipe_4=(N*(U4^2))/(2*gc);
19 printf("\n\n F(4 in. pipes) = %f ft*lbf/lbm",Fpipe_4)
    ;
20 // for 5 inch pipe
21 L=100; // [ft]
22 d=5.047; // [inch]
23 Lbyd=(L*12)/(d);
24 // valves contributes 26 diameters and six elbows

```

```

        contribute 30 diameters each; therefore
25 Lbyd=Lbyd+26+6*30;
26 N=Lbyd/45; // no. of velocity heads
27 N=N+kexit+kentrance;
28 U5=Q/62.3;
29 Fpipe_5=(N*(U5^2))/(2*gc);
30 printf("\n\n F(5 in. pipes) = %f ft*lb/ft/lbm",Fpipe_5)
;
31 // for 6 inch pipe
32 d=6.065; // [inch]
33 L=5; // [ft]
34 Lbyd=(L*12)/(d);
35 // adding the contributions due to fittings
36 Lbyd=Lbyd+1*13+2*30;
37 N=Lbyd/45;
38 N=N+0+kentrance;
39 U6=Q/90;
40 Fpipe_6=(N*(U6^2))/(2*gc);
41 printf("\n\n F(6 in. pipes) = %f ft*lb/ft/lbm",Fpipe_6)
;
42 F_largepipes=Fpipe_4+Fpipe_5+Fpipe_6;
43 printf("\n\n F(large pipes) = %f ft*lb/ft/lbm",
F_largepipes);

```

---

### Scilab code Exa 10.14 non circular conduits

```

1 clc;
2 warning("off");
3 printf("\n\n example10.14 - pg459");
4 // given
5 l=0.09238;
6 rh=0.1624*l;
7 L=300;
8 de=4*rh;
9 p=1000; // [kg/m^3]

```

```

10 mu=10^-3; // [ kg/m* sec ]
11 Uavg=1.667;
12 Nre=(de*Uavg*p)/mu;
13 f=0.0053;
14 deltap=((4*f*L)/de)*(p*(Uavg^2)*(1/2));
15 printf("\n\n -deltap = %e kg/m*s = %e N/m^2 = %f kPa
",deltap,deltap,deltap*10^-3);

```

---

### Scilab code Exa 10.15 orifice meter

```

1 clc;
2 warning("off");
3 printf("\n\n example10.15 - pg466");
4 // given
5 Q=400; // [gpm]
6 p=1.1*62.4; // [lbm/ft ^3]
7 mu=2*(6.72*10^-4); // [lb / ft *sec]
8 e=1.5*10^4;
9 // 4 inch schedule pipe
10 d=0.3355;
11 S=(%pi*(d^2))/4;
12 U4=Q/39.6;
13 ebyd=e/d;
14 w=3671/60;
15 pm=13.45*62.4;
16 g=32.1;
17 gc=32.174;
18 deltz=2.5;
19 deltap=(g/gc)*(pm-p)*(deltaz);
20 betaa=((1)/(1+[(2*p*gc)*(deltap)]*(((0.61*S)/w)^2)))^(1/4);
21 d2=betaa*d;
22 Nre2=(4*w)/(%pi*d2*mu);
23 a=(1/30)*4.026;
24 b=(1/4)*(2.013-1.21);

```

```

25 c=(1/8)*(2.42);
26 if a<b then
27   if a<c then
28     opt=a;
29   else
30     opt=c;
31 end
32 else
33   if b<c then
34     opt=b;
35   else
36     opt=c;
37 end
38 end
39 printf("\n\n The pertinent orifice details are \n
          orifice diameter = %f in \n corner taps , square
          edge\n orifice plate not over %f in thick",d2*12,
          opt);

```

---

### Scilab code Exa 10.16 venturi and nozzle

```

1 clc;
2 warning(" off");
3 printf("\n\n example10.16 - pg470");
4 // given
5 Q=400; // [gpm]
6 p=1.1*62.4; // [lbm/ft ^3]
7 mu=2*(6.72*10^-4); // [lb / ft * sec ]
8 e=1.5*10^4;
9 // 4 inch schedule pipe
10 d=0.3355;
11 S=(%pi*(d^2))/4;
12 U4=Q/39.6;
13 ebyd=e/d;
14 w=3671/60;

```

```

15 pm=13.45*62.4;
16 g=32.1;
17 gc=32.174;
18 Nre=(d*U4*p)/mu;
19 if Nre>10^4 then
20     c=0.98;
21 end
22 deltaz=2.5;
23 deltap=(g/gc)*(pm-p)*(deltaz);
24 betaa=((1)/(1+[(2*p*gc)*(deltap)]*((c*S)/w)^2)))^
    (1/4);
25 d2=betaa*d;
26 printf("\n\n The pertinentr details of the venturi
    design are\n Throat diameter = %f inch\n Approach
    angle = 25\n Divergence angle = 7",d2*12);

```

---

### Scilab code Exa 10.17 pitot tube

```

1 clc;
2 warning("off");
3 printf("\n\n example10.17 - pg477");
4 // given
5 Uzmax=3.455; // [ ft/sec ]
6 m=32;
7 a1=-0.3527;
8 a2=-0.6473;
9 rbyro=0.880;
10 UzbyUzmax=1+a1*(rbyro^2)+a2*(rbyro^(2*m));
11 Uz=Uzmax*(UzbyUzmax);
12 Uzavg=(4/9)*Uzmax+(5/18)*(Uz+Uz);
13 printf("\n\n the average velocity is \n Uzavg = %f
    ft/sec \n\n Thus, in this example there is an
    inherent error of 5.5 percent , even before any
    experimental errors are introduced",Uzavg);

```

---

# Chapter 11

## heat and mass transfer in duct flow

Scilab code Exa 11.1 conduction

```
1 clc;
2 warning("off");
3 printf("\n\n example11.1 - pg497");
4 // given
5 K_drywall=0.28; // [Btu/ft*degF] - thermal
    conductivity of dry wall
6 K_fibreglass=0.024; // [Btu/ft*degF] - thermal
    conductivity of fibre glass
7 K_concrete=0.5; // [Btu/ft*degF] - thermal
    conductivity of concrete
8 T4=0; // [degF]
9 T1=65; // [degF]
10 deltaT=T4-T1; // [degF]
11 a=1; // [ft^2] - assuming area of 1 ft^2
12 deltax1=0.5/12; // [ft]
13 deltax2=3.625/12; // [ft]
14 deltax3=6/12; // [ft]
15 R1=deltax1/(K_drywall*a); // [h*degF/Btu]
16 R2=deltax2/(K_fibreglass*a); // [h*degF/Btu]
```

```

17 R3=deltax3/(K_concrete*a); // [h*degF/Btu]
18 qx=deltaT/(R1+R2+R3);
19 q12=-qx;
20 q23=-qx;
21 q34=-qx;
22 deltaT1=(-q12)*deltax1*(1/(K_drywall*a));
23 T2=T1+deltaT1;
24 deltaT2=(-q23)*deltax2*(1/(K_fibreglass*a));
25 T3=T2+deltaT2;
26 deltaT3=(-q34)*deltax3*(1/(K_concrete*a));
27 T4=T3+deltaT3;
28 printf("\n\n T1 = %f degF\n T2 = %f degF\n T3 = %f
degF\n T4 = %f degF",T1,T2,T3,T4);

```

---

### Scilab code Exa 11.2 the resistance concept

```

1 clc;
2 warning("off");
3 printf("\n\n example11.2 - pg501");
4 // given
5 r1=(2.067/2)/(12); // [ft]
6 r2=r1+0.154/12; // [ft]
7 r3=r2+3/12; // [ft]
8 L=1; // [ft]
9 Ka=26; // [Btu/h*ft*degF]
10 Kb=0.04; // [Btu/h*ft*degF]
11 T1=50; // [degF]
12 Ra=(log(r2/r1))/(2*pi*L*Ka);
13 Rb=(log(r3/r2))/(2*pi*L*Kb);
14 R=Ra+Rb;
15 deltaT=-18; // [degF] - driving force
16 Qr=-(deltaT/(R));
17 disp(Qr);
18 deltaT1=(-Qr)*(Ra);
19 T2=T1+deltaT1;

```

