

# Integrating a Heat Balance Equation

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## 1 Introduction

One method used for heating a steady stream of ethylene glycol fluid is a shell and tube heat exchanger (Figure 1). The fluid is surrounded by a shell containing a saturated vapor which is continuously condensed so that the temperature can be maintained at  $T_s$ . The fluid has an inlet temperature of  $T_1$  and an outlet temperature of  $T_2$ .

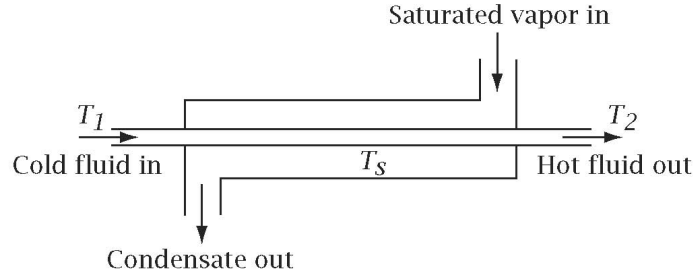


Figure 1: Heat exchanger

The length for the heat exchanger is found from a heat balance differential equation (Finney 2006)

$$\frac{dL}{dT} = \frac{mc_p}{\pi Dh(T_s - T)^2} \quad (1)$$

where

$$\begin{aligned} m &= \text{Mass flow rate (lb/hr)} \\ D &= \text{Tube diameter (ft)} \\ \omega &= \text{Local heat transfer coefficient} \\ c_p &= \text{Specific heat of ethylene glycol} \end{aligned}$$

(1) is an integrable ODE so integrating with respect to T gives L in terms of T

$$L = \frac{m}{\pi D} \int_{T_1}^{T_2} \frac{c_p dT}{h(T_s - T)^2} \quad (2)$$

The local heat transfer coefficient is found from

$$h = \frac{0.023k}{D} \left( \frac{4m}{\pi D \mu} \right)^{0.8} \left( \frac{\mu c_p}{k} \right)^{0.4} \quad (3)$$

where

$$\begin{aligned} k &= \text{Thermal conductivity of ethylene glycol (BTU/hr}\cdot^\circ\text{F)} \\ \mu &= \text{Viscosity (lb/ft}\cdot\text{hr)} \end{aligned}$$

Viscosity can be determined from the relationship

$$\ln \mu = .00002T^2 - .0225T + 5.4874 \quad (4)$$

Specific heat is given by the relationship

$$c_p = .053 + .00065T \quad (5)$$

The values of  $\mu$  and  $c_p$  were determined by a polynomial fit of experimental data (Finney 2006).

## 2 Methodology

To carry out the integration in (2) the function must be in term of  $T$ . Substituting the values of the parameters into (2) results in a graph that has a parabolic shape (figure 2).

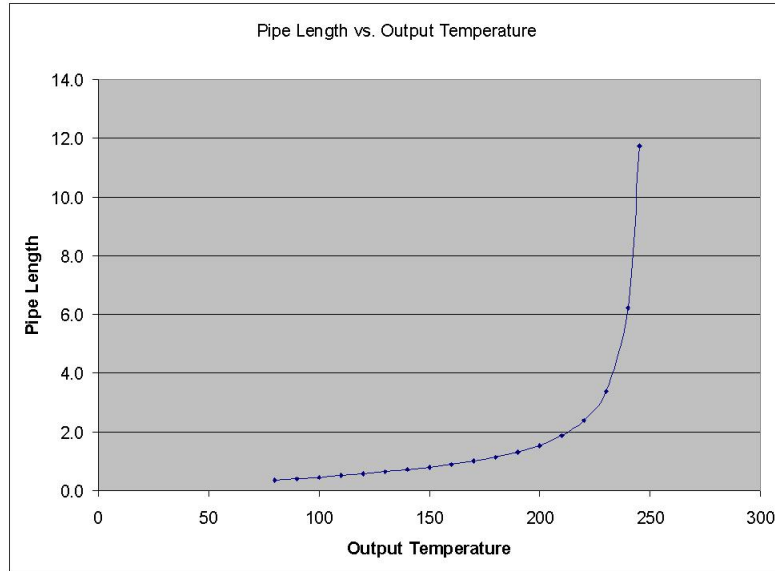


Figure 2: The area under the graph represents pipe length(total area not shown)

(2) has no easily attainable analytical solution so a numerical method must be used to evaluate the integral. In this case the method of Romberg integration based on the trapezoid rule is a sufficiently robust method. A Romberg integration routine combines an approximate integration technique (in this case the trapezoid rule) with Richardson's improvement formula. In each iteration of a Romberg integration routine the step size is scaled by some constant amount and then the new extrapolations are computed.

