THE CARNOL PROCESS FOR CO₂ MITIGATION
FROM POWER PLANTS AND THE TRANSPORTATION SECTOR

by

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May 1995

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ABSTRACT

A CO₂ mitigation process is developed which converts waste CO₂, primarily recovered from coal-fired power plant stack gases with natural gas, to produce methanol as a liquid fuel and coproduct carbon as a materials commodity. The Carnol process chemistry consists of methane decomposition to produce hydrogen which is catalytically reacted with the recovered waste CO₂ to produce methanol. The carbon is either stored or sold as a materials commodity. A process design is modelled and mass and energy balances are presented as a function of reactor pressure and temperature conditions. The Carnol process is a viable alternative to sequestering CO₂ in the ocean for purposes of reducing CO₂ emissions from coal burning power plants. Over 90% of the CO₂ from the coal burning plant is used in the process which results in a net CO₂ emission reduction of over 90% compared to that obtained for conventional methanol production by steam reforming of methane. Methanol as an alternative liquid fuel for automotive engines and for fuel cells achieves additional CO₂ emission reduction benefits. The economics of the process is greatly enhanced when carbon can be sold as a materials commodity. Improvement in process design and economics should be achieved by developing a molten metal (tin) methane decomposition reactor and a liquid phase, slurry catalyst, methanol synthesis reactor directly using the solvent saturated with CO₂ scrubbed from the power plant stack gases. The benefits of the process warrants its further development.

I. INTRODUCTION

The evidence for greenhouse gas CO₂ warming causing global climate change is continuing to mount and international agreements are being sought to limit CO₂ emissions (10). The CO₂ emissions are primarily due to fossil fuel combustion (principally coal, oil and gas) in the industrial, commercial and transportation sectors. Although much effort in the U.S. has gone into the science of climate change, relatively little effort has been expended for technologies that mitigate greenhouse gas emissions. Improvement in efficiency of energy production and utilization is recognized as a cost effective method for reducing CO₂ emission to a limited degree (9). Fuel substitution, utilizing more natural gas and oil versus coal is recognized to further reduce CO₂ emission. The use of biomass for energy production is also effective in CO₂ reduction. A more aggressive manner for reducing CO₂ emissions is the removal, recovery and disposal of CO₂ from central power plants, which primarily burn coal. A fair amount of research has gone into disposal and sequestering CO₂ in the ocean and in depleted gas wells (9). However, CO₂ sequestration presents some formidable, technical and economic problems. Much less effort has gone into precombustion fuel processing to significantly reduce CO₂ emission. Coal gasification combined cycle is one limited step in that direction but still requires CO₂ sequestration (9). The concept of extraction and disposal of carbon from fossil fuels and utilization of the hydrogen enriched fractions has been introduced with the idea that carbon is much less difficult to store and sequester than CO₂ (9). The coprocessing of fossil fuels with biomass by the Hydrocarb process (60) producing methanol as a liquified fuel can achieve zero CO₂ emission. The use of methanol as an efficient automotive fuel can further reduce CO₂ emission from the transportation sector (9). To maximize methanol production and reduce development effort, the Hynol process which coprocesses biomass with natural gas has been introduced (9) and avoids carbon sequestration while still obtaining significant CO₂ emission reduction in a cost effective manner. In this paper we describe and develop an alternative process which converts waste CO₂ primarily recovered from coal-fired power plant stack gases, with natural gas to produce methanol as a liquid fuel and carbon as a storable materials commodity co-product.
II. THE CARNOL PROCESS DESCRIPTION

The Carnol Process grew out of a preliminary investigation of alternative processes for using CO₂. The Carnol Process relies on two basic chemical reactions: The thermal decomposition of methane and the catalytic synthesis of methanol from hydrogen with CO₂.

Methane decomposition: \[3 \text{CH}_4 = 3 \text{C} + 6 \text{H}_2\]

Methanol Synthesis: \[2 \text{CO}_2 + 6 \text{H}_2 = 2 \text{CH}_3 \text{OH} + 2 \text{H}_2 \text{O}\]

Overall Process: \[3 \text{CH}_4 + 2 \text{CO}_2 = 2 \text{CH}_3 \text{OH} + 2 \text{H}_2 \text{O} + 3 \text{C}\]

Thus 1 mole of methanol is produced from the utilization of 1 mole of CO₂ which results in a net zero CO₂ emission when the methanol is burned. It takes 1.5 moles of CH₄ to produce 1 mole of methanol or to react with 1 mole of CO₂ by means of H₂. The CO₂ mitigation comes about by removing 1 mole of CO₂ from power plant stack gas (primarily coal fired) and producing but not burning the 1.5 moles of carbon per mole of methanol produced.

