CHEMCAD CALCULATOR/_PARSER MODULE

GENERAL INFORMATION

The Calculator/Parser module in CHEMCAD is a powerful utility module, which offers the power and flexibility of the high level C-like language without having to use a compiler. The module allows the user to do the following things:

- Create your own unit operation within a flowsheet.
- Retrieve stream/equipment/component information in a flowsheet.
- Perform mathematical calculation. Most mathematical functions are built into the standard library.
- The "for statement" for loop calculation, the "if else" statement for information control and the "goto" statement are allowed in the module.
- All flash functions, physical property routines and many other useful functions can be called to perform necessary calculations.
- Print the calculated results by the "printf" statement.
- Put the calculated results back to the flowsheet.

Typical applications of the calculator/parser module include:

- Calculate stream compositions or property for use by the controller.
- Calculate size/cost of process equipment.
- Calculate the flowrate of the makeup streams for better convergence.
- Simulate a special unit operation as required.

The usefulness of the module is limited only by the imagination of the user.

COSTING CALCULATIONS

All of the cost calculating routines in CHEMCAD are done using the Parser language described here. This is to enable users to modify the costing routines for their own use. The source code for these calculations is stored in the CHEMCAD program directory (typically C:\CC5) and have the following names:

- $$$aclr.sf  Air cooled heat exchanger costing
- $$$cfug.sf  Centrifuge costing
- $$$comp.sf  Compressor costing
- $$$crsh.sf  Crusher costing
- $$$crys.sf  Crystallizer costing
- $$$cycl.sf  Cyclone costing
- $$$dpip.sf  Double pipe heat exchanger costing
- $$s$evap.sf  Evaporator heat exchanger costing
- $$$expn.sf  Expander costing
DIALOG BOXES FOR PARSER UNITS

If your CALC unitop needs to use a dialog screen, it must be built according to the Screen Builder manual and documentation. Please see the screen builder documentation for more details.

For a Parser unit op, the screen files will be named in the same format as the program file, as “CALC#.MY” and “CALC#.MAP” where # is the unit ID number. For example, if your unit id number is 5, then the files would be named “calc5.my” and “calc5.map”. Please see the screen builder section of this manual for more details on these files.

PARSER LANGUAGE SYNTAX AND STRUCTURE

The Calculator/Parser module uses a C-like language to perform all calculations. It is recommended that the user be familiar with some of the basic C language syntax before writing the code for this module. Several examples at the end of this chapter should be studied before you proceed.

Data types:

- int integer
- float floating point real numbers
- double double precision real numbers

Examples:

- int i, j, k;
- double x, y, z;
- float a, b, c;

Both one dimensional array and two dimensional arrays are supported.

Examples:

- int ix[100];
- double x[50];
- float uspec[300];
- float matrix[10][10];
Intrinsic functions:

+ add
- subtract
* multiply
/ divide
exp exponential
log natural logarithm (Ln)
log10 common logarithm
pow(x,y) x to the power of y
fabs absolute value of a real number
abs absolute value of an integer
sqrt square root of a real number
sin Sine(radians)
cos Cosine(radians)
tan Tangent(radians)
asin Arc Sine(radians)
acos Arc cosine(radians)
atan Arc tangent(radians)
== equal
> greater than
>= greater or equal
< less than
<= less or equal
&& and
|| or

Loop calculations:

The "for statement" can be used for loop calculation.

Example:

```c
int i, n;
n = 30;
for( i = 0; i < n; i++)
{
    /* Program code */
    ....
}
```

Up to two nested loops are allowed.

Example:

```c
int i, j;
for( i = 0; i < 10; i++)
{
    for( j = 0; j < 20; j++)
```
Information control:

- if (expression ) expression;
  Example:
  a = 0.;
  if( x > 20. ) a = 1.;

- if( expression )
  { expressions;
  }
  Example:
  a = 0.;
  b = 0.;
  if( x > 20. )
  { a = 1.;
    b = 2.;
  }

- if( expression)
  { expressions;
  }
  else if( expression )
  { expressions;
  }
  else if( expression )
  { expressions;
  }
  else
  { expressions;
  }
  Example:
  a = 0.;
  b = 0.;
  if( x > 10. )
  {
a = 1.;
b = 1.;
}
else if( x > 20. )
{
a = 2.;
b = 2.;
}
else if( x > 30. )
{
a = 3.;
b = 3.;
}
else
{
a = 4.;
b = 4.;
}

Note: Nested "if else" up to two levels are allowed.

Example:
if( a > (b * 23.))
{
  if( c > 25. )
  {
      .....  
  }
  else
  {
      ....
  }
}

- goto LABEL;

Example:
x = 23 * y;
if( x > 100. ) goto THERE;
....
THERE:
....

Note: The goto statement can only be used to jump to the statement below to goto line. It cannot be used to jump back to the previous statement.
printf statement

printf statement can be used similar to the C language.
Up to 5 arguments can be included in one printf statement.

For debugging purpose, it is recommended that the user use the "Generate Run History" checkbox on the "Convergence" screen of the "Run" menu so that the printf and any error messages can be seen on the screen when you run the calculator module.

Example:

printf("Calculate area = %g, heat duty = %g\n",area,duty);
printf("Stream %d, Temperature = %13.2f\n",str_ID,temp);

Return Values of the Arguments:

In the standard C syntax, the return value of an argument in a function call should be passed by the address. This is omitted in the calculator module as illustrated by the following example.

Standard C:

ProgramX()
{
    float input_data;
    float output_data;
    /* Notice the address sign (&) for the output data */
    functionX(input_data,&output_data);
}

CHEMCAD Calculator:

CHEMCAD_Link Program_Name
{
    float input_data;
    float output_data;
    /* Notice the address sign (&) is omitted here */
    functionX(input_data,output_data);
}

PROGRAM STRUCTURE

A program file has to be created in the job subdirectory with the file name CALC#.TXT where # is the equipment ID number of the calculator. For example, if the calculator number is 5, you need to write a program CALC5.TXT and put this file in your job subdirectory. The program file has to have the following structure:

CHEMCAD_Link Program_Name
{
    /* Section 1 : Variable declarations */
    .... ;
Creating a dialog screen for a Parser (Calculator) unit op

If your unit has any settings, such as output pressure for a pump or # of stages for a distillation column, you will need to generate a dialog box. The construction of dialogs is covered in the Screen Builder section of this manual.