The trapezoid rule for computing the approximate integral between  $a$  and  $b$  has the general form

$$\int_a^b f(x) = h \left[ \frac{f(a)}{2} + \sum_{i=2}^{np} f[a + (i-1)h] + \frac{f(b)}{2} \right] \quad (6)$$

where  $h$  is the step size  $([b - a]/np)$  and  $np$  is the number of panels to be used. When  $np = 1$  (i.e. the first iteration) the trapezoid rule is very simple to program (figure 2)(appendix A).

Richardson's improvement formula uses a weighted average of approximate solutions (obtained

```

nt=1
h=(b-a)/nt
endpts=f(b)/(2d0)+f(a)/(2d0)
area=(h)*(endpts)

```

Figure 3: First iteration of the trapezoid rule in a Romberg integration routine

using the trapezoid rule) with the same error order to extrapolate a solution with a higher error order. The second approximation must have a different step size than the first approximation. Richardson's improvement formula has the general form

$$I_2(h_2) = \frac{r^k I_1(h_2) - I_1(h_1)}{r^k - 1} \quad (7)$$

where  $h_1 = r h_2$  and  $r > 1$ . Program `heatex` combines the trapezoid rule and Richardson's improvement formula to carry out the integration in (2)(figure 2).

```

j=0
do
  j=j+1
  h=h/r
  nt=2d0*nt
  do i=1,nt/2                !trapezoid rule
    newp=newp+f(a+(i-1)*(2d0*h)+h)
  end do
  IG(j+1,2)=(h)*(endpts+newp)
  k=1
  do i=1,j
    k=2*k                    !richardson's improvement
    IG(j+1,i+2)=((r**(k)*IG(j+1,i+1))-IG(j,i+1))/(r**(k)-1)
  end do
end do

```

Figure 4: Romberg integration routine

The stopping criteria for program `heatex` are a solution found test and a maximum iterations test (figure 5).

```

if(j>maxit-2)then
  write(*,*)"Reached max iteration"
  numit=j
  return
else if(tol<eps)then
  ans=IG(j+1,j+2)
  numit=j
  exit
  return
end if

```

Figure 5: Stopping Criteria

### 3 Application

The pipe length in the heat exchanger is determined from various parameters that are known in the problem (Table 1).

Table 1: Parameters associated with determining pipe length

Parameter	Variable	Value
Tube diameter	$D$	.086 ft
Vapor Temperature	$T_s$	250 °F
Mass flow rate	$m$	45000 lb/hr
Solution tolerance	$\varepsilon$	$10^{-10}$
Specific heat	$c_p$	J/lb·F
Viscosity	$\mu$	lb/ft·hr

To test the sensitivity of the system, different parameters can be varied (Table 2).

Table 2: Variation of Parameters

Run #	Variable	Initial value	New value	Variation
1	$D$	.086 ft	.0774	-10%
2		.086 ft	.0946	10%
3	$T_s$	250 °F	225	-10%
4		250 °F	275	10%
5	$m$	45000 lb/hr	40500	-10%
6		45000 lb/hr	49500	10%
7	$\varepsilon$	$10^{-10}$	$1 \cdot 10^{-5}$	-50%
8		$10^{-10}$	$1 \cdot 10^{-15}$	50%
9	$c_p$	no scale	.9 scale	-10%
10		no scale	1.1 scale	10%
11	$\mu$	no scale	.9 scale	-10%
12		no scale	1.1 scale	10%