Both reactions are known to take place and have been practiced in different forms on a commercial scale. Methane decomposition to form carbon black is known as the Thermal Black process. The hydrogen is not recovered in this process but is used as fuel. From an energy point of view the Thermal Black process as commercially practiced is very inefficient. A continuous catalytic methane cracking process called Hypro has been operated for hydrogen production for hydrocracking oil in a refinery, however, in this case the carbon was not recovered but was used as fuel in the process.

The catalytic methanol synthesis from CO₂ and H₂ has also been practiced commercially, however, only on a limited scale, mainly because of the lack of cost effective CO₂ feedstock. Most methanol produced currently is made by the catalytic synthesis of CO and hydrogen which is produced by the steam reforming of natural gas. There is no reduction in CO₂ emission by the use of the conventional methanol synthesis process using natural gas. In fact when coal is used to produce the synthesis gas, there is a large increase in CO₂ emission.

The reasons the Carnol process can be considered as a technically feasible and potentially cost effective means for CO₂ emission reduction are as follows:

1. Much chemical engineering development effort has recently gone into removal and recovery of CO₂ from power plant stack gases. Through the use of hindered amine absorption solvents, the energy requirement for CO₂ removal and recovery has been significantly reduced.

2. In principle, hydrogen production by methane decomposition requires the least amount of energy compared to other means of hydrogen production, such as steam reforming of methane and electrolysis of water. It only takes 18 Kcal to decompose a mole of methane. Thus the production of a mole of hydrogen requires only 5% of the energy of combustion of natural gas. The kinetics of methane decomposition has been further studied and has become better understood. High surface area carbon itself can act as an autocatalyst for improving rates of decomposition at lower temperatures.

3. Much catalyst development work has lately gone into the synthesis of methanol from CO₂ and H₂ resulting in development of improved catalysts.

4. Methanol as an alternative fuel has a number of benefits: (1) it is a liquid fuel which can be used on a large scale, (2) it can be transported and stored in accordance with the present infrastructure, (3) it can be used in stationary and automotive engines as a substitute for petroleum based fuel, thus reducing imports and improving the balance of payments and, (4) when used in internal combustion engines, it is 30% more efficient than gasoline which results in lower CO₂ emission in the transportation sector and (5) it has potential as an ideal fuel supply for efficient fuel cell power systems now under development.

5. The Carnol process converts CO₂ from power plant stacks to another useful fuel product and thus the carbon from the power plant is essentially used twice. Furthermore, the methanol fuel can obtain additional CO₂ reduction when used in the dispersed automotive sector of the economy. CO₂ emission from automotive
engines emits about one-third of the total emission of CO₂ in the U.S. which is about the same quantity of CO₂ emitted from central power plant stacks.

6. It is possible to obtain low net CO₂ emission without the use of biomass. Instead of using waste CO₂ from the atmosphere through biomass, Carnol uses waste CO₂ directly from coal burning power plant stacks.

7. It is possible to consider the large scale application of Carnol because next to coal, natural gas is abundantly available at low cost.

Based on thermodynamic principles, a first order simplified analysis of Carnol can be made using a simplified two reactor flow diagram shown in Figure 1 and given in Table 1. Hydrogen is used to provide the endothermic heat of reaction (by indirect heat transfer) for the thermal decomposition of methane so as to obtain zero CO₂ emission. The catalytic CO₂/H₂ reaction for methanol synthesis is exothermic and can produce some process steam. Table 1 indicates that there is a 61% reduction in methanol yield by the Carnol process compared to the conventional methanol process by steam reforming of methane using the same simplified procedure. However, the CO₂ emission is completely eliminated compared to conventional methanol production. Although the thermal efficiency is 49.7% compared to 81.5% by the conventional process there is available a significant quantity of carbon co-product which can be sold as a useful material on the commodity market to offset methanol costs in competing with conventional methanol cost. Thus, thermal efficiency is not the only criteria to judge the Carnol process.

### III. CARNOL PROCESS DESIGN

A process design and analysis has been made taking into account process temperature and pressure conditions. A computer simulation program was used to make a detailed mass and energy balance. The assumptions in the model are as follows:

1. Close approach to equilibrium is assumed in the methane decomposition reactor (MDR) and the methanol synthesis reactor (MSR). The equilibrium data for methane decomposition is graphically shown in Figure 2.

