# Calculation for CO<sub>2</sub> Compression and Refrigeration for Carbon Capture and Storage (CCS), Estimation of Energy Penalty

Kim, Seong Gyun

Department of Chemical Engineering, Hanyang University

CHE 1003: Chemical Engineering Thermodynamics 1

Bae, Young Chan

May 31, 2021

# **Table of Contents**

1	Intro	duction	3
	1.1	Overview on Carbon Capture and Storage (CCS)	3
	1.2	Topics Covered	4
2	Theo	oretical Background	5
	2.1	Setting Initial and Final Values and Constants	5
	2.2	Compression – Assuming Ideal-Gas State	6
	2.3	Compression – Using Lee-Kesler Equation of State	8
	2.4	Refrigeration for Low-pressure Compression	11
	2.5	Other Energy Requirements for CCS	12
3	Calc	ulation for Compression Processes	12
	3.1	High-pressure Compression to 11 MPa	12
	3.2	Low-pressure Compression to 2 MPa	13
	3.3	Refrigeration Process for Low-pressure Compression	15
4	Sum	mary of Calculation Results	17
5	Conc	clusion	19
6	Refe	rences	20
7	App	endix	22
	7.1	Saturated/Superheated Table for Ammonia	22
	7.2	P-H Diagram for Ammonia	24
	7.3	MATLAB® Implementation	25
	7.4	MATLAB® Execution Results	31

#### 1 Introduction

## 1.1 Overview on Carbon Capture and Storage (CCS)

In a world where fossil fuel supplies about 80 percent of the total energy that we produce [1], the carbon dioxide produced from burning fossil fuel is a major contributor to global warming and other environmental issues. Therefore, in order to prevent carbon dioxide from getting into the atmosphere, we would use a technology called Carbon Capture and Storage (CCS) to separate carbon dioxide from the flue gas and store it underground or under water. This would prevent carbon dioxide from getting into the atmosphere and slow down the rate of global warming. Currently, there are three main types of CCS in use. The three main types of pre-combustion capture, post-combustion capture, and oxyfuel process [2].

The first type of CCS is pre-combustion capture. Pre-combustion capture involves separation of carbon dioxide from fossil fuel before the combustion begins [3]. There are a couple of ways to do this. One widely used method is to first gasify the fossil fuel with oxygen and pressure. This synthesis gas (syngas) would be made up of carbon monoxide and hydrogen. Then steam is added to syngas and passed through a bed packed with catalysts. Through this reaction, carbon monoxide would be turned into carbon dioxide [4]. However, before syngas can be used to drive the turbine, the carbon dioxide needs to be separated beforehand. In order for this to be achieved, this process involves a physical solvent such as rectisol and selexol. Carbon dioxide is dissolved using these solvents at higher pressure and can be released when the pressure is lower. Using this technique, carbon dioxide can be stripped away and safely stored while the hydrogen gas is used to spin the turbine [4]. One big advantage of this system is that the physical solvents are available at low costs and require relatively low amounts of energy. However, the main disadvantage of this system is that it requires a chemical plant in front of the turbine which can be very difficult to achieve depending on the plant [4].

The second type of CCS is post-combustion capture. During post-combustion capture, carbon dioxide is removed from the flue gas through a thermal power plant combustion chamber. Flue gas from coal fired power plants typically have a carbon dioxide concentration of less than 15 percent and less than 8 percent from a gas fired power plant [4]. Because there are such low amounts of carbon dioxide present in flue gas, it is challenging to develop a cost-effective way to capture carbon dioxide. One method is to use amines to separate carbon dioxide from the flue gas. In order to do this, first, fossil fuel is burned as normal. Then, before the flue gas is released into the atmosphere, it passes through an absorber column. In this column, it is filled with liquid solvents called amines. These amines absorb the carbon dioxide before the flue gas is released into the atmosphere. Then super-heated steam is applied which releases the carbon dioxide from the amines. After this process, the carbon dioxide can now be safely stored [5]. One huge advantage of this method is that the required equipment can be easily attached to already existing power plants. However, the disadvantage of this method is that due to the concentration of carbon dioxide consisted in the flue gas being low, the equipment are rather large and are expensive.

The last type of CCS is oxyfuel process. As it was mentioned before, because the flue gas contains such low concentrations of carbon dioxide, it made it even more difficult to separate carbon dioxide from the flue gas. Therefore, during oxyfuel process, combustion of fossil fuel is taken place where it

is full of oxygen. When this happens, the flue gas is mostly composed of carbon dioxide and water vapor. Water is easily removed with condensation while carbon dioxide is remained to stored. Then we use this combination and run it through the turbine to produce electricity [4]. The advantage of this method is that because the volume of flue gas has decreased and the concentration of carbon dioxide has increased, the cost of this method is relatively cheaper [4].

The final step in Carbon Capture and Storage is to store the carbon dioxide. Carbon dioxide is compressed into the liquid phase and is stored in a few different ways. One way is to store them in tanks and transport them by using trucks. While another way is to use pipelines to carry carbon dioxide to its appropriate places. The pipeline would eventually lead to a deep underground layer of porous rock where the carbon dioxide would be stored. This process is known as geo-sequestration [6]. When carbon dioxide is injected into this layer of porous rock, it would spread through the gaps and fill the holes. Over time, these carbon dioxide would react with the surround rocks and create carbonate minerals such as calcium carbonate, magnesium carbonate, or ferrous carbonate [6].

Currently, most of the plants that does CCS uses the post-combustion capture method. This is because amine scrubbing is the most developed capture option. Oxyfuel capture method still needs some more development and pre-combustion method only works with a limited type of fossil fuel [5]. Therefore, we will also focus mostly on post-combustion capture method for this term paper.

## 1.2 Topics Covered

In this paper we would be covering following topics learned in the Chemical Engineering Thermodynamics I course (CHE 1003):

- Equations of state ideal-gas state, Lee-Kesler equation of state
- MCPH, MCPS functions for calculating isobaric and isochoric heat capacities
- Residual thermodynamic properties residual enthalpy, residual entropy
- Compression isentropic, isothermal, actual compression of CO<sub>2</sub>
- Refrigeration vapor-compression cycle of NH<sub>3</sub>
- Numerical methods to estimate roots open method, approximation by iteration
- Reading phase diagrams P-H diagram of NH<sub>3</sub>
- Reading saturation/superheated tables NH<sub>3</sub>

## 2 Theoretical Background

# 2.1 Setting Initial and Final Values and Constants

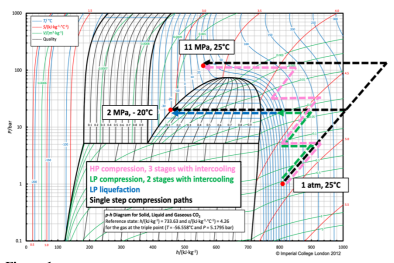
A post-combustion CCS process consists of 1) CO<sub>2</sub> separation from the flue gas, 2) high- or low-pressure compression and liquefaction processes, followed by 3) transport and storage processes. In this study, two paths have been thermodynamically analyzed and studied: the compression of CO<sub>2</sub> to a high pressure and ambient temperature (supercritical state, 11 MPa, 25 °C) suitable for pipeline transport, and the low-pressure compression and refrigeration (saturated liquid state, 2 MPa, -20 °C) suitable for tanker transport. The final property values are chosen in accordance with actual industrial processes as well as studies that have been conducted on CCS [5], [7]. The CO<sub>2</sub> stream from the separation process is assumed to enter the compression processes at ambient temperature and the initial pressure of the atmospheric pressure (25 °C, 0.101325 MPa).

**Table 1**Two paths of CO<sub>2</sub> compression and transportation

Path 1	High-pressure comp	pression	Transport by pipelines
Path 2	Low-pressure compression	Refrigeration (Liquefaction)	Transport by trucks

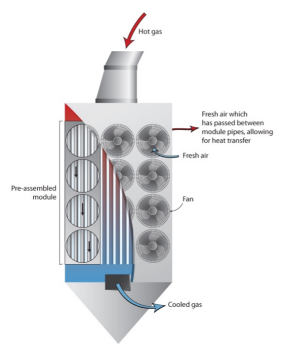
Although the polytropic efficiency of a compressor in a CCS process is reported as high as 94% in literature [7], in the present report, the compressors are assumed to perform at 65% efficiency, which is realistically set in accordance with the industrial applications and studies [5], [8], [9].

According to Nored [10], optimal compression ratio for CO<sub>2</sub> is specified by compressor manufacturers as 3.5 to 4, whereas compression ratio required for CCS ranges from 19 (to 2MPa) to



**Figure 1**P-H diagram for CO<sub>2</sub>, with compression paths annotated Original diagram prepared by Imperial College London [11]

110 (to 11MPa). Hence, in this report, multiple stages of compression and intercooling stages are planned with regard to keeping compression ratio of each compressor between 3 to 4.5. Lowering entering temperature of the flow by intercooling stages in turn lowers the energy required to compress the gas in each compressor unit. Figure 1 demonstrates three-step compression path to 11 MPa, two-step compression path to 2 MPa followed by refrigeration, and single-step paths on the P-H diagram of CO<sub>2</sub> originally prepared by Imperial College London [11].



**Figure 2.** A gas-to-air heat exchanger manufactured by Redecam group [12]

Intercooling stages between compression stages are done by gas-to-air cooling refrigerators, where the ambient air works as the refrigerant. This process is suitable for cooling CO<sub>2</sub> stream because 1) it is dry gas, moisture removed in the earlier separation process, and 2) it is observed to heat up to 500 K when compression ratio is kept optimal, which is far below the maximum temperature that gas-to-air heat exchangers can tolerate (~750 °C) [12].

For example, in the gas-to-air intercooling device manufactured by Redecam group shown in **Figure 2**, the CO<sub>2</sub> flow would enter at the hood on top, pass through the tubes and leave at the opening located at the bottom of the hopper [13].

Power requirement for operating intercooling between compression stages is included as a portion of 'parasitic' power requirements when calculating the total energy penalty.