## 4 Results

With  $T_2 = 90^\circ\text{F}$  program `heatex` gives the pipe length  $L=0.3967153601$  (figure 6).

```

The best estimate is:   0.3967153601
    # of iterations:    5

The Romberg extrapolation table is (first col. is stepsize):
      h           I 1           I 2           I 4           I 5
90.0000000000  0.4157619924  0.0000000000...0.0000000000  0.0000000000
45.0000000000  0.4016544622  0.3969519521...0.0000000000  0.0000000000
22.5000000000  0.3979631260  0.3967326806...0.0000000000  0.0000000000
11.2500000000  0.3970281530  0.3967164953...0.3967154059  0.0000000000
 5.6250000000  0.3967936117  0.3967154313...0.3967153601  0.3967153601

```

Figure 6: Abridged Romberg integration results for  $T_2 = 90^\circ$ 

With  $T_2 = 180^\circ\text{F}$  program `heatex` gives the pipe length  $L=1.1347906901$  (figure 7).

The best estimate is: 1.1347906901				
# of iterations: 6				
The Romberg extrapolation table is (first col. is stepsize):				
h	I 1	I 2	I 5	I 6
180.000000000	1.7446267397	0.0000000000...	0.0000000000	0.0000000000
90.000000000	1.3316105678	1.1939385105...	0.0000000000	0.0000000000
45.000000000	1.1905851943	1.1435767365...	0.0000000000	0.0000000000
22.500000000	1.1493974712	1.1356682301...	0.0000000000	0.0000000000
11.250000000	1.1384920326	1.1348568864...	1.1348014657	0.0000000000
5.625000000	1.1357191620	1.1347948718...	1.1347906901	1.1347906901

Figure 7: Abridged Romberg integration results for  $T_2 = 180^\circ$ 

The sensitivity analysis reveals how the pipe length and number of iterations change when the parameters are individually varied. It can be seen from the table that the pipe length is most sensitive to a change in the shell temperature. A change in the parameter epsilon results in an almost negligible change in the pile length but it has an effect on the number of iterations to find the solution. The data applies when  $T_2 = 180^\circ\text{F}$ . The data for  $T_2 = 90^\circ\text{F}$  is not included because it shows similar trends.

Table 3: Parameters associated with determining pipe length

Run #	Variable	New value	Percent varied	Pipe Length	Variation	# iterations
1	$D$	0.0774	-10%	1.0430611141	8.08%	6
2		0.0946	10%	1.2247005097	7.92%	6
3	$T_s$	225	-10%	1.8479438758	68.8%	6
4		275	10%	0.7907736591	30.32%	6
5	$m$	40500	-10%	1.1111284458	2.09%	6
6		49500	10%	1.1566295968	1.92%	6
7	$\epsilon$	$1 \cdot 10^{-5}$	-50%	1.1348014657	$\sim 0\%$	5
8		$\cdot 10^{-15}$	50%	1.1347904658	$\sim 0\%$	7
9	$c_p$	.9 scale	-10%	1.0652738588	6.13%	6
10		1.1 scale	10%	1.2015763597	5.89%	6
11	$\mu$	.9 scale	-10%	1.0879595981	4.13%	6
12		1.1 scale	10%	1.1788887985	3.89%	6

As the required output temperature ( $T_2$ ) approaches  $250^\circ\text{F}$  the pipe length approaches infinity (figure 8). Further optimization could be done to find the best pipe length for optimal fluid heating.