```
20 printf("\n\n The interface temperature is \n T2 = %f  
degF",T2);
```

---

### Scilab code Exa 11.3 the resistance concept

```
1 clc;  
2 warning("off");  
3 printf("\n\n example11.3 - pg502");  
4 // given  
5 Ra=8.502*10^-4; // [h*degF*Btu^-1]  
6 Rb=5.014; // [h*degF*Btu^-1]  
7 r1=(2.067/2)/(12); // [ft]  
8 r2=r1+0.154/12; // [ft]  
9 r3=r2+3/12; // [ft]  
10 d1=2*r1;  
11 d0=2*r3;  
12 h0=25; // [Btu/h*ft^2*degF]  
13 h1=840; // [Btu/h*ft^2*degF]  
14 L=1; // [ft] - considering 1 feet length  
15 R0=1/(h0*pi*d0*L);  
16 R1=1/(h1*pi*d1*L);  
17 R=R0+R1+Ra+Rb;  
18 disp(R);  
19 deltaT=-400; // [degF]  
20 Qr=-(deltaT)/R;  
21 disp(Qr);  
22 // the heat loss calculated above is the heat loss  
// per foot.therefore for 500 ft  
23 L=500;  
24 Qr=Qr*L;  
25 printf("\n\n the heat loss for a 500 feet pipe is \n  
qr = %e Btu/h",Qr);
```

---

### Scilab code Exa 11.5 heat and mass transfer during turbulent flow

```
1 clc;
2 warning("off");
3 printf("\n\n example11.5 - pg521");
4 // given
5 Nre=50000;
6 d=0.04; // [m] - diameter of pipe
7 // physical properties of water
8 T1=293.15; // [K]
9 T2=303.15; // [K]
10 T3=313.15; // [K]
11 p1=999; // [kg/m^3] - density of water at
    temperature T1
12 p2=996.0; // [kg/m^3] - density of water at
    temperature T2
13 p3=992.1; // [kg/m^3] - density of water at
    temperature T3
14 mu1=1.001; // [cP] - viscosity of water at
    temperature T1
15 mu2=0.800; // [cP] - viscosity of water at
    temperature T2
16 mu3=0.654; // [cP] - viscosity of water at
    temperature T3
17 k1=0.63; // [W/m*K] - thermal conductivity of water
    at temperature T1
18 k2=0.618; // [W/m*K] - thermal conductivity of water
    at temperature T2
19 k3=0.632; // [W/m*K] - thermal conductivity of water
    at temperature T3
20 cp1=4182; // [J/kg*K] - heat capacity of water at
    temperature T1
21 cp2=4178; // [J/kg*K] - heat capacity of water at
    temperature T2
22 cp3=4179; // [J/kg*K] - heat capacity of water at
    temperature T3
23 Npr1=6.94; // prandtl no. at temperature T1
24 Npr2=5.41; // prandtl no. at temperature T2
```

```

25 Npr3=4.32; // prandtl no. at temperature T3
26 // (a) Dittus -Boelter-this correction evaluates all
      properties at the mean bulk temperature ,which is
      T1
27 kmb=0.603
28 h=(kmb/d)*0.023*((Nre)^(0.8))*((Npr1)^0.4);
29 printf("\n\n (a) Dittus -Boelter\n the heat transfer
      coefficient is \n h = %f W/m^2*K = %f Btu/ft ^2*h
      ^-1*degF",h,h*0.17611);
30 // (b) Seider Tate-this correlation evaluates all
      the properties save muw at the mean bulk
      temperature
31 h=(kmb/d)*(0.027)*((Nre)^0.8)*((Npr1)^(1/3))*((mu1/
      mu3)^0.14);
32 printf("\n\n (b) Seider Tate\n the heat transfer
      coefficient is \n h = %f W/m^2*K = %f Btu/ft ^2*h
      ^-1*degF",h,h*0.17611);
33 // (c) Sleicher-Rouse equation
34 a=0.88-(0.24/(4+Npr3));
35 b=(1/3)+0.5*exp((-0.6)*Npr3);
36 Nref=Nre*(mu1/mu2)*(p2/p1);
37 Nnu=5+0.015*((Nref)^a)*((Npr3)^b);
38 h=Nnu*(kmb/d);
39 printf("\n\n (c) Sleicher-Rouse equation\n the heat
      transfer coefficient is \n h = %f W/m^2*K = %f
      Btu/ft ^2*h^-1*degF",h,h*0.17611);
40 // (d) Colbum Analogy- the j factor for heat
      transfer is calculated
41 jh=0.023*((Nref)^(-0.2));
42 Nst=jh*((Npr2)^(-2/3));
43 U=(Nre*mu1*10^-3)/(d*p1);
44 h=Nst*(p1*cp1*U);
45 printf("\n\n (d) Colbum Analogy\n the heat transfer
      coefficient is \n h = %f W/m^2*K = %f Btu/ft ^2*h
      ^-1*degF",h,h*0.17611);
46 // (e) Friend-Metzner
47 f=0.005227;
48 Nnu=((Nre)*(Npr1)*(f/2)*((mu1/mu3)^0.14))

```

```

    /(1.20+((11.8)*((f/2)^(1/2))*(Npr1-1)*(Npr1)
    ^(-1/3)))) ;
49 h=Nnu*(kmb/d) ;
50 printf("\n\n (e) Friend-Metzner\n the heat transfer
        coefficient is \n h = %f W/m^2*K = %f Btu/ft^2*h
        ^-1*degF" ,h,h*0.17611) ;
51 // (f) Numerical analysis
52 Nnu=320;
53 h=Nnu*(kmb/d) ;
54 printf("\n\n (f) Numerical analysis\n the heat
        transfer coefficient is \n h = %f W/m^2*K = %f
        Btu/ft^2*h^-1*degF" ,h,h*0.17611) ;

```

---

### Scilab code Exa 11.6 heat and mass transfer during turbulent flow

```

1 clc;
2 warning(" off");
3 printf("\n\n example11.6 - pg525");
4 // given
5 Tw=680; // [K] - temperature at the wall
6 Tb=640; // [K] - temperature at the bulk
7 Tf=(Tw+Tb)/2; // [K]
8 Nre=50000;
9 vmb=2.88*10^-7;
10 vf=2.84*10^-7;
11 Nref=Nre*(vmb/vf);
12 k=27.48;
13 d=0.04;
14 // from table 11.3 the prandtl no. is
15 Npr=8.74*10^-3
16 // constant heat flow
17 Nnu=6.3+(0.0167)*((Nref)^0.85)*((Npr)^0.93);
18 h=Nnu*(k/d);
19 printf("\n\n constant heat flow\n h = %f W/m^2*K =
        %f Btu/ft^2*h*degF" ,h,h*0.17611);

```

```

20 // constant wall temperature
21 Nnu=4.8+0.0156*((Nref)^0.85)*((Npr)^0.93);
22 h=Nnu*(k/d);
23 printf("\n\n constant wall temperature\n h = %f W/m
^2*K = %f Btu/ft ^2*h*degF",h,h*0.17611);

```

---

### Scilab code Exa 11.7 double pipe heat exchangers simple solutions