# 2.2 Compression – Assuming Ideal-Gas State

## Step 1: Setting equations for the minimum required work for compression

If CO<sub>2</sub> is assumed to be in its ideal-gas state during the compression process, the isentropic work (minimum work) for compression is given by:

$$W_{comp,isen} = \Delta H_S = R \int_{T_c}^{T_{2,isen}} \frac{C_P^{ig}}{R} dT$$

Where  $C_P^{ig}/R$  is given by a function of T:

$$C_{p}^{ig}/R = A + BT + CT^{2} + DT^{-2}$$

And A, B, C, and D are constants specifically determined for CO<sub>2</sub>:

$$A = 5.457$$
,  $B = 1.045 \times 10^{-3}$ ,  $C = 0$ ,  $D = 1.157 \times 10^{5}$ 

## **Step 2: Finding the final temperature after isentropic compression**

 $T_{2,isen}$  has to be determined by iterations with arbitrarily chosen initial value:

$$T_{2,isen}[1] = T_1 + 100$$

Then, with the mean heat capacity acquired in the previous step,  $T_{2,isen}[i+1]$  is determined:

$$T_{2,isen}[i+1] = T_1 \left(\frac{P_2}{P_1}\right)^{R/\langle C_P \rangle_S},$$

Where the exponent can be calculated as the reciprocal of:

$$\frac{\langle C_P \rangle_S}{R} = \text{MCPS} = \int_{T_1}^{T_{2,isen}[i]} \frac{A + BT + CT^2 + DT^{-2}}{R} \frac{dT}{T} \times \frac{1}{\ln\left(\frac{T_{2,isen}[i]}{T_1}\right)}$$

If the (i+1)-th value does not significantly differ from the i-th value anymore, the iteration stops and  $T_{2.isen}$  is determined:

If

$$\frac{T_{2,isen}[i+1] - T_{2,isen}[i]}{T_{2,isen}[i]} < 0.01\%$$

Then

$$T_{2,isen} = T_{2,isen}[i+1]$$

## Step 3: Evaluating the real work required from the isentropic work and efficiency

With the  $T_{2,isen}$  determined in the previous step, the isentropic work for compression can be calculated:

$$W_{comp,isen} = \Delta H_S = R \int_{T_1}^{T_{2,isen}} \frac{C_P^{ig}}{R} dT$$

Then the actual work required can be calculated by dividing minimum work by the efficiency of the compressor:

$$W_{comp} = \frac{W_{comp,isen}}{\eta_{comp}}$$

# Step 4: Finding the actual final temperature after compression

The actual final temperature can be found by iterating until the value converges, in a similar manner as in **Step 2**, setting an arbitrary initial guess somewhat higher than the initial temperature.

$$T_2[i+1] = T_1 + \frac{W_{comp}}{\langle C_P \rangle_H}$$

Where

$$\langle C_P \rangle_H = R \times MCPH = R \times \int_{T_1}^{T_2[i]} (A + BT + CT^2 + DT^{-2}) dT \times \frac{1}{T_2[i] - T_1}$$

## 2.3 Compression – Using Lee-Kesler Equation of State

Lee and Kesler proposed a three-parametric analytic equation of state that gives reasonable estimation of thermodynamic properties of real substances [14]. Mazzoccoli et al. have reported that the Lee-Kesler equation of state (LK EOS) is one of the most accurate methods for calculating the enthalpy of gas [15]. In this section, enthalpy and entropy change of CO<sub>2</sub> in the compression process are calculated by employing the concept of residual properties, which are calculated by:

$$\begin{split} \frac{H^R}{RT_c} &= \left[\frac{H^R}{RT_c}\right]^{(0)} + \frac{\omega}{\omega^{(r)}} \left\{ \left[\frac{H^R}{RT_c}\right]^{(r)} - \left[\frac{H^R}{RT_c}\right]^{(0)} \right\} \\ \frac{S^R}{R} &= \left[\frac{S^R}{R}\right]^{(0)} + \frac{\omega}{\omega^{(r)}} \left\{ \left[\frac{S^R}{R}\right]^{(r)} - \left[\frac{S^R}{R}\right]^{(0)} \right\} \end{split}$$

Where  $\omega$  denotes acentric factor of the substance of interest.

The residual properties are calculated twice with different sets of constants, for reference fluid and then for real fluid. Superscripts r and 0 denote real and reference fluid, respectively. The residual properties can be derived from the compressibility factor and directly calculated from specified reduced temperature and specific volume. The resulting equations are:

$$\frac{H^R}{RT_c} = T_r \left( Z - 1 - \frac{b_2 + \frac{2b_3}{T_r} + \frac{3b_4}{T_r^2}}{T_r V_r} - \frac{c_2 - \frac{3c_3}{T_r^2}}{2T_r V_r^2} + \frac{d_2}{5T_r V_r^5} + 3E \right)$$

$$\frac{S^R}{R} = \ln Z - \frac{b_1 + \frac{b_3}{T_r^2} + \frac{2b_4}{T_r^3}}{V_r} - \frac{c_1 - \frac{2c_3}{T_r^3}}{2V_r^2} - \frac{d_1}{5V_r^5} + 2E - T_r$$

Where

$$E = \frac{c_4}{2T_r^3 \gamma} \left( \beta + 1 - \left( \beta + 1 + \frac{\gamma}{V_r^2} \right) \exp\left( -\frac{\gamma}{V_r^2} \right) \right)$$

$$Z = \frac{P_r V_r}{T_r} = 1 + \frac{B}{V_r} + \frac{C}{V_r^2} + \frac{D}{V_r^5} + \frac{c_4}{T_r^3 V_r^3} \left( \beta + \frac{\gamma}{V_r^2} \right) \exp\left( -\frac{\gamma}{V_r^2} \right)$$

$$P_r = \frac{P}{P_c}, \qquad T_r = \frac{T}{Tc}, \qquad V_r = \frac{P_c V}{RT_c}$$

Constants calibrated by Lee and Kesler [14] are shown in the **Table 2**. Detailed explanation for derivation of the residual properties and the calculation process are not included in this report, since

it is not of the main interest of this project. Instead, calculating the work required for the compression process with the residual properties obtained by LK EOS is explained in detail.

**Table 2**Constants for Lee-Kesler equation of state [14]

Constant	Simp	le fluids Reference Fluids
$b_1$	0.1181193	0.2026579
$b_2$	0.265728	0.331511
$b_3$	0.154790	0.027655
$b_4$	0.030323	0.203488
$c_1$	0.0236744	0.0313385
$c_2$	0.0186984	0.0503618
$c_3$	0.0	0.016901
$c_4$	0.042724	0.041577
$d_1\times 10^4$	0.155488	0.48736
$d_2 \times 10^4$	0.623689	0.0740336
β	0.65392	1.226
γ	0.060167	0.03754

## Step 1: Setting equations for the minimum required work for compression

By using residual properties calculated by LK EOS, the isentropic work (minimum work) and change of entropy for compression are given by:

$$W_{comp,isen} = \Delta H = -H_1^R + \Delta H_S^{ig} + H_2^R$$

$$\Delta S_{isen} = -S_1^R + \Delta S_S^{ig} + S_2^R (=0)$$

Where  $\Delta H_S^{ig}$  is acquired in the same manner as in **2.2**.  $H_1^R$ ,  $S_1^R$  and  $H_2^R$ ,  $S_2^R$  are residual properties in the initial and the final state respectively.

## Step 2: Finding the final temperature after isentropic compression

 $T_{2,isen}$  has to be determined by iterations with arbitrarily chosen initial value. Since the compression is isentropic, iterating the calculation of entropy change with  $T_{2,isen}$  until total entropy change becomes reasonably close to 0 can approximate the value of  $T_{2,isen}$ .

In our MATLAB® code,  $T_{2,isen}$  is programmed to decrease if  $\Delta S$  is calculated to be positive and increase if  $\Delta S$  is calculated to be negative. The calculation is repeated with exponentially decreasing step size to determine the value of  $T_{2,isen}$  to the third decimal digit:

If

$$\Delta S > 0$$

Then

$$T_{2.isen}[i+1] = T_{2.isen}[i] - 10^{(2-a)}$$

If

$$\Delta S < 0$$

Then

$$T_{2.isen}[i+1] = T_{2.isen}[i] + 10^{(2-a)}$$

Where a is initially set to 1 and increased when sign change of  $\Delta S$  is observed.

For example, firstly,  $T_{isen}$  is decreased by  $10 \ (10^{(2-1)})$  in every iteration if initially calculated  $\Delta S$  is positive. Then, when  $\Delta S$  is first calculated to be negative after some iterations, a is increased by 1; now  $T_{isen}$  is increased by 1  $(10^{(2-2)})$  in every iteration until  $\Delta S$  is calculated to be positive again. This is repeated until a exceeds 5, meaning  $T_{isen}$  has been determined to the accuracy of 0.001  $(10^{(2-5)})$ . The MATLB® code for this process can be found in **Appendix 7.3**.

# Step 3: Evaluating the real work required from the isentropic work and efficiency

With the  $T_{isen}$  determined in **Step 2**, isentropic work can be calculated:

$$W_{comp,isen} = \Delta H = -H_1^R + \Delta H_S^{ig} + H_2^R$$

Then the actual work required can be calculated by dividing minimum work by the efficiency of the compressor:

$$W_{comp} = \frac{W_{comp,isen}}{\eta_{comp}}$$

To reduce running time of the code, a procedure to determine the actual final temperature  $T_2$  from the actual work is not implemented. It is not of great importance after all, since the values calculated by assuming ideal-gas state already give reasonable estimation.

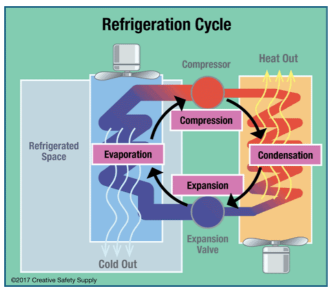
Values and units of variables used in the calculations are presented below in **Table 3**.

Table 2
Values and units of constants and variables used in the calculation

Symbol	Variable	Value	Unit
R	Gas Constant	8.314462	J/mol·K
$T_1$	Initial temperature	25	°C
$P_1$	Initial pressure	101.325	kPa
$\eta_{comp}$	Efficiency of a compressor	65.0	%
$M_{\rm CO_2}$	Molar mass of CO <sub>2</sub>	44.01	g/mol
	High-pressure compression		
	Final pressure	11,000	kPa
	Number of compressors	2-6	-
	Low-pressure compression		
	Final pressure	2,000	kPa
	Number of compressors	1-5	-

## 2.4 Refrigeration for Low-pressure Compression

CO<sub>2</sub> compressed to low pressure goes through the refrigeration process to be suitable for storage in tanks, which are then transported by trucks, etc. In this report, a vapor-compression cycle with Ammonia as the refrigerant is analyzed. The evaporator temperature is set as -30°C and the condenser temperature 26°C, in accordance with actual industrial processes [5]. Also, heat removed from unit mass of CO<sub>2</sub> stream is assumed to be 330 kJ/kg-CO<sub>2</sub>, the value found from literature [16].



**Figure 3** Vapor-compression refrigeration cycle using NH<sub>3</sub> as refrigerant [17]

Ammonia (NH<sub>3</sub>) was chosen as the refrigerant in this process for its superior thermodynamic properties, low chemical and piping cost, and environmentally friendly character [17]. Previous studies have reported that NH<sub>3</sub> does not contribute to ozone depletion and global warming when released to the atmosphere [18].

The compressor unit employed for the vapor-compression cycle is assumed to operate at 80% efficiency, same as the compressor unit studied in Example 9.1 in the textbook [19].

The cycle is analyzed in four states:

- State 1: Saturated liquid + saturated vapor (Throttle valve → Evaporator)
- State 2: Saturated vapor
   (Evaporator → Compressor)
- State 3: Superheated vapor (Compressor → Condenser)
- State 4: Saturated liquid (Condenser → Throttle valve)

# 2.5 Other Energy Requirements for CCS

Before the compression and liquefaction processes, separating CO<sub>2</sub> from the flue gas requires part of the steam from the turbines to be diverged with results in consumption of the electricity generated from the turbines. Page, Williamson, and Mage [5] have given a reasonable estimation of power loss due to CO<sub>2</sub> separation process as 0.500 MJ/kg-CO<sub>2</sub> and other parasitic requirements for devices to run the CCS process (i.e. fans, pumps, etc.) as 0.470 MJ/kg-CO<sub>2</sub>.