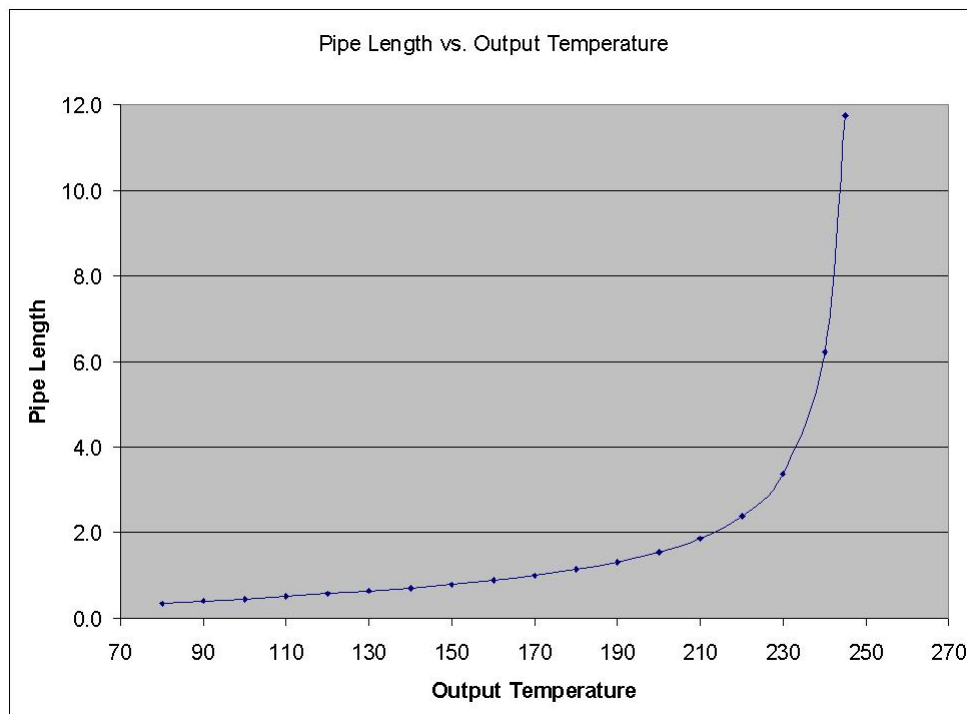


Figure 8: ( $T_2$ ) approaches 250°F as L approaches infinity

## 5 Conclusion

The following can be concluded from the analysis:

- The necessary pipe length when  $T_2 = 90^\circ\text{F}$  is  $L=0.3967153601$
- The necessary pipe length when  $T_2 = 180^\circ\text{F}$  is  $L=1.1347906901$
- Pipe length is most sensitive to a change in the shell temperature ( $T_s$ )
- Required output temperature ( $T_2$ ) approaches 250°F as the pipe length (L) approaches infinity.

## 6 References

Finney, Brad. Lab 1 handout, Humboldt State University, Spring 2006.

## Appendix A

### Source Code and Program Output

\*Please note that the program output was slightly abridged in order to fit them on one page.

```
cwb12@ere-server:~/engr326/lab1> cat lab1.f90
module con
  double precision,parameter::e=2.718281828,pi=3.1415926,m=45000,D=.086,k=.153,
    Ts=250
end module con

program heatex
  use con
  implicit none
  integer::numtrap,maxit,numit,s,j
  double precision::T1,T2,ans,eps
  character(len=30)::func
  double precision,allocatable,dimension(:,:)::I
  interface
    subroutine romtrap(nt,a,b,ans,maxit,eps,numit,IG,f)
      implicit none
      integer,intent(in)::nt,maxit
      integer,intent(out)::numit
      double precision,intent(in)::b,a,eps
      double precision,intent(out)::ans
      double precision,dimension(:,:),intent(out)::IG
      interface
        function f(x)
          double precision::x
          double precision::f
        end function f
      end interface
    end subroutine romtrap
  end interface
  function temp(x)
    double precision::x
    double precision::temp
  end function temp
end interface

!this program will integrate a pre set flow equation using an integration
!subroutine. The subroutine uses the trapezoid rule given upper and lower
!limits, a number of trapezoids and a pipe radius.
!variable list:
!T1=lower limit of integration
!T2=upper limit of integration
!numtrap=number of trapezoids
```

```

write(*,*)"This program will integrate an equation ",&
    "using a trapezoid rule integration subroutine(answer in feet).\"
write(*,*)"Enter the lower limit of integration.\"
read(*,*)T1
write(*,*)"Enter the upper limit of integration.\"
read(*,*)T2
numtrap=1
!write(*,*)"Enter the number of trapezoids to use.\"
!read(*,*)numtrap
maxit=20
allocate(I(maxit,maxit))
eps=.0000000001
call romtrap(numtrap,T1,T2,ans,maxit,eps,numit,I,temp)
write(*,\"(/,a,x,f14.10)\")\"The best estimate is:\",ans
write(*,\"(5x,a,i4,/)\")\"# of iterations:\",numit+1
write(*,\"(a)\")\"The Romberg extrapolation table is (first col. is stepsize):\"
write(*,\"(7x,a,13x)\",advance=\"no\")\"h\"
do j=1,numit+1
    write(*,\"(a,i2,11x)\",advance=\"no\")\"I\",j
end do
write(*,\"(a)\",advance=\"yes\")\" \"
do s=1,numit+1
    write(*,\"(100f14.10)\")(I(s,j),j=1,numit+2)
end do
stop
end program heatex

subroutine romtrap(nt,a,b,ans,maxit,eps,numit,IG,f)
    implicit none
    integer,intent(inout)::nt,maxit
    integer,intent(out)::numit
    double precision,intent(in)::b,a,eps
    double precision,intent(out)::ans
    double precision,dimension(:,:),intent(out)::IG
    double precision::h,intp,endpts,newp,r,tol        !local variables
    integer::i,j,k

    !variable list:
    !h=          step size
    !a,b=        upper/lower limit of integration
    !interior=   area under the curve without the endpoints
    !f=          function name

interface
    function f(x)
        double precision::x
        double precision::f
    end function f