2. A fluidized bed MDR is assumed using an indirectly heated circulating alumina heat transport system. The rate of methane thermal decomposition is adequate, for a reasonable reactor design, at temperatures of 800°C and above.

3. The MSR is a conventional ICI type gas phase methanol catalytic convertor operating at 50 atm pressure and 260°C with a 4 to 1 recycle ratio to achieve close to 100% conversion of the CO₂ feed to the MSR system.

4. A multi-stage compressor increases the pressure of the process gas from the MDR to the MSR. The compressor is driven by steam generated from the MDR combustor exhaust gas.

5. A condenser-fractionator separates the product methanol from the water and the exothermic energy from the MSR provides the steam for the fractionator.

6. Residual gas from the MSR is recycled to the MDR for either process gas or as fuel in the combustor.

7. CO₂ is supplied as gas at 1 atm from the power plant stack gas recovery system.

A number of recycled and heat transfer configurations and process variables were explored. Table 2 gives the results of a number of computer runs for the process flow sheet configuration shown in Figure 3 (designated as Carnol III) varying the MDR pressure and temperature from 1 to 50 atm and the temperature from 800 to 1100°C respectively. Increasing temperature in the MDR decreases CO₂ emission and increasing pressure in the MDR increases CO₂ emission. Decreasing pressure in the MSR also increases CO₂ emission. Table 2 indicates that at 1 atm pressure in the MDR and temperatures from 800 to 1100°C the yield (thermal efficiency) of MeOH remains at 41.1% while the CO₂ emission is reduced by 87% and higher compared to the combustion of methanol produced by the conventional steam reforming process. From a materials point of view, temperatures in the order of 800 to 900°C for the MDR is preferable. The flow sheet of Figure 3 is based on an MDR temperature of 800°C. A summary of the mass and energy balances for Carnol III is given in Table 3 and the stream compositions in Figure 3. The decrease in thermal efficiency from the simplified analysis of 49.7% indicates the inefficiencies when taking into account detailed mass and energy balances.

The CO₂ feed to the Carnol process is provided by removal and recovery from coal fired power plant stack...
gases by a monoethanolamine (MEA) solvent absorption-stripping system. The amine system has been used for CO₂ removal and recovery from process gases in ammonia and methanol plants in the U.S. for a number of decades. Recently published papers from Japan(19) using hindered amine solvents and improved absorption column packing to decrease pressure drop across the column and an integrated system with the power plant has decreased energy requirements so that there is only an 8% loss in power from a coal burning plant when recovering 90% or better of the CO₂ from its stack gases.

IV. PRELIMINARY ECONOMIC ANALYSIS

The assumptions made are as follows:

1. CO₂ is removed and recovered from a 600 MW(e) coal burning plant (40% efficiency) using amine solvent at 90% CO₂ recovery, 90% plant factor and 10% additional capacity to make up for avoidance loss.

\[ \text{CO}_2 \text{ rate} = \frac{215 \text{ lbs}}{\text{MMBTU coal}} \times 6 \times 10^5 \text{KW(e)} \times \frac{8500\text{BTU}}{\text{KW(e)}} \times \frac{hr}{yr} \times \frac{\text{Ton}}{2000\text{lb}} - 4.34 \times 10^6 \text{ T/yr} \]  

The Carnol plant capacities are shown at the bottom of Table 4 requires 400,000 MSCFD of natural gas. The methanol production rate is 8460 T/D or 61,100 Bbbl/D and the carbon produced is 5800 TD.

2. Since, the Carnol plant has two reaction steps (MDR and MSR) and the conventional plant has two steps (steam reforming of methane and MeOH synthesis), the capital investment is based on an equivalent conventional methanol world size plant estimated at $100,000/ton MeOH/Day. Thus, the total investment then is $100,000 x 8460 T/D = $846x10^4. Production cost is estimated based on factors of capital investment as follows: 19% for financing (depreciation & interest), 1% for labor, 3% for maintenance and 2% for power and miscellaneous, resulting in a total of 25% of the capital investment on an annual basis for the production cost.

3. Natural gas prices are assumed to vary between $2 and $3/MSCF ($95 to $142/ton). It should be noted that natural gas prices in the U.S. were as low as $1.50/MSCF ($71/ton) in 1994.

4. The carbon is assumed to be stored at $10/ton C. Carbon can also result in income since it has a market in tires, pigments and newsprint inks, etc. Depending on grade, carbon can sell from $100 to $1000/ton. In Table 4, carbon price balanced production cost at less than $20/ton.

