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DEPARTMENT OF CHEMISTRY & LIFE SCIENCE

CH365 2024-2025 TEXT: Smith, Van Ness, Abbott & Swihart

WRITTEN PARTIAL REVIEW II SCOPE: Lessons 10-20 15 October 2024, A-Hour TIME: 55 Minutes

References Permitted: Open notes, book, internet, Mathematica, and Excel.

INSTRUCTIONS

- 1. Do not mark this exam or open it until "begin work" is given.
- 2. You have 55 minutes to complete the exam.
- 3. Solve the problems in the space provided. Show all work to receive full credit.
- 4. There are 3 problems on 3 pages in this exam (not including the cover page). Write your name on the top of each sheet. Use the back of the pages for extra workspace as needed.
- 5. Save Mathematica files on your desktop and re-save frequently.
- 6. When finished, upload all Mathematica files to Canvas.

(TOTAL WEIGHT: 200 POINTS)

DO NOT WRITE IN THIS SPACE

PROBLEM	VALUE	CUT
A	50	
В	70	
С	80	
TOTAL CUT		
TOTAL GRADE	200	

Problem: Weight: 50

Methanol (CH₄O(g)) is fed to a combustion reactor with 100% excess air.

- (a) Balance the reaction on a basis of one mole of methanol in the feed.
- (b) Calculate the heat of reaction at 25 °C and 1.0 bar in J/mol on a basis of one mole of methanol in the feed.
- (c) Calculate the standard gas-phase isothermal heat of reaction at 300 °C and 1.0 bar for one mole of methanol in 100% excess air.

The following table contains ideal gas heat capacity polynomial coefficients and standard heats of formation at 298 K from Appendix C:

species	a	b×10 ³	c×10 ⁶	d×10 ⁻⁵	$\Delta H_{f,298}^{0}$, J/mol
CH ₄ O	2.211	12.216	-3.450	0.000	-200,660
O_2	3.639	0.506	0.000	-0.227	0
N_2	3.280	0.593	0.000	0.040	0
CO_2	5.457	1.045	0.000	-1.157	-393,509
H_2O	3.470	1.450	0.000	0.121	-241,818

Solution, part (a):

Balanced reaction in 100% excess air:

$$\begin{split} CH_4O(g) + 3 \ O_2(g) + 11.29 \ N_2(g) \\ & \to \ CO_2(g) + 2 \ H_2O(g) + 1.50 \ O_2(g) + 11.29 \ N_2(g) \end{split}$$

Solution of parts (b) and (c) in Mathematica are shown on the following page.

Cadet:

Solution, part (b):

```
(*List of stoichiometric coefficients*)
(*methanol, oxygen, nitrogen, carbon dioxide, water, oxygen, nitrogen*)
v = \{-1, -3, -11.29, 1, 2, 1.5, 11.29\};
(*List of standard Heats of Formation*)
\Delta Hf298 = \{-200660, 0, 0, -393509, -241818, 0, 0\};
(*Standard Heat of Reaction at 298 K*)
\Delta H298 = Plus @@ (v * \Delta Hf298) (*J/mol*)
-676485.
         ANS, J per mole of methanol
```

Solution, part (c):

```
(*Heat Capacity polynomial coefficients, Given, from Table C.1:*)
a = \{ 2.211, 3.639, 3.280, 5.457, 3.470, 3.639, 3.280 \};
b = \{12.216, 0.506, 0.593, 1.045, 1.450, 0.506, 0.593\} * 10^{-3};
c = \{-3.450, 0.000, 0.000, 0.000, 0.000, 0.000, 0.000\} * 10^{-6};
d = \{ 0.000, -0.227, 0.040, -1.157, 0.121, -0.227, 0.040 \} * 10^5;
\Delta a = Plus @@ (v * a);
\Delta b = Plus @@ (v * b);
\Delta c = Plus @@ (v * c);
\Delta d = Plus @@ (v * d);
(*Temperatures, K*)
T = 300 + 273.15;
T0 = 25 + 273.15;
(*Gas constant*)
R = 8.314; (*\frac{1}{mol+k}*)
(*Equation 4.21 for MDCPH*)
MDCPH = \Delta a + \frac{\Delta b}{2} \star (T + T\theta) + \frac{\Delta c}{3} \star (T^2 + T\theta^2 + T \star T\theta) + \frac{\Delta d}{T + T\theta}
1.13391
(*Equation 4.22 for corrected heat of reaction*)
(*Gives molar heat of reaction in J/mol*)
\Delta H573 = \Delta H298 + MDCPH * R * (T - T0)
-673892.5
```

ANS, J per mole of methanol

An engineering plan for a vinyl chloride plant includes a pipeline carrying ethylene (C₂H₄) at a volumetric flow rate of 25.750 MMSCFD (million standard cubic feet per day). Standard conditions are 519.67°R and 14.696 psia. At these conditions, the molar volume of C₂H₄ is 377.27 ft³/lbmol.