For a Parser unit op, the screen files will be named in the same format as the program file, as “CALC#.MY” and “CALC#.MAP” where # is the unit ID number. For example, if your unit id number is 5, then the files would be named “calc5.my” and “calc5.map”. Please see the screen builder section of this manual for more details on these files.
CHEMCAD LIBRARY FUNCTIONS

The following functions may be accessed from a Parser unit. They fall into the general categories of:

- Flowsheet Interface: Functions to move data from the flowsheet into the calculator and back
- Thermodynamics/Engineering Calcs: Functions used to calculate properties of mixtures

FLOWSHEET INTERFACE FUNCTIONS:

These functions are used to pass information into the calculator function and back.

- CC_Get_Equip_Parameters - returns the unitop array for a specified unit
- CC_Put_Equip_Parameters - overwrites the unitop array for a specified unit
- CC_Get_Process_Stream - returns a flowsheet stream data
- CC_Put_Process_Stream - overwrites a flowsheet stream with new data
- CC_Get_Input_Stream - returns stream data for a stream connected to the Calculator unit’s inlet
- CC_Put_Inlet_Stream - overwrites the stream data for a stream connected to the Calculator unit’s inlet
- CC_Get_Outlet_Stream - returns stream data for a stream connected as an outlet to the Calculator unit.
- CC_Put_Outlet_Stream - overwrites the stream data for a stream connected to the Calculator unit’s outlet
- CC_No_Of_Components - returns the number of components
- CC_Int_Array_Function - returns an array of flowsheet information
- CC_Get_Stream_Property - returns a property of a given stream
- CC_Get_Value - generic interface to pure component, stream and unit data

Function: CC_Get_Equip_Parameters

Get equipment parameters for a specified equipment ID number.

Prototype: int CC_Get_Equip_Parameters(int id, float uspec[])
Description:

A float array of uspec[300] should be declared in the file before this function is called. After
calling CC_Get_Equip_Parameters the array uspec[300] should store all parameters in internal
units for this equipment.

Example:

/* Get heat exchanger area and LMTD for equipment 12 */
CHEMCAD_Link Parser
{
  float uspec[300];
  float area_ft2, LMTD;
  /* Get parameters for equipment 12 which is an exchanger */
  CC_Get_Equip_Parameters(12,uspec);
  /* Area (var. no = 29) in ft2 */
  area_ft2 = uspec[29];
  /* LMTD (var. no = 22) in deg R */
  LMTD = uspec[22];
  ...... 
}

******************************************************************************

Function: CC_Put_Equip_Parameters

Put equipment parameters for a specified equipment ID number into the flowsheet

Prototype: int CC_Put_Equip_Parameters(int id, float uspec[])

Parameters:

input id equipment ID
  if id = 0, the current (active) equipment parameters in the memory will be updated
uspec[] an array stores all parameters for the specified equipment

Return Value:

=0 normal return
= 1 error

Description:

A float array of uspec[300] should be declared in the file before this function is called. The array
uspec[300] should store all parameters in internal units.

Example:
/* Calculate basic heat exchanger cost for equipment 12 */

CHEMCAD_Link Parser
{
    float uspec[300];
    float area_ft2, log_area, bcost;

    /* Get parameters for equipment 12 which is an exchanger */
    CC_Get_Equip_Parameters(12, uspec);

    /* Area (var. no = 29) in ft2 */
    area_ft2 = uspec[29];

    /* Log area */
    log_area = log(area_ft2);

    /* Calc base cost of exchanger */
    bcost = exp(8.821 - 0.30863 * log_area +
                0.0681 * log_area * log_area);

    /* Put cost(var. no 48) back to htxr */
    uspec[48] = bcost;
    CC_Put_Equip_Parameters(12, uspec);
}

Funcitio: CC_Get_Process_Stream

Get component flowrates, temperature, pressure, vapor fraction and enthalpy of a specified stream.

Prototype: int CC_Get_Process_Stream(int id, float xmol[], float t, float p, float vf, float h)

Parameters:
    input id stream ID
    output xmol[] An array stores component flowrates
    in lbmol/hr
    t temperature in R
    p pressure in psia
    vf mole vapor fraction
    h enthalpy in btu/hr

Return Value:
    = 0 normal return
    = 1 error

Description:
    A valid stream ID > 0 must be specified. Xmol[] must have a dimension equal to or greater than the
    number of components in the system.
Example:

`/*  Get stream 20 information */

CHEMCAD_Link Parser
{
  float xmol[50];
  float t, p, vf, h;
  /*  Get stream 1 information */
  CC_Get_Process_Stream(20,xmol, t, p, vf, h);
  ....
}

----------------------------------------------------------------------

Function:   CC_Put_Process_Stream

Put component flowrates, temperature, pressure, vapor fraction and enthalpy of a specified stream back to the flowsheet.

Prototype:  int   CC_Put_Process_Stream(int id, float xmol[], float t, float p, float vf, float h)

Parameters:

  input id stream ID
  xmol[] An array stores component flowrates
         in lbmol/hr
  t   temperature in R
  p   pressure in psia
  vf  mole vapor fraction
  h   enthalpy in btu/hr

Return Value:

  = 0  normal return
  = 1  error

Description:

A valid stream ID > 0 must be specified. Xmol[] must have a dimension equal to or greater than the number of components in the system.

Example:

`/*  Double the flowrate and enthalpy of stream 20 */

CHEMCAD_Link Parser
{
  float xmol[50];
  float t, p, vf, h;
  int  nc, i;
  /*  Get stream 1 information */
`
CC_Get_Process_Stream(20, xmol, t, p, vf, h);

/* Get the no of components in the system */
nc = CC_No_Of_Components();

/* Double the component flowrate */
for( i = 0; i < nc; i++)
{
xmol[i] = 2. * xmol[i];
}

/* Double the enthalpy */
h = 2. * h;

/* Put the calculated information back to stream 20 */
CC_Put_Process_Stream(20, xmol, t, p, vf, h);

// Function: CC_Get_Input_Stream
Get component flowrates, temperature, pressure, vapor fraction and enthalpy of a input stream for
the calculator module.