5. The cost of CO₂ to Carnol recovered from the power plant can be a highly variable quantity depending on whether there is a carbon tax, in which case Carnol can charge the power plant for disposing of the CO₂. At full cost recovery, it is estimated that $5/ton would cover the cost of CO₂ recovery, assuming 8% reduction in power plant output charged at $0.06/Kwh(e). Other CO₂ cost charges were also assumed varying from zero to $108/ton as the market income of MeOH varied.

6. The market price of MeOH has been historically around $0.45/gal ($136/ton) depending on stable natural gas feed stock costs. Recently, the MeOH market price increased to $1.30 gal ($394/ton) due to a supply shortage in its use for production of MTBE mandated as a gasoline oxygenation agent(17). This huge increase in price has a profound effect on the economics of the Carnol process. However, as soon as new MeOH capacity comes on line in the next several years, it is expected that the price will drop back to historical levels(17). At 0.45/gal MeOH competes with gasoline at $0.73/gal based on a 30% improvement in IC engine efficiency (1.54 gal MeOH is equivalent to 1 gal gasoline)(9). No credit is taken in this paper for the use of methanol as a transportation fuel which would result in an additional 33% reduction in CO₂ emission compared to gasoline.

In Table 4, production cost factors were equated to income factors and the CO₂ credit was determined in the last column and evaluated as the figure of merit for the process. The conclusions drawn from this analysis are as follows:

1. When operating the MDR at 900°C and above and the MSR at 50 atm, the CO₂ emission reduction is greater than 90% compared to CO₂ emission for methanol production by the conventional process.
2. With no cost for feedstock CO₂ to Carnol, natural gas at $2/MSCF, no credit for carbon, and methanol at
$0.45/gallon, the cost of reducing CO₂ emission is $25/ton (listed as negative credit). This is less than the average IEA estimate for removal, recovery and sequestering CO₂ in the ocean at $37/ton neglecting transportation (pipelining) costs to the ocean. At $3/MSCF the CO₂ reduction cost using Carnol increases to $55/ton which is the upper limit for ocean disposal of CO₂ neglecting pipelining to the ocean.

3. By selling the carbon as a commodity at $58 and $126/ton when natural gas cost $2 and $3/MSCF respectively, the CO₂ reduction cost is reduced to zero. Since the carbon is very pure this carbon price of $0.06/lbs or less would have an easy market to compete with current prices of carbon black of up to $0.50/lb. The U.S. market for tire carbon amounts to 2 x 10^6 tons/yr. and there are other uses for carbon at a low cost price, for example as a filler in construction materials.

4. If the power plant wants to recover its cost for recovering CO₂ up to as high as $10/ton, at a natural gas cost of $3/MSCF, carbon produced by Carnol would have to sell for $170/ton ($0.085/lb) to achieve zero CO₂ reduction cost, which is still a very reasonable possibility.

5. If the methanol can continue to demand a $1.30/gal or almost 3 times the historical price, at $3/MSCF for natural gas and CO₂ feedstock cost recovered from the power plant of $5/ton (recoverable cost) and assuming no carbon sales, a CO₂ credit of $103/ton for reducing CO₂ emission can be realized. On the other hand, if the CO₂ credit for reducing emissions is reduced to zero, the power plant could charge as much as $108/ton for feeding its CO₂ to the Carnol plant. Obviously the charges and profits could be negotiated between the power plant and the Carnol plant.

V. ADVANCED CARNOL VI PROCESS

Two recent developments have been uncovered that could significantly improve the basic Carnol process. One refers to the methane decomposition and the other to methanol synthesis.

1. The design of an efficient methane decomposition reactor (MDR) can be difficult because high temperature energy must be provided to decompose the methane and the particulate carbon must be recovered and removed in a continuous manner. As mentioned earlier, intermittent reheat batch reactors and fluidized bed reactors have their drawbacks. Recently, we have found that molten metal technology is being applied to decompose liquid and solid carbonaceous waste material to produce simple gaseous compounds using a molten iron bath (Fe) at temperatures from 1400°C up to 1650°C(14). The advantage of using a molten metal bath to decompose waste material can be readily applied to the decomposition of methane. Molten metal is a good liquid phase direct contact heat transfer media through which gaseous methane can be bubbled. The large density differences between solid carbon and the molten metal could allow efficient capture from the gas phase and separation of the carbon particles from the liquid phase by flotation. Although molten iron at temperatures up to 1600°C could completely decompose methane to its elements, carbon and hydrogen, as indicated by the equilibrium diagram in Figure 2, the use of such extreme temperatures is a disadvantage when it becomes necessary to design a thermally efficient heat recovery system. Molten tin (Sn) at a lower temperature appears to be a suitable molten metal media for an MDR for the following reasons:
   a) The liquid range for Sn is much wider than Fe; Mpt = 236°C to Bpt 2260°C.
   b) Density of liquid Sn = 7.31 gm/cc
   c) partial pressure of Sn at 1000°C = < 10^-4 atm (< 1 ppm)
   d) Molten tin in the range of 800 to 1000°C should be sufficient to decompose methane to a high degree.
   e) The molten metal tin bath may also be catalytic for decomposing methane.
   f) Carbon does not react or dissolve in liquid tin.
   g) Surface nitrided refractory metal (titanium or molybdenum) can provide adequate corrosion resistant materials for heat transfer and containment of the molten tin.
   h) The viscosity of the molten tin is low which provides for good mixing between the gaseous and liquid phases.