- (a) Calculate the molar flow rate of C₂H₄ in lbmol/sec.
- (b) Use the van der Waals (vdW) equation of state (equations 3.41, 3.44, and 3.45), constants in Table 3.1, and gas constant R = 10.73 ft³·psia/(lbmol·R) to determine the isothermal reversible work produced by increasing the molar volume of C_2H_4 by 5%. Report the work in hp.

From Appendix B, the critical temperature, pressure, and acentric factor for ethylene are $T_C = 508.14$ Rankine, and $P_C = 730.99$ psia, and $\omega = 0.087$.

Solution, part (a):

$$\frac{25,750,000 \,\text{ft}^3}{\text{day}} \cdot \frac{1 \,\text{day}}{24 \,\text{h}} \cdot \frac{1 \,\text{h}}{3600 \,\text{s}} \cdot \frac{11 \,\text{bmol}}{377.27 \,\text{ft}^3} = \frac{0.78997}{\text{MNS}} \frac{\text{lbmol}}{\text{NS}}$$

Solution, part (b):

```
T = 519.67; (*Rankine, given*)
R = 10.73; (*\frac{ft^3*psia}{lbmol*R}, given*)
\omega = 0.087; (*App. B, p.665*)
Tc = 508.14; (*Rankine, given*)
Pc = 730.99; (*psia, given*)
Tr1 = T / Tc; (*reduced temperature*)
(*Table 3.1, p.100*)
\sigma = 0;
€ = 0;
\Omega = 1/8;
\Phi = 27 / 64;
\alpha = 1;
(*Equations 3.41, 3.44, and 3.45*)
b = \Omega * \frac{R * Tc}{Pc};
a = \Phi * \frac{\alpha * R^2 * Tc^2}{Pc};
P[V_{-}] = \frac{R * T}{V - b} - \frac{a}{(V + \epsilon * b) * (V + \sigma * b)};
W = - P[V] dV (*eq 1.4, units <math>\frac{psia*ft^3}{1bmol}*)
-270.5491
```

$$W = -\int_{377.27}^{1.05*377.27} P[V] \, dV(*eq 1.4, units \frac{psia*ft^3}{1bmol}*)$$

$$-270.5491$$

$$\frac{-270.5491 \, psia*ft^3}{1bmol} * \frac{\frac{1bf}{in^2}}{psia} * (\frac{12 \, in}{ft})^2 * \frac{0.78997 \, lbmol}{s} * \frac{1.34102 \, hp}{737.562 \, \frac{ft*lbf}{s}}$$

$$\frac{-55.95719 \, hp}{ANS}$$

Problem: Weight: 80

0.200 kmol of an ideal gas in a closed system is changed from an initial state of 1.5 m³, 5.0 bar and 451.0 K to a final state of 5.0 m³, 1.5 bar, and 451.0 K by a two-step process consisting of an adiabatic expansion followed by heating at constant pressure.

Cadet:_

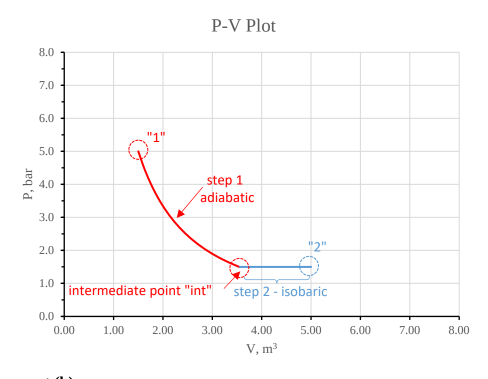
- (a) Sketch the process path on the PV axes below.
- (b) Calculate Q, W, ΔU and ΔH for the overall process in units of kJ.
- (c) Calculate the intermediate temperature after step 1.

$$R=8.314 \ J/(mol\cdot K)=0.08314 \ (bar\cdot m^3)/(kmol\cdot K)$$

$$C_P=(7/2)\cdot R, \ and$$

$$C_V=(5/2)\cdot R$$

Solution, part (a):



Solution, part (b):

The overall process is isothermal (T₂=T₁), so $\Delta \underline{\underline{H}_{Overall} = \Delta U_{Overall} = 0}_{ANS}$ ANS

Equation 3.23c gives the P-V relation for an adiabatic process. Let the intermediate state be denoted with subscript "int" with $P_{int} = P_2$ since the second step is at constant pressure.