```

1 clc;
2 warning("off");
3 printf("\n\n example11.7 - pg536");
4 // given
5 di=0.620; // [inch] - internal diameter
6 d0=0.750; // [inch] - outer diameter
7 Ai=0.1623; // [ft ^2/ft]
8 Ao=0.1963; // [ft ^2/ft]
9 wc=12*(471.3/0.9425);
10 cp=1; // [Btu/lbm*degF] - heat capacity of water
11 Tco=110;
12 Tci=50;
13 qtotal=wc*cp*(Tco-Tci);
14 deltaH_coldwater=3.6*10^5;
15 deltaH_vapourization=1179.7-269.59;
16 wh=deltaH_coldwater/deltaH_vapourization;
17 hi=80; // [Btu/h*ft ^2*degF]
18 ho=500; // [Btu/h*ft ^2*degF]
19 km=26; // [Btu/h*ft*degF]
20 Ui=1/((1/hi)+((Ai*log(d0/di))/(2*pi*km))+(Ai/(Ao*ho
    )));
21 disp(Ui)
22 deltaT1=300-50;
23 deltaT2=300-110;
24 LMTD=(deltaT1-deltaT2)/log(deltaT1/deltaT2));
25 A=qtotal/(Ui*LMTD);
26 L=A/Ai;

```

```
27 printf("\n\n the length of the heat exchanger is \n"
L = %f ft",L);
```

---

### Scilab code Exa 11.8 double pipe heat exchangers simple solutions

```
1 clc;
2 warning("off");
3 printf("\n\n example11.8 - pg537");
4 // given
5 L=30; // [ft] - length
6 Ai=0.1623*L;
7 di=0.620; // [inch] - internal diameter
8 d0=0.750; // [inch] - outer diameter
9 Ao=0.1963*L; // [ft^2/ft]
10 wc=12*(471.3/0.9425);
11 cp=1; // [Btu/lbm*degF] - heat capacity of water
12 deltaH_coldwater=3.6*10^5;
13 deltaH_vapourization=1179.7-269.59;
14 wh=deltaH_coldwater/deltaH_vapourization;
15 hi=80; // [Btu/h*ft^2*degF]
16 ho=500; // [Btu/h*ft^2*degF]
17 km=26; // [Btu/h*ft*degF]
18 Ui=1/((1/hi)+((Ai/L)*log(d0/di))/(2*pi*km))+(Ai/(
    Ao*ho));
19 deltaT1=300-50;
20 deltaT=deltaT1/(exp((Ui*Ai)/(wc*cp)));
21 Tsat=300;
22 Tc2=Tsat-deltaT;
23 printf("\n\n Therefore , the outlet temperature of
the cold fluid is \n Tc2 = %f degF",Tc2);
```

---

### Scilab code Exa 11.9 double pipe heat exchangers simple solutions

```

1 clc;
2 warning(" off");
3 printf("\n\n example11.9 - pg538");
4 // given
5 Ai=4.869;
6 wc=6000;
7 cp=1;
8 Rf=0.002;
9 Uclean=69.685;
10 Udirty=1/(Rf+(1/Uclean));
11 deltaT1=300-50;
12 deltaT2=deltaT1/(exp((Udirty*Ai)/(wc*cp)));
13 Th2=300;
14 Tc2=Th2-deltaT2;
15 printf("\n\n the outlet temperature is \n Tc2 = %f
degF",Tc2);

```

---

### Scilab code Exa 11.10 multipass heat exchangers equipment

```

1 clc;
2 warning(" off");
3 printf("\n\n example11.10 - pg544");
4 // given
5 Ui=325; // [W/m^2*K] - overall heat transfer
           coefficient
6 Thi=120; // [degC] - inlet temperature of
            hydrocarbon
7 Tho=65; // [degC] - outlet temperature of
            hydrocarbon
8 Tci=15; // [degC] - inlet temperature of water
9 Tco=50; // [degC] - outlet temperture of water
10 cp=4184; // [J/kg*K] - heat capacity of water
11 ch=4184*0.45; // [J/kg*K] - heat capacity of
                  hydrocarbon
12 wc=1.2; // [kg/sec] - mass flow rate of water

```

```

13 wh=((wc*cp)*(Tco-Tci))/((ch)*(Thi-Tho));
14 qtotal=wc*cp*(Tco-Tci);
15 // (a) - parallel double pipe
16 F=1;
17 Thi=120; // [degC] - inlet temperature of
    hydrocarbon
18 Tho=65; // [degC] - outlet temperature of
    hydrocarbon
19 Tci=15; // [degC] - inlet temperature of water
20 Tco=50; // [degC] - outlet temperture of water
21 deltaT1=Thi-Tci;
22 deltaT2=Tho-Tco;
23 LMTD=(deltaT2-deltaT1)/(log(deltaT2/deltaT1));
24 Ai=qtotal/((Ui*LMTD));
25 printf("\n\n (a) parallel double pipe \n Ai = %f m^2
    ",Ai);
26 // (b) - counter flow
27 F=1;
28 Thi=120; // [degC] - inlet temperature of
    hydrocarbon
29 Tho=65; // [degC] - outlet temperature of
    hydrocarbon
30 Tco=15; // [degC] - inlet temperature of water
31 Tci=50; // [degC] - outlet temperture of water
32 deltaT1=Thi-Tci;
33 deltaT2=Tho-Tco;
34 LMTD=(deltaT2-deltaT1)/(log(deltaT2/deltaT1));
35 Ai=qtotal/((Ui*LMTD));
36 printf("\n\n (b) counter flow \n Ai = %f m^2",Ai);
37 // (c) - 1-2 shell and tube
38 Thi=120; // [degC] - inlet temperature of
    hydrocarbon
39 Tho=65; // [degC] - outlet temperature of
    hydrocarbon
40 Tci=15; // [degC] - inlet temperature of water
41 Tco=50; // [degC] - outlet temperture of water
42 Z=(Thi-Tho)/(Tco-Tci);
43 nh=(Tco-Tci)/(Thi-Tci);

```

```

44 deltaT1=Thi-Tco;
45 deltaT2=Tho-Tci;
46 F=0.92;
47 LMTD=(F*(deltaT2-deltaT1))/(log(deltaT2/deltaT1));
48 Ai=qtotal/((Ui*LMTD));
49 printf("\n\n (c) 1-2 shell and tube \n Ai = %f m^2"
      ,Ai);
50 // (d) - 2-4 shell and tube
51 Thi=120; // [degC] - inlet temperature of
            hydrocarbon
52 Tho=65; // [degC] - outlet temperature of
            hydrocarbon
53 Tci=15; // [degC] - inlet temperature of water
54 Tco=50; // [degC] - outlet temperture of water
55 Z=(Thi-Tho)/(Tco-Tci);
56 nh=(Tco-Tci)/(Thi-Tci);
57 F=0.975;
58 LMTD=(F*(deltaT2-deltaT1))/(log(deltaT2/deltaT1));
59 Ai=qtotal/((Ui*LMTD));
60 printf("\n\n (d) 2-4 shell and tube \n Ai = %f m^2"
      ,Ai);

```

---

# Chapter 12

## transport past immersed bodies

Scilab code Exa 12.2 the laminar boundary layer

```
1 clc;
2 warning("off");
3 printf("\n\n example12.2 - pg562");
4 p=1.2047*0.06243; // [lb/ft^3]
5 mu=(18.17*10^-6)*(0.6720); // [lb/ft*sec]
6 v=mu/p;
7 x=2; // [ft]
8 U=6; // [ft/sec]
9 Nre=(x*U)/v;
10 disp("The Reynolds number is well within the laminar
      region",Nre,"Nre=");
11 del=5*x*(Nre)^(-1/2);
12 C1=0.33206;
13 Cd=2*C1*(Nre)^(-1/2);
14 L2=2; // [ft]
15 L1=1; // [ft]
16 b=1;
17 F=((2*(C1)*U*b))*((mu*p*U)^(1/2))*(((L2)^(1/2))-((L1
      )^(1/2)));
18 gc=32.174;
19 F=F/gc;
```

```
20 printf("\n\n The value of F properly expressed in  
force units is \n F=%e lbf",F);
```

---

### Scilab code Exa 12.3 turbulent boundary layer