2. Recently liquid phase catalytic synthesis of methanol has been shown to improve production of methanol because of improvement in transferring the exothermic heat of the synthesis reaction(15). The catalyst is in a
slurry form in an organic solvent carrier such as an oil or glycol. For application in the Carnol process it then becomes possible to practice liquid phase methanol synthesis by reacting hydrogen with CO₂ when it becomes absorbed in the MEA solvent during recovery from the power plant stack gases. A methanol synthesis catalyst would be carried in slurry form in the MEA absorbent. The conditions for the synthesis can be estimated from the phase diagram shown in Fig. 4 using a hindered amine solvent. For example, absorbing flue gas CO₂ from a coal burning plant at equilibrium at 40°C, produces a solution having a CO₂ to amine ratio of 0.58. Heating this solution to 120°C gives an equilibrium partial pressure of CO₂ above this solution of 100 psia (6.8 atm). By pressurizing this solution with hydrogen up to about 30 atm pressure thus providing a 3 to 1 H₂ to CO₂ pressure ratio in the presence of the slurry catalyst, methanol should be formed. Table 5 estimates the equilibrium concentration of methanol at 30 and 50 atm and at 120 and 260°C respectively, when feeding a 3 to ratio of H₂ to CO₂ mixture. This data indicates a much improved yield of methanol at the lower temperature, which results in a lower recycle ratio and improved economics of the process.

Applying the above two developments, a Carnol VI Process flowsheet is designed and is shown in Figure 5. The heat recovery around the molten tin reactor and the separation of hydrogen from the unreacted methane by pressure swing adsorption (PSA) to produce a pure H₂ stream is shown in Fig. 6. A heat and mass balance for Carnol VI indicates that the overall thermal efficiency for production of methanol is 49.7%. By storing the carbon or using it as a materials commodity, the net CO₂ emission, taking credit for CO₂ from the power plant is 13.0 KgCO₂/GJ (30.2 lbs CO₂/MMBTU) which represents an 83% reduction in CO₂ emission compared to the production of methanol by conventional process, i.e. the steam reforming of natural gas. When methanol is used in internal combustion engines an additional 33% reduction in CO₂ emission is obtained compared to the use of gasoline as automotive fuel.

VI. CONCLUSION

The Carnol process which produces methanol as a liquid fuel, can effectuate a very significant net decrease in CO₂ emission from coal fired power plants. The economic value is significantly improved when the coproduct carbon can be sold as a materials commodity. Two R and D efforts which can significantly improve the process are (1) developing a molten metal methane decomposition reactor and (2) developing a liquid phase MEA slurry catalyst reactor for directly converting CO₂ scrubbed from power plant fuel gas with hydrogen from methane decomposition to produce methanol as a liquid fuel for the automotive industry. The benefits of the process warrants its further development.

REFERENCES


### Table 1
SIMPLIFIED THERMODYNAMIC ANALYSIS OF CARNOL PROCESS

<table>
<thead>
<tr>
<th>Unit Operations</th>
<th>Reaction</th>
<th>Enthalpy, ΔH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decomposition:</td>
<td>(3\text{CH}_4 = 3 \text{C} + 6\text{H}_2)</td>
<td>(+18\text{ Kcal/mol CH}_4)</td>
</tr>
<tr>
<td>MeOH Synthesis:</td>
<td>(2\text{CO}_2 + 6\text{H}_2 = 2\text{CH}_3\text{OH} + 2\text{H}_2\text{O})</td>
<td>(-22\text{ Kcal/mol MeOH})</td>
</tr>
<tr>
<td>Combustion:</td>
<td>(\text{H}_2 + \frac{1}{2}\text{O}_2 = \text{H}_2\text{O})</td>
<td>(-68\text{ Kcal/mol H}_2)</td>
</tr>
</tbody>
</table>