Also
$$\gamma = C_P/C_V = (7R/2)/(5R/2) = (7/5) = 1.4$$
 from equation 3.24.

Solve for intermediate volume, V_{int}, using equation 3.23c from lesson 11:

$$P_{int}V_{int}^{\gamma} = P_{1}V_{1}^{\gamma} \Rightarrow V_{int} = \sqrt[\gamma]{\frac{P_{1}}{P_{int}}V_{1}^{\gamma}} = \sqrt[4]{\frac{5.0\,bar}{1.5\,bar} \left(1.5\,m^{3}\right)^{1.4}} = 3.544672\,m^{3}$$

Once V_{int} is known, the adiabatic work for step 1 is given by equation 3.26:

$$W_{Step1} = \frac{P_{int} \cdot V_{int} - P_1 \cdot V_1}{\gamma - 1} = \frac{1.5 \, bar \cdot 3.5446721 m^3 - 5.0 \, bar \cdot 1.5 \, m^3}{1.4 - 1} = -5.457479 \, bar \cdot m^3$$

Step 2 is at constant pressure, so work -P Δ V:

$$W_{Step 2} = -P \cdot \Delta V = -P \cdot (V_2 - V_{int}) = -1.5 \cdot (5.0 \,\text{m}^3 - 3.544672 \,\text{m}^3) = -2.18299 \,\text{bar} \cdot \text{m}^3.$$

The total overall work is the sum of the work of each step:

$$W_{Overall} = W_{Step1} + W_{Step2} = -5.45748 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 = -7.64047 \, bar \cdot m^3 + \left(-2.18299\right) bar \cdot m^3 + \left(-2.18299\right$$

Now convert the units to kJ:

Finally, for Q:

$$\Delta U_{\text{Overall}} = Q_{\text{Overall}} + W_{\text{Overall}} = 0 \implies Q_{\text{Overall}} = -W_{\text{Overall}} = 7\underline{\underline{64.047}}\,\text{kJ}$$
ANS

Solution, part (c):

$$PV = nRT \Rightarrow T_{int} = \frac{P_{int} \cdot V_{int}}{n \cdot R} = \frac{1.5 \text{ bar} \cdot 3.544672 \text{ m}^3}{0.200 \text{ kmol} \cdot \left(0.08314 \frac{\text{bar} \cdot \text{m}^3}{\text{kmol} \cdot \text{K}}\right)} = \frac{319.8 \text{ K}}{\text{ANS}}$$

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(TOTAL WEIGHT: 200 POINTS)

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PROBLEM	VALUE	CUT
A	50	
В	70	
С	80	
TOTAL CUT		
TOTAL GRADE	200	

Ethanol (C₂H₆O(g)) is fed to a combustion reactor with 100% excess air.

- (a) Balance the reaction on a basis of one mole of ethanol in the feed.
- (b) Calculate the heat of reaction at 25 °C and 1.0 bar in J/mol on a basis of one mole of ethanol in the feed.
- (c) Calculate the standard gas-phase isothermal heat of reaction at $30\underline{0}$ °C and 1.0 bar for one mole of ethanol in 100% excess air.

The following table contains ideal gas heat capacity polynomial coefficients and standard heats of formation at 298 K from Appendix C:

species	a	b×10 ³	c×10 ⁶	d×10 ⁻⁵	$\Delta H_{\rm f,298}^{0}$, J/mol
C ₂ H ₆ O	3.518	20.001	-6.002	0.000	-235,100
O_2	3.639	0.506	0.000	-0.227	0
N_2	3.280	0.593	0.000	0.040	0
CO_2	5.457	1.045	0.000	-1.157	-393,509
H_2O	3.470	1.450	0.000	0.121	-241,818

Solution, part (a):

Balanced reaction in 100% excess air:

$$\begin{split} C_2H_6O(g) + 6 \ O_2(g) + 22.57 \ N_2(g) \\ & \to \ 2 \ CO_2 \ (g) + 3 \ H_2O(g) + 3 \ O_2(g) + 22.57 \ N_2(g) \end{split}$$

Solution of parts (b) and (c) in Mathematica are shown on the following page.