Prototype: int CC_Get_Input_Stream(int ith_input, float xmol[], float t, float p, float vf, float h)

Parameters:
input ith_input The ith input st ream of the calculator base 1.
output xmol[] An array stores component flowrates
in lbmol/hr
t temperature in R
p pressure in psia
vf mole vapor fraction
h enthalpy in btu/hr

Return Value:
= 0 normal return
= 1 error

Description:
Xmol[] must have a dimension equal to or greater than the number of components in the system.

Example:
/* Get the first input stream information */
CHEMCAD_Link Parser
{
    float xmol[50];
float t, p, vf, h;
/* Get stream 1 information */
CC_Get_Input_Stream(1,xmol, t, p, vf, h);
.....
}

Function: CC_Get_Output_Stream

Get component flowrates, temperature, pressure, vapor fraction and enthalpy of an output stream for
the calculator module.

Prototype: int CC_Get_Output_Stream(int ith_output, float xmol[], float t, float p, float vf, float h)

Parameters:
input  ith_output  The ith output stream of the calculator base 1.
output  xmol[]  An array stores component flowrates
           in lbmol/hr
           t  temperature in R
           p  pressure in psia
           vf  mole vapor fraction
           h  enthalpy in btu/hr

Return Value:
= 0  normal return
= 1  error

Description:
Xmol[] must have a dimension equal to or greater than the number of components in the system.

Example:
/* Get the first output stream information */
CHEMCAD_Link Parser
{
  float xmol[50];
  float t, p, vf, h;
  /* Get stream 1 information */
  CC_Get_Output_Stream(1,xmol, t, p, vf, h);
  .....  
}

Function: CC_Put_Input_Stream
Put component flowrates, temperature, pressure, vapor fraction and enthalpy of a input stream back to the flowsheet.

Prototype: int CC_Put_Input_Stream(int ith_input, float xmol[], float t, float p, float vf, float h)

Parameters:
- ith_input: The ith input stream of the calculator base 1.
- xmol[]: An array stores component flowrates in lbmol/hr
  - t: temperature in R
  - p: pressure in psia
  - vf: mole vapor fraction
  - h: enthalpy in btu/hr

Return Value:
- = 0  normal return
- = 1  error

Description:
Xmol[] must have a dimension equal to or greater than the number of components in the system.

Function: CC_Put_Output_Stream

Put component flowrates, temperature, pressure, vapor fraction and enthalpy of a output stream back to the flowsheet.

Prototype: int CC_Put_Output_Stream(int ith_output, float xmol[], float t, float p, float vf, float h)

Parameters:
- ith_output: The ith output stream of the calculator base 1.
- xmol[]: An array stores component flowrates in lbmol/hr
  - t: temperature in R
  - p: pressure in psia
  - vf: mole vapor fraction
  - h: enthalpy in btu/hr

Return Value:
- = 0  normal return
- = 1  error

Description:
Xmol[] must have a dimension equal to or greater than the number of components in the system.

Example:
/* Simulate a 50/50 split divider */
CHEMCAD_Link Parser
{ float xmol[50];
    float t, p, vf, h;
    int nc, i;

    /* Get stream 1 information */
    CC_Get_Input_Stream(1, xmol, t, p, vf, h);

    /* Get the no of components in the system */
    nc = CC_No_Of_Components();

    /* Double the component flowrate */
    for( i = 0; i < nc; i++)
    {
        xmol[i] = 0.5 * xmol[i];
    }

    /* Double the enthalpy */
    h = 0.5 * h;

    /* Put the calculated information to output stream 1 */
    CC_Put_Output_Stream(1, xmol, t, p, vf, h);

    /* Put the calculated information to output stream 2 */
    CC_Put_Output_Stream(2, xmol, t, p, vf, h);
}

----------------------------------------------------------------------------------------------------

Function: CC_No_Of_Components
Get the number of components in the system.

Prototype: int CC_No_Of_Components()

Parameters:
    input None

Return Value:
    No of components in the system

Description:
    This function returns the number of components in the system.

Example:
CHEMCAD_Link Parser
{
    int nc;

    /* Get the no of components in the system */
    nc = CC_No_Of_Components();
}
Function:  CC_Get_Stream_Property

Get the property of a stream.

Prototype:  float  Get_Stream_Property(float xmol[], double t, double p, double vf, double h, int iprop)

Parameters:

input  xmol[]  An array stores component flowrates in lbmol/hr
     t    temperature in R
     p    pressure in psia
     vf   mole vapor fraction
     h    enthalpy in btu/hr
     iprop property flag

  1  = Temperature
  2  = Pressure
  3  = Mole vap frac
  4  = Enthalpy
  5  = Tot. mole rate
  6  = Tot. mass rate
  7  = Total std liq.
  8  = Total std vap.
  9  = Total act. vol
 10  = Tot. act. dens
 11  = Total Mw
 12  = Gross H value
 13  = Net H value
 14  = Reid vapor P
 15  = UOPK
 16  = VABP
 17  = MeABP
 18  = Flash point
 19  = Pour point
 20  = Total entropy
 21  = Mass vap frac
 22  = PH value
 23  = Total resv2
 24  = Total resv3
 25  = Total resv4
 26  = Vap mole rate
 27  = Vap mass rate
 28  = Vapor enthalpy
 29  = Vapor entropy
30 = Vapor Mw
31 = Vap act. dens
32 = Vap act. vol.
33 = Vap std. liq.
34 = Vap std. vap.
35 = Vapor cp
36 = Vapor Z factor
37 = Vap viscosity
38 = Vapor th. cond
39 = Vapor resv1
40 = Vapor resv2
41 = Liq. mole rate
42 = Liq. mass rate
43 = Liq. enthalpy
44 = Liq. entropy
45 = Liq. Mw
46 = Liq. act. dens
47 = Liq. act. vol.
48 = Liq. std. liq.
49 = Liq. std. vap.
50 = Liq. cp
51 = Liq. Z factor
52 = Liq. viscosity
53 = Liq. th. cond.
54 = Liq. surf tens