```

```

end interface

h=(b-a)/nt
!write(*,*)h
IG(1,1)=h
intp=0
do i=2,nt
    intp=intp+f(a+(i-1)*h)
end do
endpts=f(b)/(2d0)+f(a)/(2d0)
IG(1,2)=(h)*(endpts+intp)

r=2d0
j=0
do
    j=j+1
    h=h/r
    nt=2d0*nt
    IG(j+1,1)=h
    do i=1,nt/2
        newp=newp+f(a+(i-1)*(2d0*h)+h)
    end do
    IG(j+1,2)=(h)*(endpts+intp+newp)
    k=1
    do i=1,j
        k=2*k
        IG(j+1,i+2)=((r**(k)*IG(j+1,i+1))-IG(j,i+1))/(r**(k)-1)
    end do
    tol=abs(IG(j+1,j+2)-IG(j+1,j+1))
    if(j>maxit-2)then
        write(*,*)"Reached max iteration"
        numit=j
        return
    else if(tol<eps)then
        ans=IG(j+1,j+2)
        numit=j
        exit
        return
    end if
end do

return
end subroutine romtrap

function temp(T)
    use con
    double precision::temp,T,cp,lnu,u,ht
    cp=.53d0+.00065*T

```

```

    lnu=.00002d0*(T*T)-((.0225d0)*T)+5.4874d0
    u=e**(lnu)
    ht=((.023d0*k)/d)*(((4d0*m)/(pi*D*u))**(.8d0))*(((u*cp)/k)**(.4d0))
    temp=(m/(pi*D))*(cp/(ht*((Ts-T)**(2d0))))
    return
end function temp

cwb12@ere-server:~/engr326/lab1> ifort lab1.f90 -o lab1
cwb12@ere-server:~/engr326/lab1> lab1
This program will integrate an equation
using a trapezoid rule integration subroutine(answer in feet).
Enter the lower limit of integration.
0
Enter the upper limit of integration.
90

The best estimate is:    0.3967153601
    # of iterations:    5

The Romberg extrapolation table is (first col. is stepsize):
      h          I 1          I 2          I 3          I 5
90.0000000000  0.4157619924  0.0000000000  0.0000000000...0.0000000000
45.0000000000  0.4016544622  0.3969519521  0.0000000000...0.0000000000
22.5000000000  0.3979631260  0.3967326806  0.3967180625...0.0000000000
11.2500000000  0.3970281530  0.3967164953  0.3967154163...0.0000000000
 5.6250000000  0.3967936117  0.3967154313  0.3967153603...0.3967153601
cwb12@ere-server:~/engr326/lab1> lab1
This program will integrate an equation
using a trapezoid rule integration subroutine(answer in feet).
Enter the lower limit of integration.
0
Enter the upper limit of integration.
180

The best estimate is:    1.1347906901
    # of iterations:    6

The Romberg extrapolation table is (first col. is stepsize):
      h          I 1          I 2          I 3          I 6
180.0000000000  1.7446267397  0.0000000000  0.0000000000...0.0000000000
90.0000000000  1.3316105678  1.1939385105  0.0000000000...0.0000000000
45.0000000000  1.1905851943  1.1435767365  1.1402192849...0.0000000000
22.5000000000  1.1493974712  1.1356682301  1.1351409964...0.0000000000
11.2500000000  1.1384920326  1.1348568864  1.1348027968...0.0000000000
 5.6250000000  1.1357191620  1.1347948718  1.1347907375...1.1347906901

```

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