```
1 clc;  
2 warning("off");  
3 printf("\n\n example12.3 - pg569");  
4 U=3; // [m/sec]  
5 x1=1; // [m]  
6 x2=2; // [m]  
7 p=1/(1.001*10^-3); // [kg/m^3];  
8 mu=1*10^-3; // [kg/m*sec]  
9 Nre1=(x1*U*p)/(mu);  
10 Nre2=(x2*p*U)/(mu);  
11 tauw=(1/2)*(p*(U^2))*((2*log10(Nre1)-0.65)^(-2.3));  
12 B=1700;  
13 Cd=(0.455*(log10(Nre2))^-2.58)-(B/(Nre2));  
14 Lb=2.0;  
15 F=(1/2)*(p*(U^2))*(Lb)*(Cd);  
16 printf("\n\n the drag on the plate is \n F = %f kg*m  
/ sec^2 = %f N",F,F);
```

---

### Scilab code Exa 12.5 heat and mass transfer during boundary layer flow past a flat plate

```
1 clc;  
2 warning("off");  
3 printf("\n\n example12.5 - pg576");  
4 T=290; // [K] - temperature of flowing water  
5 U=3; // [m/sec] - free stream velocity  
6 Tf_s=285; // [K] - temperature of free stream  
7 vr=10^-3; // [m^3/kg] - volume per unit mass
```

```

8 p=1/vr; // [kg/m^3] - density of water at Tfs
9 mu=1225*10^-6; // [N*sec/m^2]
10 k=0.590; // [W/m*K]
11 Npr=8.70;
12 // (a) The length of laminar boundary
13 Nre=5*10^5;
14 xc=(Nre)*(mu/(p*U));
15 printf("\n\n (a) The length of laminar boundary is \
n xc = %f m",xc);
16 // (b) Thickness of the momentum boundary layer and
   thermal boundary layer
17 del=5*xc*((Nre)^(-1/2));
18 delh=del*((Npr)^(-1/3));
19 printf("\n\n (b) The thickness of momentum boundary
   layer is \n del = %e m\n The thickness of the
   hydrodynamic layer is \n delh = %e m",del,delh);
20 // (c) Local heat transfer coefficient
21 x=0.2042; // [ft]
22 hx=((0.33206*k)/(x))*((Nre)^(1/2))*((Npr)^(1/3));
23 printf("\n\n (c) The local heat transfer coefficient
   is \n h = %f W/m^2*K = %f Btu/hr*ft^2*degF",hx,
   hx*0.17611);
24 // (d) Mean heat transfer coefficient
25 hm=2*hx;
26 printf("\n\n (d) The mean heat transfer coefficient
   is \n h = %f W/m^2*K = %f Btu/hr*ft^2*degF",hm,hm
   *0.17611);

```

---

### Scilab code Exa 12.10 stokes flow past a sphere

```

1 clc;
2 warning("off");
3 printf("\n\n example12.10 - pg590");
4 // given
5 T=293.15; // [K]

```

```

6 pp=999; // [kg/m^3] - density of water
7 mu=0.01817*10^-3; // [kg/m*sec] - viscosity of air
8 p=1.205; // [kg/m^3] - density of air
9 d=5*10^-6; // [m] - particle diameter
10 g=9.80; // [m/sec^2]
11 rp=d/2;
12 Ut=((2*g*(rp^2))*(pp-p))/(9*mu);
13 Nre=(d*Ut*p)/(mu);
14 // clearly the flow is in the stokes law region at
   this low reynolds number; therefore , the drag
   force is
15 Fp=6*pi*mu*rp*Ut;
16 printf("\n\n The drag force is \n Fp = %e N",Fp);

```

---

### Scilab code Exa 12.11 drag coefficient correlations

```

1 clc;
2 warning("off");
3 printf("\n\n example12.11 - pg591");
4 // given
5 T=293.15; // [K]
6 pp=999; // [kg/m^3] - density of water
7 mu=0.01817*10^-3; // [kg/m*sec] - viscosity of air
8 p=1.205; // [kg/m^3] - density of air
9 d=5*10^-6; // [m] - particle diameter
10 g=9.80; // [m/sec^2]
11 rp=d/2;
12 Ut=((2*g*(rp^2))*(pp-p))/(9*mu);
13 Nre=(d*Ut*p)/(mu);
14 t=(-2*(rp^2)*pp)/(9*mu)*(log(1-0.99));
15 printf("\n\n Time for the drop of water in previous
   example from an initial velocity of zero to 0.99*
   Ut is \n t = %e sec",t);
16 printf("\n\n In other words , the drop accelerates
   almost instantaneously to its terminal velocity")

```

;

---

### Scilab code Exa 12.12 drag coefficient correlations

```
1 clc;
2 warning("off");
3 printf("\n\n example12.12 - pg 594");
4 // given
5 pp=1.13*10^4; // [kg/m^3] - density of lead particle
6 p=1.22; // [kg/m^3] - density of air
7 g=9.80; // [m/sec^2] - acceleration due to gravity
8 d=2*10^-3; // [m] - diameter of particle
9 mu=1.81*10^-5; // [kg/m*sec] - viscosity of air
10 // let us assume
11 Cd=0.44;
12 Ut=((4*d*g*(pp-p))/(3*p*Cd))^(1/2);
13 disp(Ut)
14 Nre=(Ut*d*p)/(mu);
15 // from fig 12,16 value of Cd is
16 Cd=0.4;
17 Ut=((4*d*g*(pp-p))/(3*p*Cd))^(1/2);
18 Nre=(Ut*d*p)/(mu);
19 // Within the readability of the chart Cd is
    unchanged and therefore the above obtained Cd is
    the final answer
20 printf("\n\n The terminal velocity is \n Ut = %f m/
    sec", Ut);
```

---

### Scilab code Exa 12.13 drag coefficient correlations

```
1 clc;
2 warning("off");
3 printf("\n\n example12.13 - pg595");
```

```

4 // given
5 distance=1/12; // [ft]
6 time=60; // [sec]
7 Ut=distance/time;
8 mu=1.68; // [lb/ft*sec] - viscosity
9 pp=58; // [lb/ft^3] - density of sphere
10 p=50; // [lb/ft^3] - density of polymer solution
11 g=32; // [ft/sec] - acceleration due to gravity
12 rp=((9*mu)*(Ut)*((2*g)^(-1))*((pp-p)^(-1)))^(1/2);
13 printf("\n\n The required particle diameter would be
        about %f inch",rp*2*12);
14 Nre=(rp*2*Ut*p)/(mu);
15 disp(Nre,"Nre=");
16 printf("\n\n This reynolds number is well within the
        stokes law region ; thus the design is
        reasonable");

```

---

### Scilab code Exa 12.14 liquid solid fluidization

```

1 clc;
2 warning("off");
3 printf("\n\n example12.14 - pg616");
4 // given
5 T=842; // [degF] - temperature
6 P=14.6; // [psia] - pressure
7 p=0.487; // [kg/m^3] - density of air
8 mu=3.431*10^-5; // [kg/m*sec] - viscosity of air
9 k=0.05379; // [W/m*K] - thermal conductivity
10 Npr=0.7025; // prandtl no.
11 // (a) static void fraction
12 mcoal=15*2000; // [lb] - mass of coal
13 pcoal=94; // [lbm/ft^3] - density of coal
14 d=10; // [ft]
15 L=7; // [ft]
16 area=("%pi*(d^2))/4";

```