## **3** Calculation for Compression Processes

# 3.1 High-pressure Compression to 11 MPa

Compression from atmospheric pressure to 11 MPa has the compression ratio of 108.562 (=11/0.101325). Simulating the isentropic compression processes by MATLAB®, it is observed that increasing number of compression stages results in decrease in the work required for total compression process as well as maximum temperature of the gas ( $T_2$ ). The single compression ratio was observed to be closest to optimal range when 3 or 4 compression stages are employed. Hypothetically setting the number of compression stages as big as 1000 where temperature change in each step is very small, the near-isothermal case was also calculated as reference. The results by assuming ideal-gas state and LK EOS are shown in **Table 4** and **Table 5**, respectively.

**Table 3**Result of simulation with different numbers of compression stages (11 MPa, ideal-gas state)

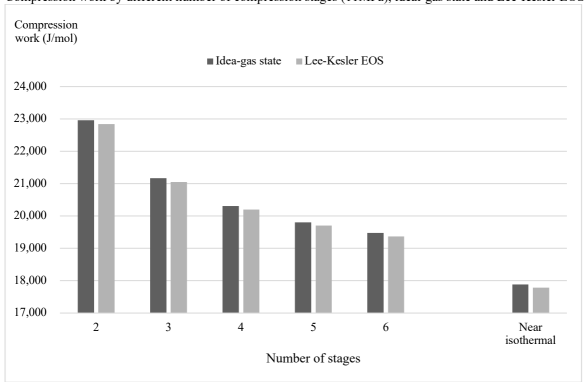
Number of compression stages	Single compression ratio	Maximum temperature (K)	Total work for compression (J/mol)
2	10.419	562.841	22964.5
3	4.770	466.986	21165.6
4	3.228	422.394	20308.4
5	2.553	396.557	19806.3
6	2.184	379.671	19476.3
1000 (near-isothermal)	1.005	298.631	17886.2

**Table 4**Result of simulation with different numbers of compression stages (11 MPa, Lee-Kesler EOS)

Number of compression stages	Single compression ratio	Total work for compression (J/mol)
2	10.419	22841.6
3	4.770	21051.1
4	3.228	20199.4
5	2.553	19700.8
6	2.184	19373.2
1000 (near-isothermal)	1.005	17788.4

The results from both calculations are presented together in **Chart 1**.

Chart 1
Compression work by different number of compression stages (11MPa), ideal-gas state and Lee-Kesler EOS



# 3.2 Low-pressure Compression to 2 MPa

Compression from atmospheric pressure to 2 MPa has the compression ratio of 19.738 (=2/0.101325). In a similar manner to **3.1**, increasing number of compression stages resulted in decrease in the work required for compression as well as maximum temperature. The single compression ratio was observed to be optimal when 2 compression stages are employed. Hypothetical near-isothermal process with 1000 compressors is included here as well. Results of

calculations by ideal-gas assumption and LK EOS are presented in **Table 6** and **Table 7**, respectively.

**Table 5**Result of simulation with different numbers of compression stages (2 MPa, ideal-gas state)

Number of compression stages	Single compression ratio	Maximum temperature (K)	Total work for compression (J/mol)
1	19.738	648.786	15607.7
2	4.443	458.714	13367.5
3	2.703	402.752	12679.7
4	2.108	375.860	12345.4
5	1.816	360.018	12147.6
1000 (near-isothermal)	1.003	298.456	11378.8

**Table 6**Result of simulation with different numbers of compression stages (2 MPa, Lee-Kesler EOS)

Number of compression stages	Single compression ratio	Total work for compression (J/mol)
1	19.738	15530.8
2	4.443	13294.9
3	2.703	12612.0
4	2.108	12280.0
5	1.816	12083.7
1000 (near-isothermal)	1.003	11271.4

The results from both calculations are presented together in **Chart 2**.

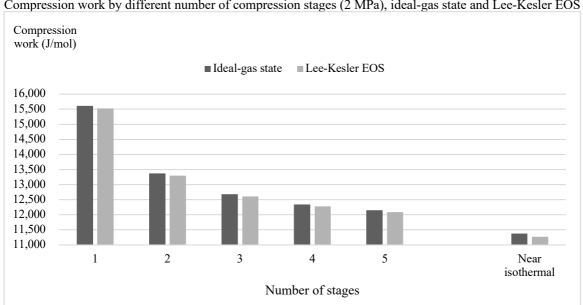


Chart 2
Compression work by different number of compression stages (2 MPa), ideal-gas state and Lee-Kesler EOS

# 3.3 Refrigeration Process for Low-pressure Compression

The vapor-compression cycle of NH<sub>3</sub> can be presented on the P-H diagram, acquired from [20]. Note that the isentropic compression is presented with the arrow with dashed line  $(2\rightarrow 3')$ .

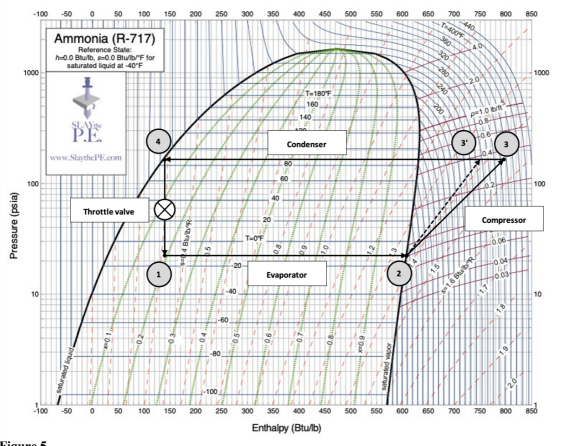


Figure 5 Vapor-compression cycle of NH<sub>3</sub> in US customary units. Original diagram included in **Appendix 7.1** [20].

## Step 1: Finding thermodynamic properties of states 2 and 4

Enthalpies from states 2 and 4 are read directly from the saturation table of NH<sub>3</sub> (**Appendix 7.1**). The entry at -30°C indicates that NH<sub>3</sub> vaporizes in the evaporator at a pressure of 1.196 bar, which corresponds to 119.6 kPa. Its properties as a saturated vapor at these conditions are:

$$H_2 = 1405.6 \frac{\text{kJ}}{\text{kg}}, \qquad S_2 = 5.785 \frac{\text{kJ}}{\text{kg} \cdot \text{K}}$$

The entry at 26°C in **Appendix 7.1** indicates that NH<sub>3</sub> condenses at 10.34 bar. Its enthalpy as a saturated liquid at these conditions is:

$$H_4 = 303.7 \frac{\text{kJ}}{\text{kg}}$$

From this, heat removed by unit mass of NH<sub>3</sub> as the refrigerant is calculated:

$$H_1 - H_2 = H_4 - H_2 = 1405.6 \frac{\text{kJ}}{\text{kg}} - 303.7 \frac{\text{kJ}}{\text{kg}} = 1101.9 \frac{\text{kJ}}{\text{kg}}$$

## Step 2: Evaluating the mass ratio of NH<sub>3</sub> and CO<sub>2</sub>

Since the basis of this analysis is 1kg of CO<sub>2</sub>, mass of NH<sub>3</sub> required for refrigeration process for 1kg of CO<sub>2</sub> stream can be calculated. Knowing that the heat removed from CO<sub>2</sub> for 1kg of CO<sub>2</sub> is 330 MJ/kg [16],

$$\frac{330 \text{ MJ}}{1 \text{ kg-CO}_2} \times \frac{1 \text{ kg-NH}_3}{1101.9 \text{ kJ}} = 0.2994 \frac{\text{kg-NH}_3}{\text{kg-CO}_2}$$

This mass ratio is later used for calculating energy consumption of refrigeration system for unit mass of CO<sub>2</sub> liquefied.

# Step 3: Finding isentropic enthalpy change $(H'_3)$ and actual enthalpy change $(H_3)$

Since the compression step is isentropic,

$$S_3' = S_2 = 5.875 \frac{\text{kJ}}{\text{kg} \cdot \text{K}}$$

Knowing the pressure value  $P_3 = 10.34$  bar and entropy value  $S_3'$ ,  $H_3'$  can be specified from the P-H diagram for NH<sub>3</sub> (**Figure 3**, or **Appendix 7.2**):

$$H_3' = 750 \frac{\text{Btu}}{\text{lb}} = 1743.3 \frac{\text{kJ}}{\text{kg}}$$

Then the isentropic enthalpy change is determined:

$$H_3' - H_2 = 1743.3 \frac{\text{kJ}}{\text{kg}} - 1405.6 \frac{\text{kJ}}{\text{kg}} = 337.7 \frac{\text{kJ}}{\text{kg}}$$

# **Step 4: Calculating real enthalpy change**

Assuming the compressor operates at efficiency of 80%,

$$H_3 - H_2 = \frac{H_3' - H_2}{0.8} = \frac{337.7 \text{ kJ/kg}}{0.8} = 422.1 \frac{\text{kJ}}{\text{kg}}$$

Which implies 422.1 kJ of work is required to operate the compressor for 1kg of NH<sub>3</sub>. By multiplying the mass ratio acquired from **Step 2**, energy requirement for 1kg of CO<sub>2</sub> is calculated and converted to the unit of MJ/kg-CO<sub>2</sub>:

$$422.1 \frac{\text{kJ}}{\text{kg-NH}_3} \times 0.2994 \frac{\text{kg-NH}_3}{\text{kg-CO}_2} = 126.4 \frac{\text{kJ}}{\text{kg-CO}_2} = 0.1264 \frac{\text{MJ}}{\text{kg-CO}_2}$$

## 4 Summary of Calculation Results

Total work requirement is calculated as sum of compression work, refrigeration work, and other energy requirements including separation (0.500MJ/kg) and parasitic requirements (0.470 MJ/kg). Energy penalty is calculated by the fraction of total work requirement to the emission intensity, i.e., energy produced per unit mass of CO<sub>2</sub> emitted to atmosphere. In this report, the emission factor of lignite coal, 95.2 tCO<sub>2</sub>/TJ, reported by Centre for Advanced Engineering [21] was taken as the reference value, which corresponds to 3.3075 MJ/kg with 31.5% efficiency. These values and relations are shown in the equations below.