= -i, flow rate of ith component
   (lbmol/hr)
= -(i+200) ith comp mass flow rate(lb/hr)
= -(i+400) ith comp std vol rate(ft3/hr)
= -(i+600) ith comp mole frac
= -(i+800) ith comp mass frac
= -(i+1000) ith comp vol. frac

Return Value:

The desired property in internal unit.

temperature = R
pressure = psia
mole flow rate = lbmol/hr
mass flow rate = lb/hr
enthalpy = btu/hr
work = btu/hr
entropy = btu/R/hr
vol flow rate = ft3/hr
density = lb/ft3
viscosity = cp
surface tens = dyne/cm
Description:
This function returns the property of a stream with given component flowrate, temperature, pressure, vapor fraction and enthalpy. The return value is always in internal unit.

Example:
/* Get the liquid viscosity of the first input stream in the calculator module */

CHEMCAD_Link Parser
{
  float xmol[50];
  float t, p, vf, h;
  float liq_vis;
  /* Get stream 1 information */
  CC_Get_Input_Stream(1,xmol, t, p, vf, h);
  /* Get liquid viscosity */
  liq_vis = CC_Get_Stream_Property(xmol,t,p,vf,h,52);

  .......
}

Function: CC_Get_Value
Get a value of the system.

Prototype: double CC_Get_Value(int id1, int id2, int id3, int id4)
Parameters:
input:
  if id1 = 1 Pure comp data
    id2 = 1 Mw
    id2 = 2 Tc
    id2 = 3 Pc
    id2 = 4 accentric factor
    id3 = comp position, base 1
  if id1 = 2 stream property data
    id2 = stream ID
    id3 = property flag, See CC_Get_Stream_Property
  if id1 = 3 Get equipment input/output stream ID
    id2 = equip ID
    id3 > 0 (id3)th input stream ID
    id3 < 0 -(id3)th output stream ID
if id1 = 4   Get cost index
    id2 = equip type, see $index.sf in cc3
    id3 = 0   Base cost index
            = 1   Current cost index

if id1=7   Get dynamic time in minutes

Return Value:
The desired value from the system

Description:
This is a generalized function that lets you get a single value from the process including pure
component properties, stream properties, and unit operation input/output stream numbers.

Example:

CHEMCAD_Link Parser
{
    int nc, i;
    float Mw[50];
    float total_vol;
    /*  Get the no of components in the system */
    nc = CC_No_Of_Components();
    /*  Get the molecular weight of each component */
    for( i = 0; i < nc; i++)
    {
        /*  Note: i+1 below is the component position base 1 */
        Mw[i] = CC_Get_Value(1,1,i+1,0);
    }
    /*  Get the actual total volumetric flowrate of stream 3 */
    total_vol = CC_Get_Value(2,3,9,0);
    ...
}

Function:   CC_Int_Array_Function
General integer array function
Prototype:  int CC_Int_Array_Function(int id1, int id2, int id3, int intArray[])

Parameter:
    id1 = 1 Get all streams in flowsheet
    id2, id3 not used
    return intArray[] contains all stream ID's
    return value is no of element in intArray
id1 = 2 Get equipment IDs in flowsheet  
id2, id3 not used  
return intArray[] contains all equipment ID's  
return value is no of element in intArray  
id1 = 3 Not used  
id1 = 4 Get component IDs  
id2, id3 not used  
return intArray[] contains all component ID's  
return value is no of element in intArray  

Return Value: Return value is no of element in intArray.

Description:

This is a generalized function that lets you get an array of integers from the flowsheet. Depends on the id1 value, the function returns stream numbers, equipment numbers or component ID numbers in intArray[].

THERMODYNAMICS/ENGINEERING FUNCTIONS:

Used to calculate the properties of mixtures

- CC_tpflash-constant temperature and pressure equilibrium
- CC_vpflash-constant vapor fraction and pressure equilibrium
- CC_vtflash-constant vapor fraction and temperature equilibrium
- CC_hspflash-adiabatic/isentropic equilibrium
- CC_keq-returns K values (Y fraction/X fraction) for a mixture
- CC_kxeq-returns K-values using ADDK for a mixture
- CC-?????
- CC_enthalpy-returns enthalpy for a mixture at conditions
- CC_hxstream-returns enthalpy cased on ADDH?
- CC_entropy-returns entropy calculation for a given mixture
- CC_zfactor-returns compressibility (z) factor for a mixture
- CC_ldense-returns liquid density
- CC_vdense-returns vapor density
- CC_vp-returns pure component vapor pressure
- CC_cp-returns heat capacity of a mixture
CC_cv-returns cv of a mixture
CC_lvisco-returns liquid viscosity of a mixture
CC_gvisco-returns vapor viscosity of a mixture
CC_lthc-returns liquid thermal conductivity of a mixture
CC_vthc-returns vapor thermal conductivity of a mixture
CC_surften-returns the surface tension of a mixture

Function: CC_tpflash

Isothermal flash calculation at specified temperature and pressure

Prototype: int CC_tpflash( float feed[], double hin, double tr, double psia,
float vapor[], float xliq[], float totalv,
float totall, float hvap, float hliq, float htotal,
float vout, float xk[], float delq, float f_ions[])

Parameters:

Input: feed[] lbmole/hr
hin btu/hr
tr flash T in degree R
psia flash P in psia

Output: vapor[] vapor out lbmole/hr
xliq[] liquid out lbmole/hr
totalv total vapor flow lbmole/hr
totall total liquid flow lbmol/hr
hvap vapor enthalpy btu/hr
hliq liquid enthalpy btu/hr
htotal total H in output streams btu/hr
vout vapor fraction
xk[] K values
delq heat duty = htotal - hin
f_ions[] ions flow rate lbmol/hr

Return Value int ret_val = 0 Normal return
= 1 Diverge

Description:

This program calculates isothermal flash at given T(R) and P(psia). The input enthalpy (hin) should be non-zero if enthalpy calculation is needed. If the electrolyte package is chosen, the routine also returns the flow rate of ions in the liquid phase.
Function: CC_vpflash
Flash calculation at specified mole vapor fraction and pressure

Prototype: int CC_vpflash(float feed[], double v, double p, double hin, double testi,
int irelod, float vapor[], float xliq[], float totalv,
float totall, float tout, float hvap, float hliq,
float htotal, float xk[], float delq, float f_ions[])

Parameters:
Input : feed[] lbmole/hr
v vapor fraction
p pressure psia
hin feed enthalpy Btu/hr
if = 0, no heat duty will be calculated
testi estimated flash temperature (R)
irelod = 0 Start from scratch.
= 1 Reload K values from xk[],
Output: vapor[] vapor product lbmole/hr
xliq[] liquid product lbmole/hr
totalv total vapor rate lbmole/hr
totall total liquid rate lbmole/hr
tout flash output temperature (R)
hvap vapor enthalpy Btu/hr
hliq liquid enthalpy Btu/hr
htotal total output enthalpy Btu/hr
xk[] flash K values
delq heat duty Btu/hr
f_ions[] ions flow rate lbmol/hr

Return Value int ret_val = 0 Normal return
= 1 Diverge

Description:
This program performs flash calculation at given vapor fraction and pressure.
The input enthalpy (hin) should be non-zero if enthalpy calculation is needed.
If input vapor fraction = 0, the routine calculates the bubble point temperature.
If input vapor fraction = 1, the routine calculates the dew point temperature.
Estimated output temperature must be given.
If the electrolyte package is chosen, the routine also returns the flow rate of ions in the liquid phase.

Function: CC_vtflash
Flash calculation at specified mole vapor fraction and temperature

Prototype: int CC_vtflash(float feed[], double vfrac, double t, double hin,
double pesti, int irelod, float vapor[], float xliq[],
float totalv, float totall, float pout, float hvap,
float hliq, float htotal, float xk[], float delq,
float f_ions[])

Parameters:

Input: feed[] lbmole/hr
   vfrac vapor fraction
   t flash temperature (R)
   hin feed enthalpy Btu/hr
   pesti estimated flash pressure (psia)
   irelod = 0, Start from scratch.
       = 1, Reload K values.

Output: vapor[] vapor product lbmole/hr
        xliq[] liquid product lbmole/hr
        totalv total vapor rate lbmole/hr
        totall total liquid rate lbmole/hr
        pout flash output pressure (psia)
        hvap vapor enthalpy Btu/hr
        hliq liquid enthalpy Btu/hr
        htotal total output enthalpy Btu/hr
        xk[] flash K values
        delq heat duty Btu/hr
        f_ions[] ions flow rate lbmol/hr

Return Value int ret_val = 0 Normal return
       = 1 Diverge

Description:

This program performs flash calculation at given vapor fraction and temperature.
The input enthalpy (hin) should be non-zero if enthalpy calculation is needed.
If input vapor fraction = 0, the routine calculates the bubble point pressure.
If input vapor fraction = 1, the routine calculates the dew point pressure.
Estimated output pressure must be given.
If the electrolyte package is chosen, the routine also returns the flow rate of ions in the liquid phase.

Function: CC_hspflash
Adiabatic/Isentropic flash calculation at specified pressure

Prototype: int CC_hspflash(float feed[], double p, double hsin, int mode,
double testi, float vapor[], float xliq[], float totalv,
float totall, float tout, float hvap, float hliq,
float htotal, float xk[], float v, float f_ions[])

Parameters:

Input: feed[] lbmole/hr
p pressure psia
hsin, mode H (mode=0)Btu/hr or S(mode=1)Btu/R/hr
testi estimated flash temperature (R)

Output: vapor[] vapor product lbmole/hr
        xliq[] liquid product lbmole/hr
totalv total vapor rate lbmole/hr
totali total liquid rate lbmole/hr
tout flash output temperature (R)
hvap vapor enthalpy Btu/hr
hliq liquid enthalpy Btu/hr
htotal total output enthalpy Btu/hr
xk[] flash K values
     v output vapor fraction
     f_ions[] ions flow rate lbmol/hr

Return Value int ret_val = 0 Normal return
          = 1 Diverge

Description:

For adiabatic flash (Constant enthalpy) calculation, set mode = 0.
For isentropic flash (Constant entropy) calculation, set mode = 1.

Parameter hsin is feed enthalpy for adiabatic calculation.
Parameter hsin is feed entropy for adiabatic calculation.

Estimated output temperature must be given.

If the electrolyte package is chosen, the routine also returns the flow rate of ions in the liquid phase.

-----------------------------------------------------------------------------------

Function: CC_keq
K value calculation routine

Prototype: void CC_keq(float yv[], float xl[], double t, double p,
                         float xkv[], float f_ions[])

Parameters:

Input: yv[] vapor mole flow rate (lbmol/hr)
       xl[] liquid mole flow rate(lbmol/hr)
       t stream temperature in degree R
       p stream pressure in psia

Output: xkv[] K values
        f_ions[] ions flow rate in liquid phase, lbmol/hr

Return Value: void

Description:
For given vapor composition, liquid composition, temperature and pressure, this routine calculates
the equilibrium K value for each component according to the K model selected by the user. If
electrolyte package is chosen, the routine also returns the flow rate of ions in the liquid phase.

Function: CC_kxeq
K value calculation routine called from user added K value routine (ADDK)

Prototype: void CC_kxeq(float yv[], float xl[], double t, double p, float xkv[])

Parameters:
  Input: yv[] vapor mole flow rate (lbmol/hr)
    xl[] liquid mole flow rate(lbmol/hr)
    t stream temperature in degree R
    p stream pressure in psia

Output: xkv[] K values

Return Value: void

Description:
Calculates K values for any thermo method. For given vapor composition, liquid composition,
temperature and pressure, this routine calculates the equilibrium K value for each component
according to the external variable modek.