```

17 Vcoal=mcoal/pcoal;
18 Vtotal=area*L;
19 e=(Vtotal-Vcoal)/(Vtotal);
20 disp(e,"(a) The void fraction is E=");
21 // (b) minimum void fraction and bed height
22 d=200; // [um] – particle diameter
23 Emf=1-0.356*((log10(d))-1);
24 // this value seems to be a lottle low and therefore
25 // 0.58 will be used
25 Emf=0.58;
26 Lmf=((L)*(1-e))/(1-Emf);
27 printf("\n\n (b) The bed height is \n Lmf = %f ft",
28 Lmf);
28 // (c) Minimum fluidization velocity
29 P1=20; // [psia]
30 P2=14.696; // [psia]
31 p1=(p*P1)/(P2);
32 // the archimides no. is
33 g=9.78; // [m/sec^2]
34 Nar=p1*g*((d*10^-6)^3)*(1506-p1)*((1/(mu)^2));
35 C1=27.2;
36 C2=0.0408;
37 Nremf=((C1^2)+C2*Nar)^(1/2)-C1;
38 Umf=(Nremf*mu)/((d*10^-6)*p1);
39 printf("\n\n (c) The minimum fluidization velocity
40 is \n Umf = %f m/sec",Umf);
40 // (d) Minimum pressure
41 deltapmf=(1506-p1)*(g)*(1-Emf)*((Lmf*12*2.54)/(100))
42 +p1*g*Lmf;
42 printf("\n\n (c) The minimum pressure drop for
43 fluidization is \n -deltapmf = %e Pa",deltapmf);
43 // (e) Particle settling velocity
44 Cd=0.44;
45 Ut=((8*((d*10^-6)/2)*g)*(1506-p1))/(3*p1*Cd))^(1/2)
46 ;
46 Nrep=(Ut*d*10^-6*p1)/(mu);
47 disp(Nrep,"Nrep=");
48 // clearly at the point of minimum velocity for fast

```

fluidization , the terminal settling velocity is  
 not in the range of Newtons law. Therefore the eq  
 . for the transition region will be tried  
 49  $U_t = ((5.923/18.5) * (((d*10^{-6}) * p_1) / (\mu))^{(0.6)})$   
 $\quad \quad \quad ^{(1/(2-0.6))}$   
 50 **printf**("\n\n (e) The particle settling velocity is \\\n Ut = %f m/sec", Ut);  
 51 // (f) Bed to wall heat transfer coefficient  
 52  $N_{refb} = (d*10^{-6}) * 2.5 * U_{mf} * p_1 * (1/\mu)$ ;  
 53  $N_{nufb} = 0.6 * N_{pr} * (N_{refb})^{(0.3)}$ ;  
 54  $h_w = N_{nufb} * (k / (d*10^{-6}))$ ;  
 55 **printf**("\n\n (f) The bed to wall heat transfer  
 coefficient is \n  $h_w = %f W/m^2*K$ ",  $h_w$ );

---

### Scilab code Exa 12.15 liquid solid fluidization

```

1 clc;  

2 warning(" off");  

3 printf("\n\n example12.5 - pg618");  

4 // given  

5 pp=249.6; // [lb/ft^3] - density of catalyst  

6 p=58; // [lb/ft^3] - density of liquid  

7 g=32.174; // [ft/sec^2]  

8 gc=32.174;  

9 Lmf=5; // [ft] - height of bed  

10 mu=6.72*10^-3; // [lbm/ft*sec] - viscosity of liquid  

11 dp=0.0157/12; // [ft] - diameter of particle  

12 emf=0.45;  

13 deltapmf=(pp-p)*(g/gc)*(1-emf)*(Lmf);  

14 Nar=(p*g*dp^3)*(pp-p)*(1/(mu)^2);  

15 C1=27.2;  

16 C2=0.0408;  

17 Nremf=((C1^2)+C2*Nar)^(1/2))-C1;  

18 Umf=Nremf*(mu/(dp*p));  

19 printf("\n\n Minimum fluidization velocity is \n Umf

```

$$= \%e \text{ ft/sec}^2, U_{mf}) ;$$

---

### Scilab code Exa 12.16 single cyclinder heat transfer

```
1 clc;
2 warning("off");
3 printf("\n\n example12.16 - pg624");
4 // given
5 d=24*10^-6; // [m] - diameter of wire
6 T=415; // [K] - operating temperature of hot wire
           anemometer
7 P=0.1; // [W] - power consumption
8 L=250*d;
9 Tair=385; // [K] - temperature of air in duct
10 A=%pi*d*L;
11 Tfilm=(T+Tair)/2;
12 // properties of air at Tfilm
13 p=0.8825; // [kg/m^3]
14 mu=2.294*10^-5; // [kg/m*s]
15 cpf=1013; // [J*kg/K]
16 kf=0.03305; // [W/m*K]
17 Npr=0.703;
18 h=P/(A*(T-Tair));
19 Nnu=(h*d)/kf;
20 function y=func(x)
21     y=Nnu-0.3-((0.62*(x^(1/2))*(Npr^(1/3)))
22                  /((1+((0.4/Npr)^(2/3)))^(1/4)))*((1+((x
23                  /(2.82*(10^5)))^(5/8)))^(4/5));
24 endfunction
25 // on solving the above function for x by using some
      root solver technique like Newton raphson method
      , we get
26 x=107.7;
27 // or
28 Nre=107.7;
```

```

27 y=func(x);
28 Um=(Nre*mu)/(d*p);
29 printf("\n\n The velocity is \n Um = %f m/sec = %f
      ft/sec",Um,Um*3.28);

```

---

### Scilab code Exa 12.17 single cyclinder heat transfer

```

1 clc;
2 warning("off");
3 printf("\n\n example12.17 - pg630");
4 // given
5 dt=0.75;
6 St=1.5*dt;
7 S1=3*dt;
8 Lw=1; // [m]
9 N=12;
10 Stotalarea=N*(St/12)*Lw;
11 Sminarea=N*((St-dt)/12)*Lw*0.3048;
12 // properties of air at 293.15 K
13 p=1.204; // [kg/m^3]
14 mu=1.818*10^-5; // [kg/m*s]
15 cp=1005; // [J*kg/K];
16 k=0.02560; // [J/s*m*K]
17 Npr=(cp*mu)/k;
18 U_inf=7; // [m/sec]
19 Umax=U_inf*(St/(St-dt));
20 w=p*Umax*Sminarea;
21 C_tubes=0.05983; // [m^2/m] - circumference of the
      tubes
22 N_tubes=96;
23 Atubes=N_tubes*C_tubes*Lw;
24 Tw=328.15; // [K]
25 Tinf=293.15; // [K]
26 Tin=293.15; // [K]
27 Tout=293.15; // [K]

```

```

28 u=100;
29 while u>10^-1
30     T=(Tin+Tout)/2
31     Told=Tout;
32     p=-(0.208*(10^-3))+(353.044/T);
33     mu=-(9.810*(10^-6))+(1.6347*(10^-6)*(T^(1/2)));
34     cp=989.85+(0.05*T);
35     k=0.003975+7.378*(10^-5)*T;
36     Npr=(cp*mu)/k;
37     dt=0.75*0.0254;
38     Gmax=w/Sminarea;
39     Nre=(dt*Gmax)/mu;
40     h=0.27*(k/dt)*(Npr^0.36)*(Nre^0.63);
41     h=h*0.98;
42     deltaT=(h*Atubes*(Tw-Tinf))/(w*cp);
43     Tout=Tin+deltaT;
44     u=abs(Tout-Told);
45 end
46 T=(Tin+Tout)/2
47 p=-(0.208*(10^-3))+(353.044/T);
48 mu=-(9.810*(10^-6))+(1.6347*(10^-6)*(T^(1/2)));
49 dt=0.75;
50 dv=(4*(St*S1-(%pi*(dt^2)*(1/4))))/(%pi*dt)
    *(0.09010/3.547);
51 de=dv;
52 Nre=(dv*24.72)/mu;
53 dv=dv/(0.09010/3.547);
54 ftb=1.92*(Nre^(-0.145));
55 Zt=S1;
56 Ltb=8*S1;
57 deltap=(ftb*(24.72^2))/(2*p*(dv/Ltb)*((St/dv)^0.4)
    *((St/Zt)^0.6));
58 printf("\n\n -deltap = %f kg/m*s = %f N/m^2 = %f
    psia",deltap,deltap,deltap*(0.1614/1113));

```

---

# Chapter 13

## unsteady state transport

Scilab code Exa 13.1 heat transfer with negligible internal resistance