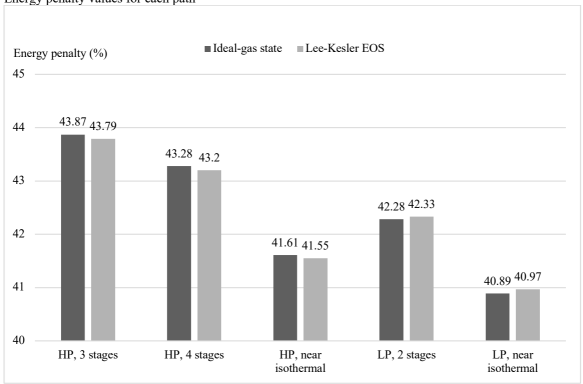
- Separation work requirement = 0.500 MJ/kg
- Parasitic work requirement = 0.470 MJ/kg
- Emission density = 3.3075 MJ/kg
- Total work requirement = Compression work
  - + Refrigeration work
  - + Other work requirements
- Energy penalty =  $\frac{\text{Total work requirement}}{\text{Emission density}} = \frac{\text{Total work requirement}}{3.3075 \, \text{MJ/kg-CO}_2} \times 100\%$

All the results calculated using these values are converted to the unit of MJ/kg-CO<sub>2</sub> and summarized in **Table 8**, **Chart 3**, and **Chart 4**.

**Table 7**Summary of the results acquired from the calculations

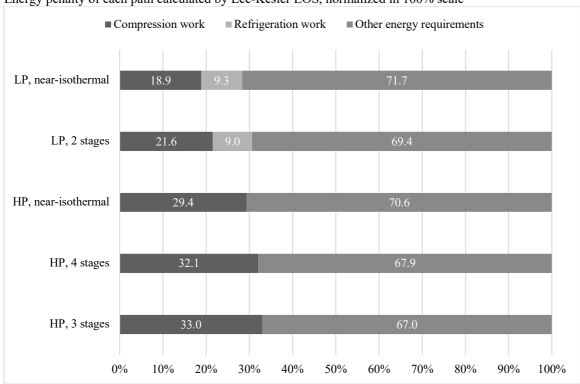
Process path	Number of compressors	Compression work (MJ/kg)	Refrigeration work (MJ/kg)	Total work requirement (MJ/kg)	Energy penalty (%)
	3	0.48093	-	1.4509	43.87
HP (11 MPa) (ideal-gas)	4	0.46145	-	1.4315	43.28
(lucal-gas)	near- isothermal	0.40641	-	1.3764	41.61
	3	0.47832	-	1.4483	43.79
HP (11 MPa) (LK EOS)	4	0.45897	-	1.4290	43.20
(LK E03)	near- isothermal	0.40419	-	1.3742	41.55
LP (2 MPa)	2	0.30374	0.12610	1.3985	42.28
(ideal-gas)	near- isothermal	0.25855	0.12640	1.3525	40.89
LP (2 MPa)	2	0.30209		1.4001	42.33
(LK EOS)	near- isothermal	0.25611	0.12640	1.3550	40.97

**Chart 3** Energy penalty values for each path



To see the contribution of energy requirement for each process in the CCS system, energy penalty values of each process calculated by Lee-Kesler EOS (compression and refrigeration) and given by literature (separation and other requirements) were shown normalized in 100% scale in **Chart 4**. The

energy requirement for separation process and parasitic requirements take up roughly 70% of the total energy penalty, while the rest 30% is attributed to compression and refrigeration processes.



**Chart 4** Energy penalty of each path calculated by Lee-Kesler EOS, normalized in 100% scale

## 5 Conclusion

This analysis result demonstrated that implementation of a post-combustion CCS to a coal power plant would result in significant energy penalty, requiring 40-45% of the energy generated from the plant. Even in the hypothetical case where the compression is done in a nearly isothermal way, more than 40% of energy penalty was calculated. This result roughly agrees with previously conducted thermodynamic and economic evaluations on post-combustion CCS [1], [5], [22].

It must be noted that our calculation is carried out under rather optimistic assumptions, 80% of compressor efficiency in the refrigeration process and 100% of CO<sub>2</sub> from flue gas being captured. Thus, more thorough and comprehensive analysis on CCS could demonstrate even higher energy penalty.

According to Page et al [5], The type of power plant from which carbon dioxide is captured has a large influence on the energy penalty. A standard lignite coal power plant with 31.5% efficiency was considered in this report. However, recently developed of ultra-supercritical power plants have been reported to achieve the net efficiency as high as 47.6% with emission factor 5.14 MJ/kg without CCS (cf. 3.31 MJ/kg, emission factor of standard lignite coal power plant used in our calculation) [23], which exhibits strong potential to reduce CCS cost as well.

Despite the inevitable energy and efficiency penalties CCS imposes, as pointed out by energy analysts Baylin-Stern and Berghout [24], "achieving net-zero goals will be virtually impossible without CCS." Although CCS technologies have not yet reached the point where it is economically attractive as shown in this report, efforts to lower the cost of the technology and accelerate more deployment have been made in both policymaking and technology developments as clean energy transition with mitigation of the impact of carbon emission to the global environment remains one of the biggest challenges in our generation.

## 6 References

- [1] F. P. Berlin, "Comparative Evaluation of Power Plants with CO 2 Capture: Thermodynamic, Economic and Environmental Performance Comparative Evaluation of Power Plants with CO 2 Capture: Thermodynamic, Economic and Environmental Performance."
- [2] S. Jackson and E. Brodal, "Optimization of the energy consumption of a carbon capture and sequestration related carbon dioxide compression processes," *Energies*, vol. 12, no. 9, 2019, doi: 10.3390/en12091603.
- [3] W. L. Theo, J. S. Lim, H. Hashim, A. A. Mustaffa, and W. S. Ho, "Review of pre-combustion capture and ionic liquid in carbon capture and storage," *Appl. Energy*, vol. 183, pp. 1633–1663, 2016, doi: 10.1016/j.apenergy.2016.09.103.
- [4] M. K. Mondal, H. K. Balsora, and P. Varshney, "Progress and trends in CO2 capture/separation technologies: A review," *Energy*, vol. 46, no. 1, pp. 431–441, 2012, doi: 10.1016/j.energy.2012.08.006.
- [5] S. C. Page, A. G. Williamson, and I. G. Mason, "Carbon capture and storage: Fundamental thermodynamics and current technology," *Energy Policy*, vol. 37, no. 9, pp. 3314–3324, 2009, doi: 10.1016/j.enpol.2008.10.028.
- [6] R. Leuning, D. Etheridge, A. Luhar, and B. Dunse, "Atmospheric monitoring and verification technologies for CO 2 geosequestration," vol. 2, pp. 401–414, 2008, doi: 10.1016/j.ijggc.2008.01.002.
- [7] F. Petrakopoulou, "Comparative Evaluation of Power Plants with CO2 Capture: Thermodynamic, Economic and Environmental Performance," 2010.
- [8] W. Fu, L. Wang, and Y. Yang, "Optimal design for double reheat coal-fired power plants with post-combustion CO2 capture: A novel thermal system integration with a carbon capture turbine," *Energy*, vol. 221, p. 119838, 2021, doi: 10.1016/j.energy.2021.119838.
- [9] H. A. Muhammad *et al.*, "A comprehensive thermodynamic performance assessment of CO2 liquefaction and pressurization system using a heat pump for carbon capture and storage (CCS) process," *Energy Convers. Manag.*, vol. 206, no. September 2019, 2020, doi: 10.1016/j.enconman.2020.112489.
- [10] J. J. Moore, M. G. Nored, R. S. Gernentz, and K. Brun, "Novel concepts for the compression of large volumes of carbon dioxide," in *Proceedings of the ASME Turbo Expo*, 2008, vol. 7, pp. 645–653, doi: 10.1115/GT2008-50924.

- [11] "p-h Diagram for Solid, Liquid and Gaseous CO2," *Imperial College London*, 2012. https://www.imperial.ac.uk/media/imperial-college/research-centres-and-groups/thermophysics/Chart-p-h-CO2.pdf.
- [12] Y. Mori, "HIGH TEMPERATURE HEAT EXCHANGERS.," in *Heat Transfer, Proceedings of the International Heat Transfer Conference*, 1986, vol. 1, pp. 259–268, doi: 10.1615/ihtc8.2470.
- [13] "Gas-to-Air Heat Exchangers Redecam Group." http://www.redecam.com/heat-exchangers/(accessed May 27, 2021).
- [14] B. I. Lee and M. G. Kesler, "A generalized thermodynamic correlation based on three-parameter corresponding states," *AIChE J.*, vol. 21, no. 3, pp. 510–527, 1975, doi: 10.1002/aic.690210313.
- [15] M. Mazzoccoli, B. Bosio, and E. Arato, "Analysis and comparison of equations-of-state with p-ρ-T experimental data for CO2 and CO2-mixture pipeline transport," *Energy Procedia*, vol. 23, pp. 274–283, 2012, doi: 10.1016/j.egypro.2012.06.052.
- [16] ASHRAE, Handbook of Fundamentals, SI Edition. 2017.
- [17] "Ammonia Refrigeration," *Creative Saftey Supply*, 2017. https://www.creativesafetysupply.com/articles/ammonia-refrigeration/ (accessed May 28, 2021).
- [18] S. M. Miner, "Ammonia as a refrigerant of choice," *ASHRAE J.*, vol. 32, no. 12, pp. 43–44, 1990.
- [19] J. M. Smith, H. C. Van Ness, M. M. Abbott, and M. T. Swihart, *Introduction to Chemical Engineering Thermodynamics*, 8th ed. McGraw-Hill Education, 2018.
- [20] "Pressure-Enthalpy Diagram for Ammonia US Customary System," *SLAY THE P.E. FREE RESOURCES*, 2020. https://www.slaythepe.com/uploads/7/2/3/9/72392189/ammonia\_p-h.pdf (accessed May 28, 2021).
- [21] Centre for Advanced Engineering, Energy Information Handbook. 2011.
- [22] R. Wennersten, Q. Sun, and H. Li, "The future potential for Carbon Capture and Storage in climate change mitigation An overview from perspectives of technology, economy and risk," *J. Clean. Prod.*, vol. 103, pp. 724–736, 2015, doi: 10.1016/j.jclepro.2014.09.023.
- [23] B. Tramošljika, P. Blecich, I. Bonefačić, and V. Glažar, "Advanced ultra-supercritical coal-fired power plant with post-combustion carbon capture: Analysis of electricity penalty and CO2 emission reduction," *Sustain.*, vol. 13, no. 2, pp. 1–20, 2021, doi: 10.3390/su13020801.
- [24] "Is carbon capture too expensive? Analysis IEA." https://www.iea.org/commentaries/is-carbon-capture-too-expensive (accessed May 30, 2021).