**Carnol Process Analysis**
- Moles CH\(_4\) to produce 2 moles MeOH = 3.000
- Moles CH\(_4\) to produce combustion H\(_2\) for Ht transfer to CH\(_4\) = 0.455
- Moles MeOH per total mole CH\(_4\) = 2.00/3.455 = 0.579
- Higher Ht of combustion of MeOH = 182,000 Kcal/Mol
- Higher Ht of combustion of CH\(_4\) = 212,000 Kcal/Mol
- Carnol MeOH thermal efficiency = 49.7%
- Carbon produced per mol MeOH = 3.455/2.0 = 1.728

CO\(_2\) emission = 2 mol CO\(_2\) (from stack gas) + 2 mol CO\(_2\) from MeOH combustion = 0

**Conventional Process Analysis\(^{11}\)**
- Moles MeOH produced per mol CH\(_4\) = 0.95
- Thermal efficiency = 81.5%
- Moles CO\(_2\) produced per mol CH\(_4\) = 1.05

**Relative to Conventional Methanol**
- Carnol process CO\(_2\) reduction = 100%
- Yield of Carnol MeOH to conventional = 61%
<table>
<thead>
<tr>
<th>Computer Run No.</th>
<th>MDR P atm/t°C</th>
<th>MSR P atm/t°C</th>
<th>CO₂ Feed Stock Kg</th>
<th>MeOH Thermal Eff. %</th>
<th>MeOH Carbon Eff. %</th>
<th>CO₂ Emission LBS MMBTU HHV MeOH</th>
<th>CO₂* Emission Reduction</th>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>1/800</td>
<td>50/260</td>
<td>156.6</td>
<td>41.1</td>
<td>50.3</td>
<td>22.7</td>
<td>87.4</td>
</tr>
<tr>
<td>5</td>
<td>1/900</td>
<td>50/260</td>
<td>147.1</td>
<td>41.1</td>
<td>50.4</td>
<td>10.2</td>
<td>94.3</td>
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<tr>
<td>6</td>
<td>1/1000</td>
<td>50/260</td>
<td>143.1</td>
<td>41.1</td>
<td>50.4</td>
<td>5.0</td>
<td>98.2</td>
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<td>10</td>
<td>1/1100</td>
<td>50/260</td>
<td>142.5</td>
<td>41.5</td>
<td>50.8</td>
<td>2.7</td>
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<td>1/800</td>
<td>30/260</td>
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<td>30/120</td>
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<td>23.0</td>
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<td>50/260</td>
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<td>10/900</td>
<td>10/120</td>
<td>133.3</td>
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<td>38.6</td>
<td>90.9</td>
<td>49.5</td>
</tr>
<tr>
<td>7</td>
<td>30/1000</td>
<td>50/260</td>
<td>122.2</td>
<td>30.3</td>
<td>145</td>
<td>Increase in Emission</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>10/1000</td>
<td>10/260</td>
<td></td>
<td></td>
<td></td>
<td>-- NO BALANCE OBTAINED --</td>
<td></td>
</tr>
</tbody>
</table>

*Emission reduction is compared to production of methanol by conventional steam reforming of natural gas which produces 180 LBS CO₂/MMBTU of methanol energy (HHV). Thermal efficiency for a conventional steam reforming plant for methanol production = 64%
Table 3
CARNOL PROCESS III DESIGN
PROCESS SIMULATION - MASS 7 ENERGY BALANCES