```
1 clc;
2 warning("off");
3 printf("\n\n example13.1 - pg651");
4 // given
5 h=12; // [W/m^2*K] - heat transfer coefficeint
6 k=400; // [W/m*K] - thermal conductivity
7 // (a) for sphere
8 r=5*10^-2; // [m] - radius of copper sphere
9 Lc=((4*pi*((r)^3))/3)/(4*pi*((r)^2));
10 Nbi=h*Lc*(1/k);
11 printf("\n\n (a) The biot no. is \n Nbi = %e",Nbi);
12 // (b) for cyclinder
13 r=0.05; // [m] - radius of cyclinder
14 L=0.3; // [m] - height of cyclinder
15 Lc=(pi*((r)^2)*L)/(2*pi*r*L);
16 Nbi=h*Lc*(1/k);
17 printf("\n\n (b) The biot no. is \n Nbi = %e",Nbi);
18 // (c) for a long square rod
19 L=.4; // [m] - length of copper rod
20 r=0.05; // [m] - radius of a cyclinder havimg same
           cross sectional area as that of square
```

```

21 x= (%pi*r^2)^(1/2));
22 Lc=((x^2)*L)/(4*x*L);
23 Nbi=h*Lc*(1/k);
24 printf("\n\n ( c ) The biot no. is \n Nbi = %e",Nbi);

```

---

**Scilab code Exa 13.6** generalized chart solution for finite slab and cylinder

```

1 clc;
2 warning("off");
3 printf("\n\n example13_6 - pg684");
4 // given
5 d=1*0.0254; // [m]
6 Lr=d/2; // [m];
7 Lz=(1.2/2)*(0.0254);
8 x=Lz;
9 r=Lr;
10 k=0.481;
11 h=20;
12 mr=k/(h*Lr);
13 mz=k/(h*Lz);
14 nr=r/Lr;
15 nz=x/Lz;
16 t=1.2; // [sec]
17 alpha=1.454*10^-4;
18 Xr=(alpha*t)/(Lr^2);
19 Xz=(alpha*t)/(Lz^2);
20 // using the above value of m,n,X the value for Ycz
   and Ycr from fig 13.14 is
21 Ycr=0.42;
22 Ycz=0.75;
23 Yc=Ycr*Ycz;
24 T_infinity=400; // [K]
25 To=295;
26 Tc=T_infinity-(Yc*(T_infinity-To));

```

```
27 printf("\n\n The temperature t the centre is \n Tc =\n %f K",Tc);
```

---

**Scilab code Exa 13.7** generalized chart solution for finite slab and cylinder

```
1 clc;
2 warning("off");
3 printf("\n\n example13_7 - pg684");
4 // given
5 T_x0=300; // [K]
6 Tw=400; // [K]
7 L=0.013; // [m]
8 alpha=2.476*(10^-5); // [m^2/sec]
9 h=600; // [W/m^2*K]
10 pcp=3.393*(10^6); // [J/m^3*K]
11 L=0.013; // [m]
12 deltax=L/10;
13 betaa=0.5;
14 deltat=0.03;
15 deltat=betaa*((deltax)^2)*(1/alpha);
16 T_infinity=400; // [K]
17 // to be sure that the solution is stable , it is
   customary to truncate this number
18 deltat=0.03; // [sec]
19 // betaa=alpha*deltat*((1/deltax)^2);
20   for i=1:11
21     Told(i)=300;
22 end
23 a=((2*h*deltat)/(pcp*deltax));
24 b=((2*alpha*deltat)/(pcp*((deltax)^2)));
25 for j=1:11
26 Tnew(1)=(T_infinity*0.08162)+(Told(1)
   *(1-0.08162-0.8791))+(Told(2)*0.8791)
27 for k=1:9
```

```

28     Tnew(k+1)=(betaa*Told(k+2))+((1-2*betaa)*(Told(k
+1)))+(betaa*Told(k));
29 end
30 Tnew(11)=((2*betaa)*(Told(10)))
31 Told=Tnew;
32 end
33 disp(Told);

```

---

### Scilab code Exa 13.9 semi infinite slab

```

1 clc;
2 warning("off");
3 printf("\n\n example 13_9 - pg700");
4 // given
5 p=2050; // [kg/m^3] - density of soil
6 cp=1840; // [J/kg*K] - heat capacity of soil
7 k=0.52; // [W/m*K] - thermal conductivity of soil
8 alpha=0.138*10^-6; // [m^2/sec]
9 t=4*30*24*3600; // [sec] - no. of seconds in 4
months
10 Tx=-5; // [degC]
11 Tinf=-20; // [degC]
12 T0=20; // [degC]
13 // from the fig 13.24 the dimensionless distance Z
is
14 Z=0.46;
15 // then the depth is
16 x=2*((alpha*t)^(1/2))*Z
17 printf("\n\n the depth is \n x = %f m = %f ft",x,x
*(3.6/1.10));

```

---

### Scilab code Exa 13.10 cylinder

```

1 clc;
2 warning(" off");
3 printf("\n\n example13.10 - pg701");
4 // given
5 d=0.01; // [m] - diameter of cylindrical porous
    plug
6 D=2*10^-9; // [m^2/sec] - diffusion coefficient
7 t=60*60; // [sec]
8 r=d/2;
9 m=0;
10 Ca_inf=0;
11 Ca_0=10;
12 X=(D*t)/((r)^2);
13 // from fig 13.14 the ordinate is
14 Y=0.7;
15 Ca_c=Ca_inf-Y*(Ca_inf-Ca_0);
16 printf("\n\n the concentration of KCL at the centre
    after 60 min is \n Ca = %f kg/m^3",Ca_c);

```

---

# Chapter 14

## estimation of transport coefficients

Scilab code Exa 14.1 kinetic theory of gases

```
1 clc;
2 warning(" off");
3 printf("\n\n example14.1 - pg726");
4 // given
5 T=40+273.15; // [K] - temperature
6 P=1; // [atm] - pressure
7 sigma=3.711*10^-10; // [m]
8 etadivkb=78.6; // [K]
9 A=1.16145;
10 B=0.14874;
11 C=0.52487;
12 D=0.77320;
13 E=2.16178;
14 F=2.43787;
15 Tstar=T/(etadivkb);
16 // using the formula si=(A/(Tstar^B))+(C/exp(D*Tstar)
17 //)+(E/exp(F*Tstar))
17 si=(A/(Tstar^B))+(C/exp(D*Tstar))+(E/exp(F*Tstar));
18 M=28.966; // [kg/mole] - molecular weight
```

```

19 // using the formula mu=(2.6693*(10^-26))*((M*T)
20 ^^(1/2))/((sigma^2)*si))
21 mu=(2.6693*(10^-26))*(((M*T)^(1/2))/((sigma^2)*si));
22 printf("\n\n The viscosity of air is \n mu=%eNs/m^2=
%fc",mu,mu*10^3);

```

---

### Scilab code Exa 14.2 non uniform gas theory

```

1 clc;
2 warning("off");
3 printf("\n\n example14.2.sce - pg726");
4 T=40+273.15; // [K] - temperature
5 P=1; // [atm] - pressure
6 // thermal conductivit of air
7 sigma=3.711*10^-10; // [m]
8 etadivkb=78.6; // [K]
9 A=1.16145;
10 B=0.14874;
11 C=0.52487;
12 D=0.77320;
13 E=2.16178;
14 F=2.43787;
15 Tstar=T/(etadivkb);
16 // using the formula si=(A/(Tstar^B))+(C/exp(D*Tstar)
17 ))+(E/exp(F*Tstar))
18 si=(A/(Tstar^B))+(C/exp(D*Tstar))+(E/exp(F*Tstar));
19 // using the formula K=(8.3224*(10^-22))*(((T/M)
20 ^^(1/2))/((sigma^2)*si))
21 M=28.966; // [kg/mole] - molecular weight of air
22 k=(8.3224*(10^-22))*(((T/M)^(1/2))/((sigma^2)*si));
23 printf("\n\n Thermal conductivity of air is \n k=%fW
/m*K",k);
24 printf("\n\n Agreement between this value and
original value is p[oor;the Chapman-Enskog theory
is in erroo when applied to thermal conductivity

```