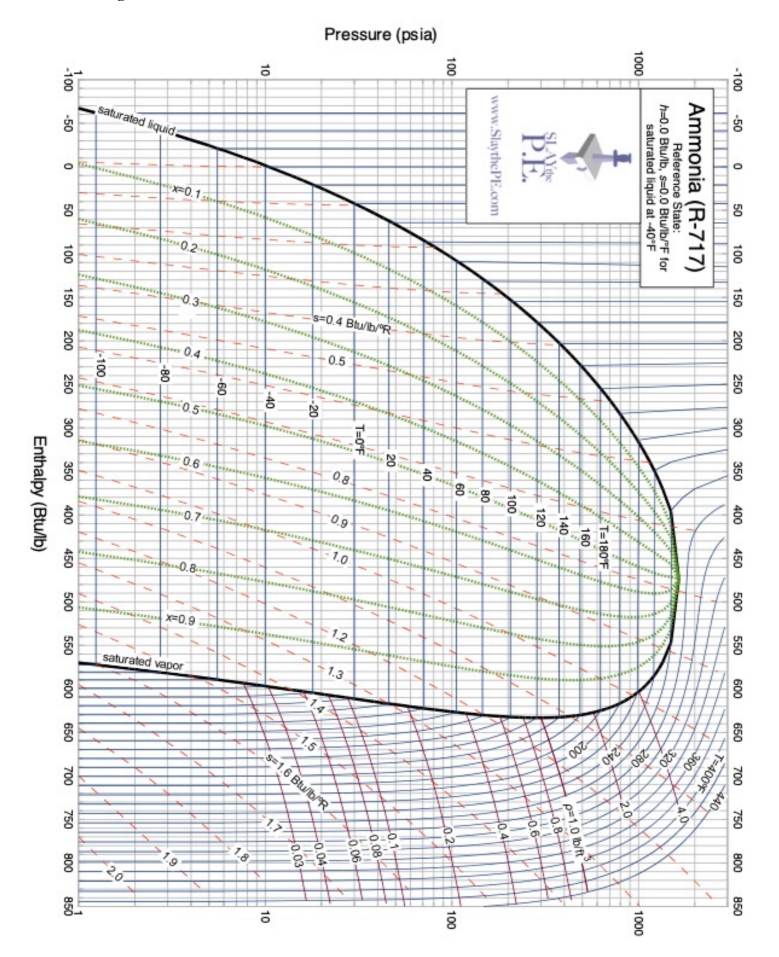
# 7 Appendix

# 7.1 Saturated/Superheated Table for Ammonia

		Satur	ated Proper		Super	heated Properties (t - t <sub>s</sub> )				
		Specific Enthalpy		Specific	Entropy	Abs. Tem (50		Abs. Tempera (100 K)	Abs. Temperature (100 K)	
Temperature - t <sub>s</sub> - (°C)	Pressure - p <sub>s</sub> - (bar)	Volume $-v_i - (m^3/kg)$	saturated liquid - h <sub>f</sub> -	saturated vapor - h <sub>g</sub> -	saturated liquid - s <sub>f</sub> -	saturated vapor - s <sub>g</sub> -	Specific Enthalpy - h -	Specific Entropy	Specific Enthalpy - h - (kJ/kg)	Specific Entropy
			(kJ/kg)	(kJ/kg)	(kJ/kgK)	(kJ/kgK)	(kJ/kg)	(kJ/kgK)	(KJ/Kg)	(kJ/kgK)
-50	0.4089	2.625	-44.4	1373.3	-0.194	6.159	1479.8	6.592	1585.9	6.948
-45	0.5454	2.005	-22.3	1381.6	-0.096	6.057	1489.3	6.486	1596.1	6.839
-40	0.7177	1.552	0	1390.0	0	5.962	1498.6	6.387	1606.3	6.736
-35	0.9322	1.216	22.3	1397.9	0.095	5.872	1507.9	6.293	1616.3	6.639
-30	1.196	0.9633	44.7	1405.6	0.188	5.785	1517.0	6.203	1626.3	6.547
-28	1.317	0.8809	53.6	1408.5	0.224	5.751	1520.7	6.169	1630.3	6.512
-26	1.447	0.8058	62.6	1411.4	0.261	5.718	1524.3	6.135	1634.2	6.477
-24	1.588	0.7389	71.7	1414.3	0.297	5.686	1527.9	6.103	1638.2	6.444
-22	1.740	0.6783	80.8	1417.3	0.333	5.655	1531.4	6.071	1642.2	6.411
-20	1.902	0.6237	89.8	1420.0	0.368	5.623	1534.8	6.039	1646.0	6.379
-18	2.077	0.5743	98.8	1422.7	0.404	5.593	1538.2	6.008	1650.0	6.347
-16	2.265	0.5296	107.9	1425.3	0.440	5.563	1541.7	5.978	1653.8	6.316
-14	2.465	0.4890	117.0	1427.9	0.475	5.533	1545.1	5.948	1657.7	6.286
-12	2.680	0.4521	126.2	1430.5	0.510	5.504	1548.5	5.919	1661.5	6.256
-10	2.908	0.4185	135.4	1433.0	0.544	5.475	1551.7	5.891	1665.3	6.227
-8	3.153	0.3879	144.5	1435.3	0.579	5.447	1554.9	5.863	1669.0	6.199
-6	3.413	0.3599	153.6	1437.6	0.613	5.419	1558.2	5.836	1672.8	6.171
-4	3.691	0.3344	162.8	1439.9	0.647	5.392	1561.4	5.808	1676.4	6.143
-2	3.983	0.3110	172.0	1442.2	0.681	5.365	1564.6	5.782	1680.1	6.116
0	4.295	0.2895	181.2	1444.4	0.715	5.340	1567.8	5.756	1683.9	6.090
2	4.625	0.2699	190.4	1446.5	0.749	5.314	1570.9	5.731	1687.5	6.065
4	4.975	0.2517	199.7	1448.5	0.782	5.288	1574.0	5.706	1691.2	6.040
6	5.346	0.2351	209.1	1450.6	0.816	5.263	1577.0	5.682	1694.9	6.015
8	5.736	0.2198	218.5	1452.5	0.849	5.238	1580.1	5.658	1698.4	5.991
10	6.149	0.2056	227.8	1454.3	0.881	5.213	1583.1	5.634	1702.2	5.967
12	6.585	0.1926	237.2	1456.1	0.914	5.189	1586.0	5.611	1705.7	5.943
14	7.045	0.1805	246.6	1457.8	0.947	5.165	1588.9	5.588	1709.1	5.920
16	7.529	0.1693	256.0	1459.5	0.979	5.141	1591.7	5.565	1712.5	5.898
18	8.035	0.1590	265.5	1461.1	1.012	5.118	1594.4	5.543	1715.9	5.876
20	8.570	0.1494	275.1	1462.6	1.044	5.095	1597.2	5.521	1719.3	5.854
22	9.134	0.1405	284.6	1463.9	1.076	5.072	1600.0	5.499	1722.8	5.832
24	9.722	0.1322	294.1	1465.2	1.108	5.049	1602.7	5.478	1726.3	5.811
26	10.34	0.1245	303.7	1466.5	1.140	5.027	1605.3	5.458	1729.6	5.790
28	10.99	0.1173	313.4	1467.8	1.172	5.005	1608.0	5.437	1732.7	5.770
30	11.67	0.1175	323.1	1468.9	1.204	4.984	1610.5	5.417	1735.9	5.750
32	12.37	0.1100	332.8	1469.9	1.235	4.962	1613.0	5.397	1739.3	5.731
34	13.11	0.0986	342.5	1470.8	1.267	4.940	1615.4	5.378	1742.6	5.711
36	13.89	0.0931	352.3	1471.8	1.298	4.919	1617.8	5.358	1745.7	5.692
38	14.70	0.0880	362.1	1472.6	1.329	4.898	1620.1	5.340	1748.7	5.674
40	15.54	0.0833	371.9	1472.0	1.360	4.877	1622.4	5.321	1751.9	5.655
42	16.42	0.0833	381.8	1473.8	1.391	4.856	1624.6	5.302	1755.0	5.637
44	17.34	0.0786	391.8	1473.8	1.422	4.835	1626.8	5.284	1758.0	5.619
46	18.30	0.0746	401.8	1474.5	1.422	4.833	1629.0	5.266	1761.0	5.602
70	10.30	0.0700	411.9	1474.3	1.433	4.793	1629.0	5.248	1764.0	5.584

	Saturated Properties								heated Properties (t - t <sub>s</sub> )	
Tamananatana	Pressure $-p_s$ - $(bar)$	D	D	Specific	Specific Enthalpy Specific Entropy		Abs. Temperature (50 K)		Abs. Temperature (100 K)	
Temperature - t <sub>s</sub> - (°C)		Volume - v <sub>i</sub> - (m³/kg)	saturated liquid - h <sub>f</sub> - (kJ/kg)	saturated vapor - h <sub>g</sub> - (kJ/kg)	saturated liquid - S <sub>f</sub> - (kJ/kgK)	saturated vapor - s <sub>g</sub> - (kJ/kgK)	Specific Enthalpy - h - (kJ/kg)	Specific Entropy - s - (kJ/kgK)	Specific Enthalpy - h - (kJ/kg)	Specific Entropy - s - (kJ/kgK)
50	20.33	0.0635	421.9	1474.7	1.515	4.773	1633.1	5.230	1766.8	5.567