<table>
<thead>
<tr>
<th>UNIT</th>
<th>CARNOL III+ H₂ - RICH GAS FUEL FOR MDR</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>MDR</td>
<td></td>
</tr>
<tr>
<td>Pressure, atm</td>
<td>1</td>
</tr>
<tr>
<td>Temperture, °C</td>
<td>800</td>
</tr>
<tr>
<td>CH₄ Feedstock, Kg</td>
<td>100</td>
</tr>
<tr>
<td>Preheat Temp, °C</td>
<td>640</td>
</tr>
<tr>
<td>CH₄ Fuel for MDR, Kg</td>
<td>-</td>
</tr>
<tr>
<td>CH₄ Conversion, %</td>
<td>91.9</td>
</tr>
<tr>
<td>Carbon Produced, Kg</td>
<td>68.8</td>
</tr>
<tr>
<td>Heat Load, Kcal</td>
<td>82,091</td>
</tr>
<tr>
<td>Purge Gas for Fuel, Kmol</td>
<td>2.4</td>
</tr>
<tr>
<td>MSR</td>
<td></td>
</tr>
<tr>
<td>Pressure, atm</td>
<td>50</td>
</tr>
<tr>
<td>Temp., °C</td>
<td>260</td>
</tr>
<tr>
<td>CO₂ Feedstock, Kg</td>
<td>156.6</td>
</tr>
<tr>
<td>CO₂ Conversion, %</td>
<td>90.9</td>
</tr>
<tr>
<td>Methanol Prod., Kg</td>
<td>100.6</td>
</tr>
<tr>
<td>Water Cond., Kg</td>
<td>58.7</td>
</tr>
<tr>
<td>Energy for Gas Compression to MSR</td>
<td></td>
</tr>
<tr>
<td>Energy, Kcal</td>
<td>75,114</td>
</tr>
<tr>
<td>Performance</td>
<td></td>
</tr>
<tr>
<td>Ratio, Methanol/CH₄, Kg/Kg</td>
<td>1.01</td>
</tr>
<tr>
<td>Carbon Efficiency MeOH, %</td>
<td>50.3</td>
</tr>
<tr>
<td>Thermal Eff. MeOH, %</td>
<td>41.1</td>
</tr>
<tr>
<td>Thermal Eff. C + MeOH, %</td>
<td>81.8</td>
</tr>
<tr>
<td>CO₂ Emission, Lbs/MM BTU</td>
<td>22.7</td>
</tr>
<tr>
<td>CO₂ Emission, Kg/GJ</td>
<td>9.8</td>
</tr>
</tbody>
</table>
Table 4

PRELIMINARY CARNOL PROCESS ECONOMICS

Costs shown in $10^8$/yr and (Unit Costs)

Investable Capital Cost (IC) = $8.46 \times 10^8$

<table>
<thead>
<tr>
<th>PRODUCTION COST FACTORS</th>
<th>=</th>
<th>INCOME FACTORS</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25 IC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$10^8$ IC</td>
<td></td>
<td>$10^8$ yr</td>
</tr>
<tr>
<td>Natural Gas</td>
<td></td>
<td>(MSCF)</td>
</tr>
<tr>
<td>2.12</td>
<td>2.60</td>
<td>($2)</td>
</tr>
<tr>
<td>2.12</td>
<td>3.90</td>
<td>($3)</td>
</tr>
<tr>
<td>2.12</td>
<td>2.60</td>
<td>($2)</td>
</tr>
<tr>
<td>2.12</td>
<td>3.90</td>
<td>($3)</td>
</tr>
<tr>
<td>2.12</td>
<td>2.60</td>
<td>($2)</td>
</tr>
<tr>
<td>2.12</td>
<td>3.90</td>
<td>($3)</td>
</tr>
<tr>
<td>2.12</td>
<td>3.90</td>
<td>($3)</td>
</tr>
<tr>
<td>2.12</td>
<td>3.90</td>
<td>($3)</td>
</tr>
<tr>
<td>2.12</td>
<td>3.90</td>
<td>($3)</td>
</tr>
</tbody>
</table>

*Based on the following plant capacities
1) CO₂ rate, 90% recovered from a 600 MW(e) net (650 MW(e)) gross coal fired power plant = 4.34 x 10⁶ T/yr
2) CH₄ rate = 2.77 x 10⁶ T/yr = 400,000 MSCF/D
3) MeOH produced = 2.78 x 10⁶ T/yr = 61,100 Bbl/D. = 8460 T/D
Table 5

METHANOL SYNTHESIS EQUILIBRIUM

Input: $\text{H}_2$ 3 mole
      $\text{CO}_2$ 1 mole

<table>
<thead>
<tr>
<th>$P$ (atm)</th>
<th>30</th>
<th>30</th>
<th>50</th>
<th>50</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T$ (°C)</td>
<td>120</td>
<td>260</td>
<td>120</td>
<td>260</td>
</tr>
<tr>
<td>CO (mole)</td>
<td>0.0007</td>
<td>0.1365</td>
<td>0.0004</td>
<td>0.1089</td>
</tr>
<tr>
<td>$\text{CO}_2$ (mole)</td>
<td>0.4459</td>
<td>0.7686</td>
<td>0.3285</td>
<td>0.6865</td>
</tr>
<tr>
<td>H$_2$O (mole)</td>
<td>0.5541</td>
<td>0.2314</td>
<td>0.6715</td>
<td>0.3135</td>
</tr>
<tr>
<td>H$_2$ (mole)</td>
<td>1.3391</td>
<td>2.5787</td>
<td>0.9862</td>
<td>2.2773</td>
</tr>
<tr>
<td>MeOH (mole)</td>
<td>0.5534</td>
<td>0.0949</td>
<td>0.6711</td>
<td>0.2046</td>
</tr>
<tr>
<td>Total (mole)</td>
<td>2.8932</td>
<td>3.8101</td>
<td>2.6577</td>
<td>3.5908</td>
</tr>
</tbody>
</table>
Fig. 1
Simplified Carnol Process for Producing Methanol From Natural Gas and CO₂ for Zero CO₂ Emission
Mass Flows Are In Moles
Fig. 2