```

        of polyatomic gases");
23 // thermal conductivity of argon
24 sigma=3.542*10^-10; // [m]
25 etadivkb=93.3; // [K]
26 A=1.16145;
27 B=0.14874;
28 C=0.52487;
29 D=0.77320;
30 E=2.16178;
31 F=2.43787;
32 Tstar=T/(etadivkb);
33 // using the formula si=(A/(Tstar^B))+(C/exp(D*Tstar))
   //)+(E/exp(F*Tstar))
34 si=(A/(Tstar^B))+(C/exp(D*Tstar))+(E/exp(F*Tstar));
35 // using the formula K=(8.3224*(10^-22))*(((T/M)
   ^^(1/2))/((sigma^2)*si))
36 M=39.948; // [kg/mole] - molecular weight of argon
37 k=(8.3224*(10^-22))*(((T/M)^(1/2))/((sigma^2)*si));
38 printf("\n\n Thermal conductivity of argon is \n k=
   %fW/m*K",k);
39 printf("\n\n The thermal conductivity from Chapman-
   Enskog theory agrees closely with the
   experimental value of 0.0185; note that argon is
   a monoatomic gas");

```

---

### Scilab code Exa 14.3 non uniform gas theory

```

1 clc;
2 warning("off");
3 printf("\n\n example14.3 - pg727");
4 T=40+273.15; // [K] - temperature
5 P=1; // [atm] - pressure
6 Cp=1005; // [J/kg*K] - heat capacity
7 M=28.966; // [kg/mole] - molecular weight
8 R=8314.3; // [atm*m^3/K*mole] - gas constant

```

```

9 // using the formula Cv=Cp-R/M
10 Cv=Cp-R/M;
11 y=Cp/Cv;
12 mu=19.11*10^-6; // [kg/m*sec] - viscosity of air
13 // using the original Eucken correlation
14 k_original=mu*(Cp+(5/4)*(R/M));
15 printf("\n\n From the original Eucken correlation \n
k=%fW/m*K",k_original);
16 // using the modified Eucken correlation
17 k_modified=mu*(1.32*(Cp/y)+(1.4728*10^4)/M);
18 printf("\n\n From the modified Eucken correlation \n
k=%fW/m*K",k_modified);
19 printf("\n\n As discussed , the value from the
modified Eucken equation is highre than the
experimental value(0.02709) , and the value
predicted by the original Eucken equation is
lower than the experimental value , each being
about 3 percent different , in this case");

```

---

### Scilab code Exa 14.4 non uniform gas theory

```

1 clc;
2 warning(" off");
3 printf("\n\n example14.4 - pg728");
4 // given
5 D=7.66*10^-5; // [m^2/sec] - diffusion coefficient
of the helium nitrogen
6 P=1; // [atm] - pressure
7 // (a) using the Chapman-Enskog
8 T(1)=323; // [K]
9 T(2)=413; // [K]
10 T(3)=600; // [K]
11 T(4)=900; // [K]
12 T(5)=1200; // [K]
13 Ma=4.0026;

```

```

14 sigma_a=2.551*10^-10; // [m]
15 etaabykb=10.22; // [K]
16 Mb=28.016;
17 sigma_b=3.798*10^-10; // [m]
18 etabbykb=71.4; // [K]
19 sigma_ab=(1/2)*(sigma_a+sigma_b);
20 etaabbykb=(etaabykb*etabbykb)^(1/2);
21 Tstar=T/(etaabbykb);
22 siD=[0.7205;0.6929;0.6535;0.6134;0.5865];
23 patm=1;
24 // using the formula Dab=1.8583*10^-27*((T^3)*((1/Ma)+(1/Mb)))^(1/2)/(patm*sigma_ab*siD)
25 Dab=(1.8583*(10^-(27))*((T^3)*((1/Ma)+(1/Mb)))^(1/2))/(patm*(sigma_ab^(2))*siD)
26 printf("\n\n (a)");
27 for i=1:5;
28     printf("\n at T=%fK; Dab=%em^2/sec",T(i),Dab(i));
29 end
30 // (b) using experimental diffusion coefficient and
31 // Chapman-Enskog equation
32 for i=1:4
33     D(i+1)=D(1)*((T(i+1)/T(1))^(3/2))*(siD(1)/(siD(i+1)));
34 end
35 printf("\n\n (b)");
36 for i=1:5;
37     printf("\n at T=%fK; Dab=%em^2/sec",T(i),Dab(i));
38 end
39 // (c)
40 for i=1:4
41     Dab(i+1)=D(1)*(T(i+1)/T(1))^(1.75);
42 end
43 printf("\n\n (c)");
44 for i=1:5;
45     printf("\n at T=%fK; Dab=%em^2/sec",T(i),Dab(i));
46 end

```

---

### Scilab code Exa 14.5 non uniform gas theory

```
1 clc;
2 warning(" off");
3 printf("\n\n example14.5 - pg730");
4 // given
5 T=323; // [K] - temperature
6 P=1; // [atm] - pressure
7 Dab_experimental=7.7*10^-6; // [m^2/sec]
8 DPM_A=1.9; // dipole moment of methyl chloride
9 DPM_B=1.6; // dipole moment of sulphur dioxide
10 Vb_A=5.06*10^-2; // liquid molar volume of methyl
chloride
11 Vb_B=4.38*10^-2
12 Tb_A=249; // normal boiling point of methyl
chloride
13 Tb_B=263; // normal boiling point of sulphur
dioxide
14 del_A=((1.94)*(DPM_A)^2)/(Vb_A*Tb_A);
15 del_B=((1.94)*(DPM_B)^2)/(Vb_B*Tb_B);
16 del_AB=(del_A*del_B)^(1/2);
17 sigma_A=(1.166*10^-9)*(((Vb_A)/(1+1.3*(del_A)^2))
^(1/3));
18 sigma_B=(1.166*10^-9)*(((Vb_B)/(1+1.3*(del_B)^2))
^(1/3));
19 etaabykb=(1.18)*(1+1.3*(del_A^2))*(Tb_A);
20 etabbykb=(1.18)*(1+1.3*(del_B^2))*(Tb_B);
21 sigma_AB=(1/2)*(sigma_A+sigma_B);
22 etaabbykb=(etaabykb*etabbykb)^(1/2);
23 Tstar=T/(etaabbykb);
24 sigmaDnonpolar=1.602;
25 sigmaDpolar=sigmaDnonpolar+(0.19*(del_AB^2))/Tstar;
26 patm=1;
27 Ma=50.488; // [kg/mole] - molecular weight of methyl
```

```

chloride
28 Mb=64.063; // [kg/mole] - molecular weight of
    sulphur dioxide
29 D_AB=(1.8583*(10^-(27))*(((T^3)*((1/Ma)+(1/Mb)))^
    ^((1/2)))/(patm*(sigma_AB^(2))*sigmaDpolar);
30 printf("\n\n Dab=%em^2/sec",D_AB);
31 printf("\n\n The Chapman Enskog prediction is about
    8 percent higher");

```

---

### Scilab code Exa 14.6 empirical correlations for gases

```

1 clc;
2 warning("off");
3 printf("\n\n example14.6 - pg732");
4 // given
5 T=423.2; // [K] - temperature
6 P=5; // [atm] - pressure
7 Ma=4.0026; // [kg/mole] - molecular weight of helium
8 Mb=60.09121; // [kg/mole] - molecular weight of
    propanol
9 Dab_experimental=1.352*10^-5; // [m^2/sec] -
    experimental value of diffusion coefficient of
    helium-propanol system
10 // the diffusion volumes for carbon , hydrogen and
    oxygen are-
11 Vc=16.5;
12 Vh=1.98;
13 Vo=5.48;
14 V_A=3*Vc+8*Vh+Vo;
15 V_B=2.88;
16 patm=5;
17 // using the FSG correlation
18 Dab=(10^-7)*(((T^1.75)*((1/Ma)+(1/Mb))^((1/2)))/(patm
    *((V_A)^(1/3)+(V_B)^(1/3))^2));
19 printf("\n\n Dab=%em^2/sec",Dab);

```