# 7.2 P-H Diagram for Ammonia



## 7.3 MATLAB® Implementation

## 7.3.1 MATLAB® Code – ideal-gas state

```
clear all, close all, clc
global R A B C D;
R = 8.314462; % l*kPa / mol*K or J/mol*K
A = 5.457; B = 1.045 * 10^{-3}; C = 0; D = -1.157 * 10^{5};
P initial = 101.325 * 1; % kPa
P = 11000;
T1 = 25 + 273.15; % ambient temperature
Tref = -30 + 273.15; % refrigerated for storage (low pressure)
EperCO2 = 10.5*0.315; % MJ/kg
fprintf(' << Calculation assuming ideal-gas state >>\n\n')
fprintf(' Initial temperature %.2f K\n', T1)
fprintf(' Initial pressure
                                                                                       %.3f kPa\n', P_initial);
fprintf(' Final pressure
                                                                                      %.1f kPa\n', P final);
n = 3; % number of compressors in the chain
r_total = P_final / P_initial ; % total compression ratio
r = r total^{-}(1/n);
                                                                         % single compression ratio
 \begin{array}{lll} & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ 
fprintf(' Single compression ratio %.3f\n', r);
eff_comp = 0.65; % single compressor efficiency
fprintf(' Compressor efficiency %.f %% \n', 100*eff_comp);
P = zeros(n+1,1);
for i = 1:n+1
         P(i) = P_{initial} * r^{(i-1)};
end
fprintf(' Ambient temperature
                                                                                     %.2f K\n', T1);
fprintf(' Refrigeration temperature %.2f K\n', Tref);
W_{comp_total} = 0;
\overline{W} other = 0.970*44.01*1000;
fprintf('\n\n >> Compression to %d MPa\n', P final/1000);
for j = 1:n
         P1 = P(j);
         P2 = P(j+1);
         T2 isen = zeros(100,1);
         T2_{isen(1)} = T1 + 100;
         for i = 1:10000
                   T2 isen(i+1) = T1*(P2/P1)^(1/MCPS(T1,T2 isen(i)));
                   if abs( (T2 isen(i+1) - T2 isen(i)) / T2 isen(i) ) < 0.0001
                            MCPS(T1,T2 isen(i+1));
                            T2 is = T2_{isen(i+1)};
                            clear T2 isen
                             break
                   end
         end
         CpH = R * MCPH(T1,T2_is); % heat capacity
         Ws_comp_isen = CpH * (T2_is - T1); % minimum work of a compressor
Ws_comp = Ws_comp_isen / eff_comp; % actual required work of a compressor
         T2 real = zeros(100,1);
         T2 real(1) = T2 is;
         for i = 1:10000
                   T2 real(i+1) = T1 + Ws comp / (R*MCPH(T1,T2 real(i)));
                   if abs( (T2\_real(i+1) - T2\_real(i))/T2\_real) < 0.0001
                     T2 rl = T2 real(i+1);
```

```
clear T2 real
            break
        end
    end
    W comp total = W comp total + Ws comp;
    fprintf('\n - Compression step %d: ',j)
    fprintf('%.3f kPa to %.3f kPa', P1, P2);
end
fprintf('\n');
fprintf('\n
                 Isentropic work %.1f J/mol', n*Ws_comp_isen);
fprintf('\n Real compression work %.1f J/mol', W_comp_total);
fprintf('\n
                                    %.5f MJ/kg', W_comp_total/44.01/1000);
fprintf('\n
                        Highest T %.3f K\n', T2 rl);
if P final < 7000
    W ref = 0.1264*44.01*1000; % molar mass of CO2: 44.01 g/mol
    fprintf('\n >> Refrigeration\n');
    fprintf('\n
                                T hot
                                        %.3f K',T1);
                               T cold %.3f K', Tref);
    fprintf('\n
   fprintf('\n
                  Refrigeration work %.2f J/mol', W_ref);
   fprintf('\n
                                       %.5f MJ/kg', W ref/44.01/1000);
    fprintf('\n
                       Coefficient of')
   fprintf('\n
                        Performance %.2f\n', Tref/(T1-Tref))
else
   fprintf('\n');
    W ref = 0;
end
fprintf('\n >> Total work required\n');
fprintf('\n
                 Compression %.1f J/mol', W_comp_total);
if P final<7000</pre>
fprintf('\n
               Refrigeration %.1f J/mol',W ref);
end
fprintf('\n
                         Other %.1f J/mol', W_other);
fprintf('\n
                         Total %.1f J/mol', W other+W comp total+W ref);
                               %.4f MJ/kg',(\overline{W}_other+\overline{W}_comp_total+\overline{W}_ref)/44.01/1000);
fprintf('\n
fprintf('\n Emission intensity %.4f MJ/kg',EperCO2);
fprintf('\n\n >> Energy penalty\n');
fprintf('\n %.2f %%',(W_other+W_comp_total+W_ref)/44.01/1000/EperCO2*100);
fprintf('\n\n');
function MCPS = MCPS(T1, T2)
    global A B C D;
    MCPS = A + (B + (C + D/(T1^2*T2^2))*((T1+T2)/2))*(T2-T1)/log(T2/T1);
end
function MCPH = MCPH(T1,T2)
    global A B C D;
    MCPH = A + (B/2)*(T2+T1) + (C/3)*(T2^2 + T1^2 + T1*T2) + D/(T1*T2);
end
```

## 7.3.2 MATLAB® Code – Lee-Kesler equation of state

```
clear all, close all, clc
global R AA BB CC DD;
R = 8.314462; % l*kPa / mol*K or J/mol*K
AA = 5.457; BB = 1.045 * 10^{-3}; CC = 0; DD = -1.157 * 10^{5}; %Cp ig const. for CO2
P_initial = 101.325; % kPa
                        % kPa
P final = 11000;
T1 = 25 + 273.15; % ambient temperature
Tref = -30 + 273.15; % refrigerated for storage (low pressure)
EperCO2 = 10.5*0.315; % MJ/kg
fprintf(' << Calculation using Lee-Kelser EOS >>\n\n')
fprintf(' Initial temperature %.2f K\n', T1)
fprintf(' Initial pressure
                                        %.3f kPa\n', P_initial)
fprintf(' Final pressure
                                        %.2f kPa\n', P final);
n = 3;
r total = P final / P initial ;
r = r total^{(1/n)};
 \begin{array}{lll} & \text{fprintf(' Number of steps} & \text{%d } \\ & \text{n', n)} \\ & \text{fprintf(' Total compression ratio} & \text{%.3f } \\ & \text{n', r_total)} \\ \end{array} 
fprintf(' Single compression ratio %.3f \n', r)
eff_{comp} = 0.65;
fprintf(' Compressor efficiency %.f %% \n', 100*eff_comp);
P comp = zeros(n+1,1);
for i = 1:n+1
    P comp(i) = P initial * r^(i-1);
end
                                       %.2f K\n', T1);
fprintf(' Ambient temperature
fprintf(' Refrigeration temperature %.2f K\n', Tref);
W \text{ other} = 0.970*44.01*1000;
fprintf('\n\n >> Compression to %d MPa\n', P final/1000);
w = 0.22394; w0 = 0.3978; Tc = 304.1282; Pc = 7377.3;
T2(1) = 500;
HR = zeros(2,1);
SR = zeros(2,1);
a = 1;
for j = 1:100
    Tmat = [T1, T2(j)];
    Pmat = [P_comp(1), P_comp(2)];
    for i=1:2 % To calculate with two sets of T and P
        T = Tmat(i);
        P = Pmat(i);
        Tr = T/Tc;
        Pr = P/Pc;
        % Calculating for simple fluids
        b1 = 0.1181193;
        b2 = 0.265728;
        b3 = 0.154790;
        b4 = 0.030323;
        c1 = 0.0236744;
        c2 = 0.0186984;
        c3 = 0.0;
        c4 = 0.042724;
        d1 = 0.155488*10^{-4};
        d2 = 0.623689*10^{-4};
        beta = 0.65392;
        gamma = 0.060167;
```

```
B = b1 - b2/Tr - b3/Tr^2 - b4/Tr^3;
                  C = c1 - c2/Tr + c3/Tr^3;
                  D = d1 + d2/Tr;
                   % Numerically solving the EOS to obtain Vr (molar volume)
                  LK eos = Pr*Vr/Tr == 1 + (B/Vr) + (C/Vr^2) + (D/Vr^5) + (c4/(Tr^3*Vr^2))*(beta + C/Vr^5)
gamma/Vr^2) *exp(- gamma/Vr^2);
                 Vr0 = vpasolve(LK_eos, Vr);
                  Vr = Vr0;
                  % Calculating the compressibility for simple fluids
                  Z0 = Pr*Vr0/Tr;
                  z = z0;
                  % Calculating the residual properties for simple fluids
                  E = (c4/(2*Tr^3*gamma))*(beta+1-(beta+1+gamma/Vr^2)*exp(-gamma/Vr^2));
                  HR RTc 0 = \text{Tr} \cdot (Z-1-(b2 + 2*b3/\text{Tr} + 3*b4/\text{Tr}^2)/(\text{Tr}^*\text{Vr}) - (c2- 3*c3/\text{Tr}^2)/(2*\text{Tr}^*\text{Vr}^2) +
d2/(5*Tr*Vr^5)+3*E);
                  SR R 0 = log(Z) - ((b1 + b3/Tr<sup>2</sup> + 2*b4/Tr<sup>3</sup>)/Vr) - ((c1- (2*c3)/(Tr<sup>3</sup>))/(2*Vr<sup>2</sup>)) -
(d1/(5*Vr^{5})) + 2*E;
                  % Calculations for reference fluids
                  b1 = 0.2026579;
                  b2 = 0.331511;
                  b3 = 0.027655;
                  b4 = 0.203488;
                  c1 = 0.0313385;
                  c2 = 0.0503618;
                  c3 = 0.016901;
                  c4 = 0.041577;
                  d1 = 0.48736*10^{-4};
                  d2 = 0.0740336*10^{-4};
                  beta = 1.226;
                  gamma = 0.03754;
                  B = b1 - b2/Tr - b3/Tr^2 - b4/Tr^3;
                  C = c1 - c2/Tr + c3/Tr^3;
                  D = d1 + d2/Tr;
                  % Numerically solving the EOS to obtain Vr (molar volume)
                  svms Vr
                  LK eos = Pr*Vr/Tr == 1 + (B/Vr) + (C/Vr^2) + (D/Vr^5) + (c4/(Tr^3*Vr^2))*(beta + C/Vr^2)
gamma/Vr^2) *exp(-gamma/Vr^2);
                  Vr1 = vpasolve(LK eos, Vr, 10);
                  Vr = Vr1;
                  % Calculating the compressibility factor for reference fluids
                  Z1 = Pr*Vr1/Tr;
                  z = z1;
                  % Calculating the residual properties for reference fluids
                  E = (c4/(2*Tr^3*gamma))*(beta+1-(beta+1+gamma/Vr^2)*exp(-gamma/Vr^2));
                   \text{HR RTc 1} = \text{Tr}^*(\text{Z}-1-(\text{b2} + 2\text{*b3/Tr} + 3\text{*b4/Tr}^2)/(\text{Tr}^*\text{Vr}) - (\text{c2}-3\text{*c3/Tr}^2)/(2\text{*Tr}^*\text{Vr}^2) + (\text{c2}-3\text{*c3/Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2\text{*Tr}^2)/(2
d2/(5*Tr*Vr^5)+3*E);
                  SR R 1 = \log(Z) - ((b1 + b3/Tr^2 + 2*b4/Tr^3)/Vr) - ((c1-(2*c3)/(Tr^3))/(2*Vr^2)) -
(d1/(5*Vr^5)) + 2*E;
                   % Calculating the properties for the substance of interest
                  Z = Z0 + (w/w0) * (Z1-Z0);
                  Vr = Z*Tr/Pr;
                  HR(i) = R*Tc* (HR RTc 0 + (w/w0) * (HR RTc 1 - HR RTc 0));
                  SR(i) = R^* (SR_R_0 + (w/w0) * (SR_R_1 - SR_R_0));
         end
         delta_H = - HR(1) + HR(2) + R * MCPH(Tmat(1), Tmat(2)) * (Tmat(2)-Tmat(1));
         S tot(j) = delta S;
         fprintf('.')
```

```
if j > 1 \&\& S tot(j) * S tot(j-1) < 0
        a = a+1;
        if a > 5
            break
        end
    end
    if S_tot(j) > 0
        \overline{T}2(j+1) = T2(j) - 10^{(2-a)};
    elseif delta S < 0</pre>
        T2(j+1) = T2(j) + 10^{(2-a)};
end
W comp isen = n * delta H;
W_comp = W_comp_isen / eff_comp;
S tot2 = n * delta S;
fprintf('\n');
\text{fprintf($^{\prime}$}\backslash n
                       Change in S %.4f J/mol K', S_tot2);
fprintf('\n
                       Change in H %.1f J/mol \n', W_comp_isen);
fprintf('\n Isentropic work %.if 0/Mol', W_comp);
fprintf('\n Real compression work %.lf J/mol', W_comp);
fprintf('\n %.5f MJ/kg', W_comp/44.01/1000);
fprintf('\n
                         Highest T %.3f K\n', T2(\overline{j}));
if P final < 7000
    W ref = 0.1264*44.01*1000; % molar mass of CO2: 44.01 g/mol
    fprintf('\n >> Refrigeration\n');
    %.3f K',T1);
                                 T hot
    fprintf('\n
                                 T cold %.3f K', Tref);
    fprintf('\n
                  Refrigeration work
                                         %.2f J/mol', W_ref);
                                          %.5f MJ/kg', W ref/44.01/1000);
    fprintf('\n
                        Coefficient of')
    fprintf('\n
    fprintf('\n
                           Performance %.2f\n', Tref/(T1-Tref))
else
    W ref = 0;
    fprintf('\n');
end
fprintf('\n >> Total work required\n');
fprintf('\n
                  Compression %.1f J/mol', W_comp);
if P final<7000</pre>
fprintf('\n Refrigeration %.1f J/mol', W ref);
end
fprintf('\n
                          Other %.1f J/mol', W other);
fprintf('\n
                          Total %.1f J/mol', W_other+W_comp+W_ref);
fprintf('\n
                                 %.4f MJ/kg',(\overline{W}_other+\overline{W}_comp+\overline{W}_ref)/44.01/1000);
fprintf('\n Emission intensity %.4f MJ/kg',EperCO2);
fprintf('\n\n >> Energy penalty\n');
fprintf('\n\n');
function MCPS = MCPS(T1, T2)
    global AA BB CC DD;
    MCPS = AA + (BB + (CC + DD/(T1^2*T2^2))*((T1+T2)/2))*(T2-T1)/log(T2/T1);
end
function MCPH = MCPH(T1,T2)
    global AA BB CC DD;
    MCPH = AA + (BB/2)*(T2+T1) + (CC/3)*(T2^2 + T1^2 + T1*T2) + DD/(T1*T2);
end
```