Cadet:

Solution, part (b):

```
(*List of stoichiometric coefficients*)
(*methanol, oxygen, nitrogen, carbon dioxide, water, oxygen, nitrogen*)
v = {-1, -6, -22.57, 2, 3, 3, 22.57};
(*List of standard Heats of Formation*)
ΔHf298 = {-235100, 0, 0, -393509, -241818, 0, 0};
(*Standard Heat of Reaction at 298 K*)
ΔH298 = Plus @@ (v * ΔHf298) (*J/mol*)
-1.277372 × 10<sup>6</sup>
ANS, J per mole of ethanol
```

Solution, part (c):

```
(*Heat Capacity polynomial coefficients, Given, from Table C.1:*)
a = { 3.518, 3.639, 3.280, 5.457, 3.470, 3.639, 3.280};
b = \{20.001, 0.506, 0.593, 1.045, 1.450, 0.506, 0.593\} * 10^{-3};
c = \{-6.002, 0.000, 0.000, 0.000, 0.000, 0.000, 0.000\} * 10^{-6};
d = \{ 0.000, -0.227, 0.040, -1.157, 0.121, -0.227, 0.040 \} * 10^5;
\Delta a = Plus @@ (v * a);
\Delta b = Plus @@ (v * b);
\Delta c = Plus @@ (v * c);
\Delta d = Plus @@ (v * d);
(∗Temperatures, K∗)
T = 300 + 273.15;
T0 = 25 + 273.15;
(*Gas constant*)
R = 8.314; (*\frac{1}{mol_{+}K}*)
(*Equation 4.21 for MDCPH*)
MDCPH = \Delta a + \frac{\Delta b}{2} * (T + T0) + \frac{\Delta c}{3} * (T^2 + T0^2 + T * T0) + \frac{\Delta d}{T + T0}
0.7535926
(*Equation 4.22 for corrected heat of reaction*)
(*Gives molar heat of reaction in J/mol*)
\Delta H573 = \Delta H298 + MDCPH * R * (T - T0)
-1.275649 \times 10^6
            ANS, J per mole of ethanol
```

An engineering plan for a vinyl chloride plant includes a pipeline carrying ethylene (C₂H₄) at a volumetric flow rate of 7.175 MMSCFD (million standard cubic feet per day). Standard conditions are 519.67°R and 14.696 psia. At these conditions, the molar volume of C₂H₄ is 376.68 ft³/lbmol.

- (a) Calculate the molar flow rate of C₂H₄ in lbmol/sec.
- (b) Use the Peng-Robinson (PR) equation of state (equations 3.41, 3.44, and 3.45), constants in Table 3.1, and gas constant R = 10.73 ft³·psia/(lbmol·R) to determine the isothermal reversible compression work required to reduce the molar volume of C_2H_4 by 10%. Report the work in hp.

From Appendix B, the critical temperature, pressure, and acentric factor for ethylene are $T_C = 508.14$ Rankine, and $P_C = 730.99$ psia, and $\omega = 0.087$.

Solution, part (a):

$$\frac{17,571,000 \, \text{ft}^3}{\text{day}} \cdot \frac{1 \, \text{day}}{24 \, \text{h}} \cdot \frac{1 \, \text{h}}{3600 \, \text{s}} \cdot \frac{11 \text{bmol}}{376.68 \, \text{ft}^3} = \underbrace{0.53990 \, \frac{\text{lbmol}}{\text{s}}}_{\text{ANS}}$$

Solution, part (b):

```
T = 519.67; (*Rankine, given*)
 R = 10.73; (*\frac{ft^3*psia}{lbmol*R}, given*)
 \omega = 0.087; (*App. B, p.665*)
 Tc = 508.14; (*Rankine, given*)
 Pc = 730.99; (*psia, given*)
 Tr1 = T / Tc; (*reduced temperature*)
  (*Table 3.1, p.100*)
 \sigma = 1;
 \epsilon = 0;
 \Omega = 0.07780;
 \Phi = 0.45724;
 \alpha = (1 + (0.37464 + 1.54226 * \omega - 0.26992 * \omega^{2}) * (1 - \sqrt{Tr1}));
 (*Equations 3.41, 3.44, and 3.45*)
 b = \Omega * \frac{R * Tc}{Pc};
 a = \Psi * \frac{\alpha * R^2 * Tc^2}{Pc};
 P = \frac{R * T}{V - b} - \frac{a}{(V + \epsilon * b) * (V + \alpha + b)};
```

$$W = -\int_{376.68}^{6.9 + 376.68} P \, dV(\star eq 1.4, units \frac{psia + ft^3}{1bmol} \star)$$

$$583.0076$$

$$\frac{583.0076 \, psia \star ft^3}{1bmol} \star \frac{\frac{1bf}{in^2}}{psia} \star \left(\frac{12 \, in}{ft}\right)^2 \star \frac{0.53990 \, lbmol}{s} \star \frac{1.34102 \, hp}{737.562 \, \frac{ft + 1bf}{s}}$$

$$82.4113 \, hp$$

$$ANS$$

Problem: Weight: 80

0.500 kmol of an ideal gas in a closed system is changed from an initial state of 7.0 m³, 3.0 bar and 505.2 K to a final state of 3.0 m³, 7.0 bar, and 505.2 K by a two-step process consisting of an adiabatic compression followed by cooling at constant pressure.