The following table shows the value of modek and corresponding model:

<table>
<thead>
<tr>
<th>modek</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Polynomial K</td>
</tr>
<tr>
<td>2</td>
<td>Grayson Streed</td>
</tr>
<tr>
<td>3</td>
<td>SRK</td>
</tr>
<tr>
<td>4</td>
<td>API SRK</td>
</tr>
<tr>
<td>5</td>
<td>UNIFAC</td>
</tr>
<tr>
<td>6</td>
<td>K table</td>
</tr>
<tr>
<td>7</td>
<td>Wilson</td>
</tr>
<tr>
<td>8</td>
<td>Ideal Vapor Pressure</td>
</tr>
<tr>
<td>9</td>
<td>Peng-Robinson</td>
</tr>
<tr>
<td>10</td>
<td>NRTL</td>
</tr>
<tr>
<td>11</td>
<td>ESSO</td>
</tr>
<tr>
<td>12</td>
<td>Amine</td>
</tr>
<tr>
<td>13</td>
<td>Sour Water</td>
</tr>
<tr>
<td>14</td>
<td>UNIQUAC</td>
</tr>
<tr>
<td>15</td>
<td>Margules</td>
</tr>
<tr>
<td>16</td>
<td>Regular Solution</td>
</tr>
<tr>
<td>17</td>
<td>Van Laar</td>
</tr>
<tr>
<td>18</td>
<td>ADDK</td>
</tr>
<tr>
<td>19</td>
<td>Henry's law</td>
</tr>
</tbody>
</table>

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Function: CC_enthalpy
Enthalpy calculation routine.

Prototype: double CC_enthalpy(float xmol[], double t, double p, int iphase, float hx)

Parameters:
- Input: xmol[] component mole flow rate(lbmole/hr)
  - t  temperature in degree R
  - p  pressure in psia
  - iphase 0 = Liquid, 1 = Vapor

Output hx  Btu/hr

Return Value: function return value is enthalpy in double precision.

Description:
For given component flow rate, temperature and pressure, this routine calculates the enthalpy of the stream according to the enthalpy model selected by the user. This routine only calculates the enthalpy of a single phase stream. For a two-phase stream, the flash routine such as tpflash should be used to determine the overall stream enthalpy.

Function: CC_hxstream
Enthalpy calculation routine called from ADDH.

Prototype: void CC_hxstream(float xmol[], double t, double p, int iphase, float hx)

Parameters:
- Input: xmol[] component mole flow rate(lbmole/hr)
  - t  temperature in degree R
  - p  pressure in psia
  - iphase 0 = Liquid, 1 = Vapor

Output hx  Btu/hr
Return Value: void

Description:
Calculates enthalpy based on any enthalpy model. For given component flow rate, temperature and pressure, this routine calculates the enthalpy of the stream according to the external variable modeh.

The following table shows the value of modeh and corresponding model:

<table>
<thead>
<tr>
<th>modeh</th>
<th>H model</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Polynomial H</td>
</tr>
<tr>
<td>2</td>
<td>Redlich-Kwong</td>
</tr>
<tr>
<td>3</td>
<td>SRK</td>
</tr>
<tr>
<td>4</td>
<td>API SRK</td>
</tr>
<tr>
<td>5</td>
<td>Peng-Robinson</td>
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<td>6</td>
<td>Lee-Kesler</td>
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<tr>
<td>7</td>
<td>Latent Heat</td>
</tr>
<tr>
<td>8</td>
<td>Amine</td>
</tr>
<tr>
<td>9</td>
<td>No Enthalpy</td>
</tr>
<tr>
<td>10</td>
<td>Enthalpy Table</td>
</tr>
<tr>
<td>11</td>
<td>ADDH</td>
</tr>
<tr>
<td>12</td>
<td>Mixed Model</td>
</tr>
</tbody>
</table>

Function: CC_entropy
Entropy calculation routine.

Prototype: void CC_entropy(float xmol[], double t, double p, int iphase, float sx)

Parameters:
- Input: xmol[] component mole flow rate(lbmole/hr)
- t temperature in degree R
- p pressure in psia
- iphase 0 = Liquid, 1 = Vapor
- Output sx Btu/R/hr

Return Value: void

Description:
For given component flow rate, temperature and pressure, this routine calculates the entropy of the stream.

Function: CC_zfactor
Z factor calculation routine.

Prototype: void CC_zfactor(float xmol[], double t, double p, int iphase, float z, int ierr)
Parameters:

Input: xmol[] component mole flow rate (lbmole/hr)
       t       temperature in degree R
       p       pressure in psia
       0phase = Liquid, 1 = Vapor

Output: 
       z          Z factor
       ierr      = 0 normal return
                   > 0 error

Return Value:   void

Description:

For given component flow rate, temperature and pressure, this routine calculates Z factor of the stream.

-------------------------------------------------------------------------------

Function:   CC_ldense

Liquid density routine.

Prototype:  void CC_ldense(float xliq[], double t, double p, float dens)

Parameters:

Input : xliq[] component mole flow rate in lbmol/hr
       t       temperature in R
       p       pressure in psia

Output: dens liquid density in lb/ft3

Return Value:   void

Description:

For given component flow rate, temperature and pressure, this routine calculates the liquid density of the mixture.

-------------------------------------------------------------------------------

Function:   CC_vdense

Vapor density routine.

Prototype:  void CC_vdense(float xvap[], double t, double p, float dens)

Parameters:

Input : xvap[] component mole flow rate in lbmol/hr
       t       temperature in R
       p       pressure in psia

Output: dens Vapor density in lb/ft3

Return Value:   void
Description:
For given component flow rate, temperature and pressure, this routine calculates the vapor density of the mixture.

----------------------------------------------------------------------
Function: CC_vp
Function to calculate the pure component vapor pressure
Prototype: float CC_vp(int i, double t)

Parameters:
Input: i component POSITION. (base 0 = first component)
       t temperature in degree R

Return Value: float, vapor pressure in psia

Description:
Given component i and temperature, this routine calculates the vapor pressure.