## 7.4 MATLAB® Execution Results

## 7.4.1 Ideal-gas state, compression to 11 MPa, 2 stages

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.0 kPa
Number of compressors 2
Total compression ratio 108.562
Single compression ratio 10.419
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

## >> Compression to 11 MPa

- Compression step 1: 101.325 kPa to 1055.734 kPa - Compression step 2: 1055.734 kPa to 11000.000 kPa

Isentropic work 14926.9 J/mol Real compression work 22964.5 J/mol 0.52180 MJ/kg Highest T  $562.841~\mathrm{K}$ 

## >> Total work required

Compression 22964.5 J/mol Other 42689.7 J/mol Total 65654.2 J/mol 1.4918 MJ/kg Emission intensity 3.3075 MJ/kg

>> Energy penalty

45.10 %

## 7.4.2 Ideal-gas state, compression to 11 MPa, 3 stages

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.0 kPa
Number of compressors 3
Total compression ratio 108.562
Single compression ratio 4.770
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

## >> Compression to 11 MPa

Compression step 1: 101.325 kPa to 483.365 kPa
 Compression step 2: 483.365 kPa to 2305.866 kPa
 Compression step 3: 2305.866 kPa to 11000.000 kPa

Isentropic work 13757.6 J/mol
Real compression work 21165.6 J/mol
0.48093 MJ/kg
Highest T 466.986 K

## >> Total work required

Compression 21165.6 J/mol
Other 42689.7 J/mol
Total 63855.3 J/mol
1.4509 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

43.87 %

## 7.4.3 Ideal-gas state, compression to 11 MPa, 4 stages

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.0 kPa
Number of compressors 4
Total compression ratio 108.562
Single compression ratio 3.228
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

## >> Compression to 11 MPa

Compression step 1: 101.325 kPa to 327.066 kPa
Compression step 2: 327.066 kPa to 1055.734 kPa
Compression step 3: 1055.734 kPa to 3407.797 kPa
Compression step 4: 3407.797 kPa to 11000.000 kPa

Isentropic work 13200.5 J/mol Real compression work 20308.4 J/mol 0.46145 MJ/kg Highest T 422.394 K

## >> Total work required

Compression 20308.4 J/mol Other 42689.7 J/mol Total 62998.1 J/mol 1.4315 MJ/kg Emission intensity 3.3075 MJ/kg

>> Energy penalty

43.28 %

## 7.4.4 Ideal-gas state, compression to 11 MPa, 5 stages

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.0 kPa
Number of compressors 5
Total compression ratio 108.562
Single compression ratio 2.553
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

## >> Compression to 11 MPa

Compression step 1: 101.325 kPa to 258.733 kPa
Compression step 2: 258.733 kPa to 660.674 kPa
Compression step 3: 660.674 kPa to 1687.028 kPa
Compression step 4: 1687.028 kPa to 4307.819 kPa
Compression step 5: 4307.819 kPa to 11000.000 kPa

Real compression work 12874.1 J/mol Real compression work 19806.3 J/mol 0.45004 MJ/kg Highest T 396.557 K

## >> Total work required

Compression 19806.3 J/mol Other 42689.7 J/mol Total 62496.0 J/mol 1.4200 MJ/kg Emission intensity 3.3075 MJ/kg

>> Energy penalty

42.93 %

## Ideal-gas state, compression to 11 MPa, 6 stages

<< Calculation assuming ideal-gas state >>

298.15 K Initial temperature Initial pressure 101.325 kPa 11000.0 kPa Final pressure Number of compressors 108.562 Total compression ratio Single compression ratio 2.184 Compressor efficiency 65 % Ambient temperature 298.15 K Refrigeration temperature 243.15 K

## >> Compression to 11 MPa

- Compression step 1: 101.325 kPa to 221.307 kPa - Compression step 2: 221.307 kPa to 483.365 kPa - Compression step 3: 483.365 kPa to 1055.734 kPa - Compression step 4: 1055.734 kPa to 2305.866 kPa - Compression step 5: 2305.866 kPa to 5036.320 kPa - Compression step 6: 5036.320 kPa to 11000.000 kPa

Isentropic work 12659.6 J/mol Real compression work 19476.3 J/mol  $0.44254 \, \text{MJ/kg}$ 

Highest T 379.671 K

## >> Total work required

Compression 19476.3 J/mol Other 42689.7 J/mol Total 62166.0 J/mol 1.4125 MJ/kg Emission intensity 3.3075 MJ/kg

>> Energy penalty

42.71 %

## 7.4.6 Ideal-gas state, compression to 11 MPa, 1000 stages

<< Calculation assuming ideal-gas state >>

```
298.15 K
Initial temperature
Initial pressure
                          101.325 kPa
Final pressure
                          11000.0 kPa
Number of compressors
                          1000
Total compression ratio
                         108.562
Single compression ratio 1.005
Compressor efficiency
                          65 %
Ambient temperature
                          298.15 K
Refrigeration temperature 243.15 K
```

- >> Compression to 11 MPa
- Compression step 1: 101.325 kPa to 101.801 kPa Compression step 2: 101.801 kPa to 102.279 kPa

:

- Compression step 999: 10897.361 kPa to 10948.560 kPa
- Compression step 1000: 10948.560 kPa to 11000.000 kPa  $\,$

Isentropic work 11626.1 J/mol
Real compression work 17886.2 J/mol
0.40641 MJ/kg
Highest T 298.631 K

>> Total work required

Compression 17886.2 J/mol
Other 42689.7 J/mol
Total 60575.9 J/mol
1.3764 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.61 %

## 7.4.7 Ideal-gas state, compression to 2 MPa and refrigeration, 1 stage

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.0 kPa
Number of compressors 1
Total compression ratio 19.738
Single compression ratio 19.738
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

## >> Compression to 2 MPa

- Compression step 1: 101.325 kPa to 2000.000 kPa

Isentropic work 10145.0 J/mol Real compression work 15607.7 J/mol 0.35464 MJ/kg Highest T 648.786 K

## >> Refrigeration

T\_hot 298.150 K
T\_cold 243.150 K
Refrigeration work 5562.86 J/mol 0.12640 MJ/kg
Coefficient of Performance 4.42

## >> Total work required

Compression 15607.7 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 63860.2 J/mol
1.4510 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

43.87 %

#### 7.4.8 Ideal-gas state, compression to 2 MPa and refrigeration, 2 stages

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.0 kPa
Number of compressors 2
Total compression ratio 19.738
Single compression ratio 4.443
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

# >> Compression to 2 MPa

- Compression step 1: 101.325 kPa to 450.167 kPa - Compression step 2: 450.167 kPa to 2000.000 kPa

Isentropic work 8688.9 J/mol Real compression work 13367.5 J/mol 0.30374 MJ/kg Highest T 458.714 K

## >> Refrigeration

 $T_hot$  298.150 K  $T_cold$  243.150 K Refrigeration work 5562.86 J/mol 0.12640 MJ/kg Coefficient of Performance 4.42

## >> Total work required

Compression 13367.5 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 61620.1 J/mol
1.4001 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

42.33 %

## Ideal-gas state, compression to 2 MPa and refrigeration, 3 stages

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K Initial pressure 101.325 kPa 2000.0 kPa Final pressure Number of compressors Total compression ratio 19.738 Single compression ratio 2.703 Compressor efficiency 65 % Ambient temperature 298.15 K Refrigeration temperature 243.15 K

#### >> Compression to 2 MPa

- Compression step 1: 101.325 kPa to 273.834 kPa - Compression step 2: 273.834 kPa to 740.046 kPa - Compression step 3: 740.046 kPa to 2000.000 kPa

Isentropic work 8241.8 J/mol Real compression work 12679.7 J/mol  $0.28811 \, \text{MJ/kg}$ Highest T 402.752 K

#### >> Refrigeration

T hot 298.150 K T cold 243.150 K Refrigeration work 5562.86 J/mol  $0.12640 \, \text{MJ/kg}$ Coefficient of Performance 4.42

#### >> Total work required

Compression 12679.7 J/mol Refrigeration 5562.9 J/mol Other 42689.7 J/mol Total 60932.3 J/mol  $1.3845 \, \text{MJ/kg}$ 

Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.86 %

## 7.4.10 Ideal-gas state, compression to 2 MPa and refrigeration, 4 stages

<< Calculation assuming ideal-gas state >>

```
298.15 K
Initial temperature
Initial pressure
                          101.325 kPa
                          2000.0 kPa
Final pressure
Number of compressors
Total compression ratio
                         19.738
Single compression ratio 2.108
Compressor efficiency
                          65 %
Ambient temperature
                          298.15 K
Refrigeration temperature 243.15 K
```

#### >> Compression to 2 MPa

```
- Compression step 1: 101.325 kPa to 213.572 kPa
- Compression step 2: 213.572 kPa to 450.167 kPa
- Compression step 3: 450.167 kPa to 948.859 kPa
- Compression step 4: 948.859 kPa to 2000.000 kPa
```

Isentropic work 8024.5 J/mol Real compression work 12345.4 J/mol  $0.28051 \, \text{MJ/kg}$ 

Highest T 375.860 K

## >> Refrigeration

T hot 298.150 K T cold 243.150 K Refrigeration work 5562.86 J/mol  $0.12640 \, \text{MJ/kg}$ Coefficient of

Performance 4.42

#### >> Total work required

Compression 12345.4 J/mol Refrigeration 5562.9 J/mol Other 42689.7 J/mol Total 60597.9 J/mol 1.3769 MJ/kg

Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.63 %

## 7.4.11 Ideal-gas state, compression to 2 MPa and refrigeration, 5 stages

<< Calculation assuming ideal-gas state >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.0 kPa
Number of compressors 5
Total compression ratio 19.738
Single compression ratio 1.816
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