EQUILIBRIUM DATA

\[ \text{CH}_4 = \text{C} + 2\text{H}_2 \]

Y\text{CH}_4 - Mol Fraction

Total Pressure atm

100 10 30 50 1

0 200 400 600 800 1000 1200

TEMPERATURE - degrees Centigrade
**CO₂ Mitigation Technology Carnol-III + Process**

Fig. 3

- **Combustor**
  - 1 atm
  - 2.8 kmol
  - Alumina or Hex.
  - MDR: 1 atm, 800 °C
  - 640 °C
  - 260 °C
  - 0.2 kmol
  - 2.86 kmol
  - 188 °C
  - CO₂ Feedstock From Power Plant Flue Gas: 156.6 kg, 20 °C

- **MDR** Carbon to Storage: 68.8 kg
- **COMP.**
  - 59.5 kmol
  - 197 °C
  - MSR: 50 atm, 260 °C
  - 200 °C
  - 138 °C
  - 59.5 kmol
  - CIRC.

- **CON**
  - 50 atm, 50 °C
  - MeOH: 101 kg
  - H₂O: 58.7 kg

**Gas Stream**

<table>
<thead>
<tr>
<th>Rate-Kmol</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temp-°C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Comp. mol%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CO</td>
<td>3.35</td>
<td>3.68</td>
<td></td>
</tr>
<tr>
<td>CO₂</td>
<td>12.86</td>
<td>14.15</td>
<td></td>
</tr>
<tr>
<td>CH₄</td>
<td>4.25</td>
<td>17.34</td>
<td>19.08</td>
</tr>
<tr>
<td>H₂O</td>
<td>4.76</td>
<td>0.14</td>
<td></td>
</tr>
<tr>
<td>H₂</td>
<td>95.75</td>
<td>56.67</td>
<td>62.34</td>
</tr>
<tr>
<td>MeOH</td>
<td>5.02</td>
<td>0.60</td>
<td></td>
</tr>
</tbody>
</table>

- **Carbon Efficiency**: 50.3%
- **Thermal Efficiency**: 41.1%
- **CO₂ Emission**: 22.7 lbs/MMBTU
- Basis 100 kg CH₄ Feed
Fig. 4

CO₂ - H₂O - Amine Phase Equilibrium
For Liquid Phase Methanol Synthesis
(Taken From Suda et al, Nov. 1993, Sweden)

KS-1 Extrapol. to 120 °C - ppCO₂ = 100 psia

Coal Fired Plant
15% CO₂ Flue Gas
pp CO₂ = 2.25 psia

CO₂/Amine Ratio = 0.58
CO₂ In Solution/Amine (mole ratio)
Fig. 5

Carnol VI Process for CO₂ Mitigation Technology
Combining CO₂ Recovery From Power Plants With Liquid Metal Methane Decomposition and Liquid Phase Methanol Synthesis

MEA Scrubber with MeOH Catalyst Slurry 1 ATM-40 °C
Liquid Phase Methanol Convertor 30 ATM 120 °C
MeOH-H₂O Fractionator 30 ATM
PSA - H₂/CH₄ Sep. Compressor From 1-10 ATM to 30 ATM
Molten-Metal Tin Methane Decomp. Reactor 1-10 ATM 800 °C - 900 °C

Feedstock

Process Chemistry:

\[ \frac{3}{2} \text{CH}_4 = \frac{3}{2} \text{C} + 3\text{H}_2 \] → Product → MeOH Synthesis

\[ 3\text{H}_2 + \text{CO}_2 = \text{CH}_3\text{OH} + \text{H}_2\text{O} \] → MeOH Synthesis

Nat. Gas Decomp.

Flue Gas

Product
Fig. 6

Molten Metal Methane Decomposition Reactor For Carnol VI Process

Basis 1.0 g-mol CH₄ Process Gas