```
20 printf("\n\n The FSG correlation agrees to about 2  
percent with the experimental value");
```

---

### Scilab code Exa 14.7 viscosity

```
1 clc;  
2 warning("off");  
3 printf("\n\n example14.7 - pg736");  
4 // given  
5 beta0=-6.301289;  
6 beta1=1853.374;  
7 clf;  
8 xtitle("Temperature variation of the viscosity of  
water","(1/T)*10^3,K^-1","viscosity,cP");  
9 x=[2.2,0.2,3.8]';  
10 y=[(beta0+beta1*x)];  
11 plot2d(x,y);  
12 // at T=420;  
13 T=420; // [K]  
14 x=1/T;  
15 y=beta0+beta1*x;  
16 mu=exp(y);  
17 printf("\n\n mu=%fcP",mu);  
18 printf("\n\n The error is seen to be 18 percent.AT  
midrange 320(K), the error is approximately 4  
percent");
```

---

### Scilab code Exa 14.8 thermal conductivity

```
1 clc;  
2 warning("off");  
3 printf("\n\n example14.8 - pg737");  
4 // given
```

```

5 M=153.82; // [kg/mole] – molecular weight of ccl4
6 T1=349.90; // [K] – temperature1
7 T2=293.15; // [K] – temperature 2
8 cp1=0.9205; // [KJ/kg*K] – heat capacity at
    temperature T1
9 cp2=0.8368; // [KJ/kg*K] – heat capacity at
    temperature T2
10 p1=1480; // [kg/m^3] – density at temperature T1
11 p2=1590; // [kg/m^3] – density at temperature T2
12 Tb=349.90; // [K] – normal boiling point
13 pb=1480; // [kg/m^3] – density at normal boiling
    point
14 cpb=0.9205; // [KJ/kg*K] – heat capacity at normal
    boiling point
15 k1=(1.105/(M^(1/2)))*(cp1/cpb)*((p1/pb)^(4/3))*(Tb/
    T1);
16 printf("\n\n The estimated thermal conductivity at
    normal boiling point is \n k=%f W*m^-1*K^-1",k1);
17 k2=(1.105/(M^(1/2)))*(cp2/cpb)*((p2/pb)^(4/3))*(Tb/
    T2);
18 printf("\n\n The estimated thermal conductivity at
    temperature %f K is \n k=%f W*m^-1*K^-1",T2,k2);
19 printf("\n\n The estimated value is 3.4 percent
    higher than the experimental value of 0.1029 W*m
    ^-1*K^-1");

```

---

### Scilab code Exa 14.9 diffusion coefficient

```

1 clc;
2 warning("off");
3 printf("\n\n example14.9 – pg743");
4 // given
5 T=288; // [K] – temperature
6 M1=60.09; // [kg/mole] – molecular weight of
    proponal

```

```

7 M2=18.015; // [kg/mole] - molecular weight of water
8 mu1=2.6*10^-3; // [kg/m*sec] - viscosity of propanol
9 mu2=1.14*10^-3; // [kg/m*sec] - viscosity of water
10 Vc=14.8*10^-3; // [m^3/kmol] - molar volume of
    carbon
11 Vh=3.7*10^-3; // [m^3/kmol] - molar volume of
    hydrogen
12 Vo=7.4*10^-3; // [m^3/kmol] - molar volume of
    oxygen
13 Vp=3*Vc+8*Vh+Vo; // molar volume of propanol
14 phi=2.26; // association factor for diffusion of
    propanol through water
15 Dab=(1.17*10^-16*(T)*(phi*M2)^(1/2))/(mu2*(Vp^0.6));
16 printf("\n\n The diffusion coefficient of propanol
    through water is \n Dab=%e m^2/sec",Dab);
17 phi=1.5; // association factor for diffusion of
    water through propanol
18 Vw=2*Vh+Vo; // [molar volume of water
19 Dab=(1.17*10^-16*(T)*(phi*M1)^(1/2))/(mu1*(Vw^0.6));
20 printf("\n\n The diffusion coefficient of water
    through propanol is \n Dab=%e m^2/sec",Dab);

```

---

# Chapter 15

## non newtonial phenomena

**Scilab code Exa 15.1** rheological characteristics of material time independent behaviour

```
1 clc;
2 warning(" off");
3 printf("\\n\\n example15.1 - pg760");
4 // given
5 r=[10 20 50 100 200 400 600 1000 2000]
6 tau=[2.2 3.1 4.4 5.8 7.4 9.8 11.1 13.9 17.0]
7 tau=tau*(10^-4);
8 clf;
9 xtitle(" basic shear diagram for the fluid ", "shear
    rate", "shear stress");
10 plot2d("11", r, tau);
11 // the data falls nearly on a straight line
12 // from the graph the slope and the intercept are
13 slope=0.3841;
14 intercept=9.17046;
15 // from the relation tau=K*(-r)^n;
16 K=exp(intercept);
17 n=slope
18 disp(K, "K=", n, "n=");
19 printf("\\n\\n The fluid is pseudo plastic , since the
```

```
slope is less than 1 ");
```

---

### Scilab code Exa 15.2 capillary viscometer

```
1 clc;
2 warning(" off");
3 printf("\n\n example15.2 - pg774");
4 // given
5 a=[651 1361 2086 5089 7575 11140 19270 25030]
6 tau=[3.71 7.49 11.41 24.08 -35.21 46.25 77.50 96.68]
7 clf;
8 xtitle("capillary shear diagram for polyisobutylene
         L-80 in cyclohexane","pseudoshear rate","wall
         shear stress");
9 plot2d("11",a,tau);
10 // from the graph
11 beta0=-4.3790154;
12 beta1=0.8851;
13 K'=exp(beta0);
14 n'=beta1;
15 printf("\n\n The final rheological model is \n tauw
         = %f*(8*Uz,avg/do)^%f",K',n');
```

---

### Scilab code Exa 15.3 capillary viscometer

```
1 clc;
2 warning(" off");
3 printf("\n\n example15.3 - pg774");
4 // given
5 // from example 15.2
6 n'=0.8851;
7 K'=0.01254;
8 n=n';
```

```

9 K=K' /((3*n+1)/(4*n));
10 disp(n,"n=");
11 printf("\n K = %f N/m^2",K);

```

---

### Scilab code Exa 15.4 capillary viscometer

```

1 clc;
2 warning("off");
3 printf("\n\n example15.4 - pg775");
4 // given
5 a=[10 20 50 100 200 400 600 1000 2000];
6 tau=[2.24 3.10 4.35 5.77 7.50 9.13 11.0 13.52 16.40]
7 tau=tau*10^-4;
8 clf;
9 xtitle("capillary shear diagram for a commercial
         polyethylene melt at 190 degC","pseudoshear rate"
         ,"wall shear stress");
10 plot2d("11",a,tau);
11 // such a plot suggests a second order polynomila
     of the type y=betao+beta1*x+beta2*x^2;
12 // where y=ln(tauw) and x=ln(8*Uz, avg/do)=ln(a);
13 // from the graph
14 betao=8.96694;
15 beta1=0.48452520;
16 beta2=0.010923041;
17 n=beta1+2*beta2*a;
18 phiw=((3*n+1)/(4*n))*(a);
19 mu=tau/phiw;
20 printf("\n\n 8*Uz, avg/do      n      (3*n+1)/(4*n)
          phiw      mu");
21 for i=1:9
22     printf("\n %f      %f      %f      %f      %f"
           ,a(i),n(i),(3*n(i)+1)/(4*n(i)),phiw(i),mu);
23 end

```

---