#### >> Compression to 2 MPa

Compression step 1: 101.325 kPa to 183.984 kPa
Compression step 2: 183.984 kPa to 334.073 kPa
Compression step 3: 334.073 kPa to 606.603 kPa
Compression step 4: 606.603 kPa to 1101.456 kPa
Compression step 5: 1101.456 kPa to 2000.000 kPa

Isentropic work 7895.9 J/mol Real compression work 12147.6 J/mol 0.27602 MJ/kg Highest T 360.018 K

#### >> Refrigeration

T\_hot 298.150 K 
T\_cold 243.150 K 
Refrigeration work 5562.86 J/mol 
0.12640 MJ/kg 
Coefficient of 
Performance 4.42

#### >> Total work required

Compression 12147.6 J/mol Refrigeration 5562.9 J/mol Other 42689.7 J/mol Total 60400.1 J/mol 1.3724 MJ/kg Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.49 %

## 7.4.12 Ideal-gas state, compression to 2 MPa and refrigeration, 1000 stages

<< Calculation assuming ideal-gas state >>

```
Initial temperature
                          298.15 K
Initial pressure
                          101.325 kPa
Final pressure
                         2000.0 kPa
                     1000
Number of compressors
Total compression ratio
                         19.738
Single compression ratio 1.003
Compressor efficiency
                         65 %
Ambient temperature
                          298.15 K
Refrigeration temperature 243.15 K
```

# >> Compression to 2 MPa

- Compression step 1: 101.325 kPa to 101.628 kPa - Compression step 2: 101.628 kPa to 101.931 kPa

:

- Compression step 999: 1988.105 kPa to 1994.044 kPa - Compression step 1000: 1994.044 kPa to 2000.000 kPa

Isentropic work 7396.2 J/mol
Real compression work 11378.8 J/mol
0.25855 MJ/kg
Highest T 298.456 K

>> Refrigeration

T\_hot 298.150 K
T\_cold 243.150 K
Refrigeration work 5562.86 J/mol 0.12640 MJ/kg
Coefficient of Performance 4.42

>> Total work required

Compression 11378.8 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 59631.4 J/mol
1.3550 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

40.97 %

# 7.4.13 Lee-Kesler equation of state, compression to 11 MPa, 2 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.00 kPa
Number of steps 2
Total compression ratio 108.562
Single compression ratio 10.419
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

# >> Compression to 11 MPa

Change in S -0.0002 J/mol K Change in H 14847.0 J/mol

Isentropic work 14847.0 J/mol Real compression work 22841.6 J/mol 0.51901 MJ/kg Highest T 477.813 K

## >> Total work required

Compression 22841.6 J/mol
Other 42689.7 J/mol
Total 65531.3 J/mol
1.4890 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

45.02 %

# 7.4.14 Lee-Kesler equation of state, compression to 11 MPa, 3 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.00 kPa
Number of steps 3
Total compression ratio 108.562
Single compression ratio 4.770
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

# >> Compression to 11 MPa

Change in S -0.0003 J/mol K Change in H 13683.2 J/mol

Isentropic work 13683.2 J/mol Real compression work 21051.1 J/mol 0.47832 MJ/kg Highest T 411.903 K

# >> Total work required

Compression 21051.1 J/mol
Other 42689.7 J/mol
Total 63740.8 J/mol
1.4483 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

43.79 %

#### 7.4.15 Lee-Kesler equation of state, compression to 11 MPa, 4 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.00 kPa
Number of steps 4
Total compression ratio 108.562
Single compression ratio 3.228
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

## >> Compression to 11 MPa

Change in S -0.0004 J/mol K Change in H 13129.6 J/mol

Isentropic work 13129.6 J/mol Real compression work 20199.4 J/mol 0.45897 MJ/kg Highest T 381.464 K

# >> Total work required

Compression 20199.4 J/mol Other 42689.7 J/mol Total 62889.1 J/mol  $1.4290~\mathrm{MJ/kg}$  Emission intensity 3.3075 MJ/kg

>> Energy penalty

43.20 %

# 7.4.16 Lee-Kesler equation of state, compression to 11 MPa, 5 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.00 kPa
Number of steps 4
Total compression ratio 108.562
Single compression ratio 3.228
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

# >> Compression to 11 MPa

Change in S -0.0004 J/mol K Change in H 13129.6 J/mol

Isentropic work 13129.6 J/mol Real compression work 20199.4 J/mol 0.45897 MJ/kg Highest T 381.464 K

## >> Total work required

Compression 20199.4 J/mol
Other 42689.7 J/mol
Total 62889.1 J/mol
1.4290 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

43.20 %

## 7.4.17 Lee-Kesler equation of state, compression to 11 MPa, 6 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.00 kPa
Number of steps 6
Total compression ratio 108.562
Single compression ratio 2.184
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

#### >> Compression to 11 MPa

......

Change in S -0.0002 J/mol K Change in H 12592.6 J/mol

Isentropic work 12592.6 J/mol Real compression work 19373.2 J/mol 0.44020 MJ/kg Highest T 352.482 K

# >> Total work required

Compression 19373.2 J/mol Other 42689.7 J/mol Total 62062.9 J/mol 1.4102 MJ/kg Emission intensity 3.3075 MJ/kg

>> Energy penalty

42.64 %

# 7.4.18 Lee-Kesler equation of state, compression to 11 MPa, 1000 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 11000.00 kPa
Number of steps 1000
Total compression ratio 108.562
Single compression ratio 1.005
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

#### >> Compression to 11 MPa

Change in S -0.0120 J/mol K Change in H 11562.5 J/mol

Isentropic work 11562.5 J/mol
Real compression work 17788.4 J/mol
0.40419 MJ/kg
Highest T 298.464 K

# >> Total work required

Compression 17788.4 J/mol Other 42689.7 J/mol Total 60478.1 J/mol 1.3742 MJ/kg Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.55 %

## 7.4.19 Lee-Kesler equation of state, compression to 2 MPa and refrigeration, 1 stage

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.00 kPa
Number of steps 1
Total compression ratio 19.738
Single compression ratio 19.738
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

# >> Compression to 2 MPa

Change in S 0.0000 J/mol K Change in H 10095.0 J/mol

Isentropic work 10095.0 J/mol Real compression work 15530.8 J/mol 0.35289 MJ/kg Highest T 537.474 K

>> Refrigeration

T\_hot 298.150 K 
T\_cold 243.150 K 
Refrigeration work 5562.86 J/mol 
0.12640 MJ/kg 
Coefficient of 
Performance 4.42

>> Total work required

Compression 15530.8 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 63783.3 J/mol
1.4493 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

43.82 %

## 7.4.20 Lee-Kesler equation of state, compression to 2 MPa and refrigeration, 2 stages

<< Calculation using Lee-Kelser EOS >>

```
Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.00 kPa
Number of steps 2
Total compression ratio 19.738
Single compression ratio 4.443
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K
```

#### >> Compression to 2 MPa

Change in S -0.0002 J/mol K Change in H 8641.7 J/mol

Isentropic work 8641.7 J/mol
Real compression work 13294.9 J/mol
0.30209 MJ/kg
Highest T 406.242 K

>> Refrigeration

T\_hot 298.150 K 
T\_cold 243.150 K 
Refrigeration work 5562.86 J/mol 
0.12640 MJ/kg 
Coefficient of 
Performance 4.42

>> Total work required

Compression 13294.9 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 61547.5 J/mol
1.3985 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

42.28 %

## 7.4.21 Lee-Kesler equation of state, compression to 2 MPa and refrigeration, 3 stages

<< Calculation using Lee-Kelser EOS >>

```
Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.00 kPa
Number of steps 3
Total compression ratio 19.738
Single compression ratio 2.703
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K
```

#### >> Compression to 2 MPa

Change in S -0.0003 J/mol K Change in H 8197.8 J/mol

Isentropic work 8197.8 J/mol
Real compression work 12612.0 J/mol
0.28657 MJ/kg
Highest T 368.113 K

#### >> Refrigeration

T\_hot 298.150 K 
T\_cold 243.150 K 
Refrigeration work 5562.86 J/mol 
0.12640 MJ/kg 
Coefficient of 
Performance 4.42

#### >> Total work required

Compression 12612.0 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 60864.5 J/mol
1.3830 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.81 %

## 7.4.22 Lee-Kesler equation of state, compression to 2 MPa and refrigeration, 4 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.00 kPa
Number of steps 4
Total compression ratio 19.738
Single compression ratio 2.108
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

# >> Compression to 2 MPa

Change in S -0.0004 J/mol K Change in H 7982.0 J/mol

Isentropic work 7982.0 J/mol Real compression work 12280.0 J/mol 0.27903 MJ/kg Highest T  $349.908 \ \mathrm{K}$ 

Highest 1 349.900

# >> Refrigeration

T\_hot 298.150 K 
T\_cold 243.150 K 
Refrigeration work 5562.86 J/mol 
0.12640 MJ/kg 
Coefficient of 
Performance 4.42

#### >> Total work required

Compression 12280.0 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 60532.6 J/mol
1.3754 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.59 %

## 7.4.23 Lee-Kesler equation of state, compression to 2 MPa and refrigeration, 5 stages

<< Calculation using Lee-Kelser EOS >>

```
Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.00 kPa
Number of steps 5
Total compression ratio 19.738
Single compression ratio 1.816
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K
```

#### >> Compression to 2 MPa

Change in S -0.0002 J/mol K Change in H 7854.4 J/mol

Isentropic work 7854.4 J/mol Real compression work 12083.7 J/mol 0.27457 MJ/kg Highest T 339.234 K

## >> Refrigeration

T\_hot 298.150 K 
T\_cold 243.150 K 
Refrigeration work 5562.86 J/mol 
0.12640 MJ/kg 
Coefficient of 
Performance 4.42

#### >> Total work required

Compression 12083.7 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 60336.3 J/mol
1.3710 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

41.45 %

## 7.4.24 Lee-Kesler equation of state, compression to 2 MPa and refrigeration, 1000 stages

<< Calculation using Lee-Kelser EOS >>

Initial temperature 298.15 K
Initial pressure 101.325 kPa
Final pressure 2000.00 kPa
Number of steps 1000
Total compression ratio 19.738
Single compression ratio 1.003
Compressor efficiency 65 %
Ambient temperature 298.15 K
Refrigeration temperature 243.15 K

## >> Compression to 2 MPa

Change in S -0.1064 J/mol K Change in H 7326.4 J/mol

Isentropic work 7326.4 J/mol Real compression work 11271.4 J/mol 0.25611 MJ/kg Highest T 298.349 K

## >> Refrigeration

T\_hot 298.150 KT\_cold 243.150 KRefrigeration work 5562.86 J/mol 0.12640 MJ/kgCoefficient of
Performance 4.42

#### >> Total work required

Compression 11271.4 J/mol
Refrigeration 5562.9 J/mol
Other 42689.7 J/mol
Total 59523.9 J/mol
1.3525 MJ/kg
Emission intensity 3.3075 MJ/kg

>> Energy penalty

40.89 %