----------------------------------------------------------------------
Function: CC_cp
Function to calculate the heat capacity of a mixture
Prototype: float CC_cp(float xmol[], double t, double p, int iphase)

Parameters:
Input:
       xmol[] mole flow lbmol/hr
       t temperature R
       p pressure psia
       iphase 0 = liquid phase
              1 = vapor phase

Return Value: float cp in Btu/R/lbmole

Description:
Given composition, temperature and pressure, this function calculates the heat capacity of the stream.

----------------------------------------------------------------------
Function: CC_cv
Function to calculate the cv of a mixture
Prototype: float CC_cv(float xmol[], double t, double p, int iphase)

Parameters:
Input:
       xmol[] mole flow lbmol/hr
Calculator/Parser Module

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- **t** temperature R
- **p** pressure psia
- **iphase** 0 = liquid phase
  1 = vapor phase

Return Value: float cv in Btu/R/lbmole

Description:

Given composition, temperature and pressure, this function calculates the cv of the stream.

---------------------------------------------

Function: **CC_lvisco**

Function to calculate the liquid viscosity of a mixture

Prototype: float CC_lvisco(float xmol[], double t, double p)

Parameters:

- **input**: xmol[] mole flow rate in lbmol/hr
  - **t** temperature in R
  - **p** pressure in psia

Return Value: float viscosity in CP

Description:

Given composition, temperature and pressure, this function calculates the viscosity of the liquid mixture.

---------------------------------------------

Function: **CC_gvisco**

Function to calculate the vapor viscosity of a mixture

Prototype: float CC_gvisco(float xmol[], double t, double p)

Parameters:

- **input**: xmol[] mole flow rate in lbmol/hr
  - **t** temperature in R
  - **p** pressure in psia

Return Value: float viscosity in CP

Description:

Given composition, temperature and pressure, this function calculates the viscosity of the vapor mixture.

---------------------------------------------

Function: **CC_lthc**

Function to calculate the liquid thermal conductivity of a mixture

Prototype: float CC_lthc(float xmol[], double t, double p)
Parameters:
  - input: xmol[] mole flow rate in lbmol/hr
  - t temperature in R
  - p pressure in psia

Return Value: float liquid thermal conductivity in Btu/hr ft² F / ft

Description:
  Given composition, temperature and pressure, this function calculates the thermal conductivity of the
  liquid mixture.

-------------------------------------------------------------------------------

Function: CC_vthc
  Function to calculate the vapor thermal conductivity of a mixture

Prototype: float CC_vthc(float xmol[], double t, double p)

Parameters:
  - input: xmol[] mole flow rate in lbmol/hr
  - t temperature in R
  - p pressure in psia

Return Value: float vapor thermal conductivity in Btu/hr ft² F / ft

Description:
  Given composition, temperature and pressure, this function calculates the thermal conductivity of the
  vapor mixture.

-------------------------------------------------------------------------------

Function: CC_surften
  Function to calculate the liquid surface tension of a mixture

Prototype: float CC_surften(float xmol[], double t, double p)

Parameters:
  - input: xmol[] mole flow rate in lbmol/hr
  - t temperature in R
  - p pressure in psia

Return Value: float liquid surface tension in dyne/cm

Description:
  Given composition, temperature and pressure, this function calculates the surface tension of the
  liquid mixture.

-------------------------------------------------------------------------------
SAMPLE FILES:

/*
   Cost estimation for heat exchangers
   
   Type
     = 0  Fixed head
     = 1  Kettle reboiler
     = 2  U-tube
   Material
     = 0  Carbon steel
     = 1  Stainless steel 316
     = 2  Stainless steel 304
     = 3  Stainless steel 347
     = 4  Nickel 200
     = 5  Monel 400
     = 6  Inconel 600
     = 7  Incoloy 825
     = 8  Titanium
     = 9  Hastelloy

*/

CHEMCAD_Link Parser
{
    int ID;
    int type, material;
    float p_design, fd, g1, g2, fm, p1, p2, fp, C;
    float area_ft2, log_area, bcost, uspec[300];
    int in_str1, in_str2;

    type = 0;
    material = 0;

    /* Get current equip parameters, 0 arg */
    CC_Get_Equip_Parameters(0,uspec);

    bcost = 0.;
    ID = uspec[1];
    printf("Preliminary Shell and Tube Heat Exchanger Cost Estimation\n\n");
    printf("Exchanger Cost for Equip. %d\n",ID);

    /* Area in ft2 */
    area_ft2 = uspec[29];

    /* Check range and give warnings */
    if( area_ft2 < 150. )
    {
        printf("Area %g < Amin (150 ft2)\n",area_ft2);
    }
    if( area_ft2 > 12000. )
}
{   printf("Area %g > Amax (12000 ft2)\n",area_ft2);
}
if( area_ft2 > 0. )
{
    /* Log area */
    log_area = log(area_ft2);
    /* Calc base cost of exchanger */
    bcost = exp(8.821 - 0.30863 * log_area + 0.0681 * log_area * log_area);
    printf("Area(ft2) = %g\n",area_ft2);
    printf("Base Cost = $%g\n",bcost);
    /* Calc type correction fd */
    if( type == 2 )
    {
        printf("U-tube\n");
        fd = exp(-0.9816 + 0.0830 * log_area);
    }else if( type == 1 )
    {
        printf("Kettle reboiler\n");
        fd = 1.35;
    }else
    {
        printf("Fixed head\n");
        fd = exp(-1.1156 + 0.0906 * log_area);
    }
    /* Calc material correction factor */
    if( material == 1 )
    {
        printf("Material = Stainless steel 316\n");
        g1 = 0.8603;
        g2 = 0.23296;
    }else if( material == 2 )
    {
        printf("Material = Stainless steel 304\n");
        g1 = 0.8193;
        g2 = 0.15984;
    }else if( material == 3 )
    {
        printf("Material = Stainless steel 347\n");
        g1 = 0.6116;
        g2 = 0.22186;
```c
} else if( material == 4 )
{
    printf("Material = Nickel 200\n");
    g1 = 1.5092;
    g2 = 0.60859;
}
else if( material == 5 )
{
    printf("Material = Monel 400 \n");
    g1 = 1.2989;
    g2 = 0.43377;
}
else if( material == 6 )
{
    printf("Material = Inconel 600\n");
    g1 = 1.204;
    g2 = 0.50764;
}
else if( material == 7 )
{
    printf("Material = Incoloy 825\n");
    g1 = 1.1854;
    g2 = 0.49706;
}
else if( material == 8 )
{
    printf("Material = Titanium\n");
    g1 = 1.5420;
    g2 = 0.42913;
}
else if( material == 9 )
{
    printf("Material = Hastelloy\n");
    g1 = 0.1549;
    g2 = 0.51774;
}
else
{
    printf("Material = Carbon steel\n");
    g2 = 0.0;
    g1 = 1.0;
}
f = g1 + g2 * log_area;
/* Get 1st inlet stream ID */
in_str1 = CC_Get_Value(3,ID,1,0);
```