Cadet:_

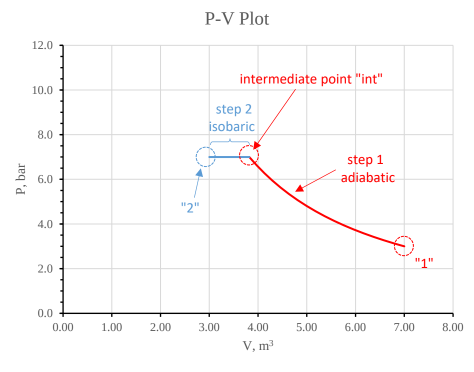
- (a) Sketch the process path on the PV axes below.
- (b) Calculate Q, W, ΔU and ΔH for the overall process in units of kJ.
- (c) Calculate the intermediate temperature after step 1.

$$R=8.314 \text{ J/(mol}\cdot\text{K)}=0.08314 \text{ (bar}\cdot\text{m}^3)/(\text{kmol}\cdot\text{K)}$$

$$C_P=(7/2)\cdot\text{R, and}$$

$$C_V=(5/2)\cdot\text{R}$$

Solution, part (a):



Solution, part (b):

The overall process is isothermal (T2=T1), so $\Delta H_{Overall} = \Delta U_{Overall} = 0 \text{ kJ}$ ANS

Equation 3.23c gives the P-V relation for an adiabatic process. Let the intermediate state be denoted with subscript "int" with $P_{int} = P_2$ since the second step is at constant pressure.

Also
$$\gamma = C_P/C_V = (7R/2)/(5R/2) = (7/5) = 1.4$$
 from equation 3.24.

Cadet:_

Solve for intermediate volume, V_{int}, using equation 3.23c from lesson 11:

$$P_{int}V_{int}^{\gamma} = P_{1}V_{1}^{\gamma} \Rightarrow V_{int} = \sqrt[\gamma]{\frac{P_{1}}{P_{int}}V_{1}^{\gamma}} = \sqrt[1.4]{\frac{3.0\,bar}{7.0\,bar}(7.0\,m^{3})^{1.4}} = 3.821708\,m^{3}$$

Once V_{int} is known, the adiabatic work for step 1 is given by equation 3.26:

$$W_{Step1} = \frac{P_{int} \cdot V_{int} - P_1 \cdot V_1}{\gamma - 1} = \frac{7.0 \, bar \cdot 3.8217081 \, m^3 - 7.0 \, bar \cdot 3.0 \, m^3}{1.4 - 1} = +14.37989 \, bar \cdot m^3$$

Step 2 is at constant pressure, so work -P Δ V:

$$W_{Step 2} = -P \cdot \Delta V = -P \cdot (V_2 - V_{int}) = -7.0 \cdot (3.0 \text{ m}^3 - 3.821708 \text{ m}^3) = +5.751956 \text{ bar} \cdot \text{m}^3.$$

The total overall work is the sum of the work of each step:

$$W_{Overall} = W_{Step1} + W_{Step2} = 14.37989 \, bar \cdot m^3 + 5.751956 \, bar \cdot m^3 = 20.13184 \, bar \cdot m^3$$

Now convert the units to kJ:

$$W_{\text{Overall}} = 20.1318 \,\text{bar} \cdot \text{m}^3 \left(\frac{1 \,\text{J}}{10^{-5} \,\text{bar} \cdot \text{m}^3} \right) \cdot \left(\frac{1 \,\text{kJ}}{1000 \,\text{J}} \right) = \frac{2013.18 \,\text{kJ}}{\text{MNS}}$$

Finally, for Q:

$$\Delta U_{\text{Overall}} = Q_{\text{Overall}} + W_{\text{Overall}} = 0 \implies Q_{\text{Overall}} = -W_{\text{Overall}} = -2013.18 \text{ kJ}$$
ANS

Solution, part (c):

$$PV = nRT \Rightarrow T_{int} = \frac{P_{int} \cdot V_{int}}{n \cdot R} = \frac{7.0 \text{ bar} \cdot 3.821708 \text{ m}^3}{0.500 \text{ kmol} \cdot \left(0.08314 \frac{\text{bar} \cdot \text{m}^3}{\text{kmol} \cdot \text{K}}\right)} = \frac{643.54 \text{ K}}{\text{ANS}}$$