/* Get 2nd inlet stream ID */
in_str2 = CC_Get_Value(3,ID,2,0);

/* Inlet pressure in psig */
p1 = CC_Get_Value(2,in_str1,2,0) - 14.696;
p2 = CC_Get_Value(2,in_str2,2,0) - 14.696;
p_design = p1;
if( p2 > p1 ) p_design = p2;
printf("Design pressure (psig) = %g\n",p_design);
if( p_design <= 300. )
{
    fp = 0.7771 + 0.04981 * log_area;
}
else if( p_design >= 600. )
{
    fp = 1.1400 + 0.12088 * log_area;
}
else
{
    fp = 1.0305 + 0.07140 * log_area;
}
C = fd * fm * fp * bcost;
printf("Exchanger cost = \$%g\n",C);
/* Put cost back to htxr */
uspec[48] = C;
CC_Put_Equip_Parameters(0,uspec);
}
else
{
    printf("No area data available.\n");
}

/* Sample calculator program
This program simulate a component separator
The flow rate, temperature, and pressure of the second output stream are specified and fixed. The program put the difference of the input stream and second output stream into the first output stream and perform an adiabatic flash calculation.
*/
CHEMCAD_Link CALC1
{
int nc, i, j;
float in1[100];
float tin;
float pin;
float vin;
float hin;
float out1[100], out2[100];
float tv, tl, hv, hl;
float t1out, t2out, p1out, p2out, h1out, h2out, v1out, v2out;
float vapor[100], xliq[100], xk[100], f_ions[100];

// Get input stream
CC_Get_Input_Stream(1, in1, tin, pin, vin, hin);

// Get 2nd output stream
CC_Get_Output_Stream(2, out2, t2out, p2out, v2out, h2out);

// Calc the difference and put in first output stream
nc = CC_No_Of_Components();
for (i = 0; i < nc; i++)
{
    out1[i] = in1[i] - out2[i];
    if (out1[i] < 0.)
    {
        printf("Error comp %d, output > input\n", i);
        out1[i] = 0.;
    }
}

// Assume adiabatic
h1out = hin - h2out;

// Call adiabatic flash calculation to get the output temperature
// and vapor fraction
CC_hspflash(out1, p1out, h1out, 0, tin, vapor, xliq, tv, tl, t1out, hv,
        hl, h1out, xk, v1out, f_ions);

// Put the results to output 1
CC_Put_Output_Stream(1, out1, t1out, p1out, v1out, h1out);
}
This program simulates the water entrainment into the oil phase. The calculator block has two input and two output streams. The two input streams come from the liquid outputs of a three phase flash calculation.

The first input is the oil stream that contains small amount of water dissolved in the oil phase. The second input stream contains all the free water. This program allows the user to specify the volumetric fraction of water to be carried into the oil phase and calculate the new compositions for the two output streams.

*/
CHEMCAD_Link CALC
{
    float factor;
    float t, p, vf, h_oil;
    float uspec[300];
    float oil_mol[100], wtr_mol[100];
    float entr_mol[100];
    float oilvol, entrain_wtrvol, h_wtr, wtrvol;
    int oilstr;
    int wstr;
    int nc;
    int i;
    /* Get the equipment parameters of the calculator */
    CC_Get_Equip_Parameters(0,uspec);
    factor = uspec[2];
    oilstr = 1;
    wstr = 2;
    /* Get oil stream */
    CC_Get_Input_Stream(oilstr, oil_mol, t, p, vf, h_oil);
    /* Calc oil volume rate */
    oilvol = CC_Get_Stream_Property(oil_mol,t,p,0.,h_oil,47);
    printf("oilvol = %g\n",oilvol);
    /* Calc entrained water volume */
    entrain_wtrvol = oilvol * factor;
    printf("factor = %g\n",factor);
    printf("entrain_wtrvol = %g\n",entrain_wtrvol);
    /* Get water stream */
    CC_Get_Input_Stream(wstr, wtr_mol, t, p, vf, h_wtr);
    /* Calc water volume rate */
    wtrvol = CC_Get_Stream_Property(wtr_mol,t,p,0.,h_wtr,47);
    printf("wtrvol = %g\n",wtrvol);
    if( wtrvol <= 0. )
    {
        
    
}
wtrvol = 0.00001;

/* Calc total wtr moles */
cn = CC_No_Of_Components();
for( i = 0; i < nc; i++)
{
    entr_mol[i] = wtr_mol[i] / wtrvol * entrain_wtrvol;
    /* Entrained flow rate can not be larger than the water flow rate */
    if( entr_mol[i] > wtr_mol[i] ) entr_mol[i] = wtr_mol[i];
}

/* New oil stream */
for( i = 0; i < nc; i++)
{
    oil_mol[i] = oil_mol[i] + entr_mol[i];
}

/* New oil enthalpy */
CC_enthalpy(oil_mol, t, p, 0, h_oil);

/* New water stream */
for( i = 0; i < nc; i++)
{
    wtr_mol[i] = wtr_mol[i] - entr_mol[i];
}

/* New water enthalpy */
CC_enthalpy(wtr_mol, t, p, 0, h_wtr);

/* Put oil back to process */
CC_Put_Output_Stream(oilstr, oil_mol, t, p, 0., h_oil);

/* Put water back to process */
CC_Put_Output_Stream(wtrstr, wtr_mol, t, p, 0., h_wtr);