SOLAR THERMOCHEMICAL ENERGY CONVERSION AND TRANSPORT

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The high temperature catalytic dissociation of $SO_3$ and the $CO_2$-$CH_4$ reforming-methanation cycle are important chemical processes being considered in the development and application of solar-thermal energy conversion, transport, and storage systems. Separate facilities for...
evaluating chemical converter-heat exchangers at temperatures to 1000°C with high flow rates of SO\textsubscript{2} and of mixtures of CO\textsubscript{2} and CH\textsubscript{4} feedstocks have been assembled and operated on the NMSU campus. A number of receiver elements (chemical reactors) have been tested in these laboratory facilities in an effort to optimize catalyst parameters and catalyst reactor configurations. These tests led to the design and fabrication of both low power and high power solar energy receivers which were operated successfully at the White Sands Solar Furnace. Energy delivery methanation reactor design and parametric studies led to the fabrication and operation of a laboratory closed-loop, energy conversion, transport, and delivery system. These latter experiments met with limited but promising success. Carbon deposition, though a problem, is believed to be controllable with the optimization of catalyst parameters and feedstock composition.
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I. INTRODUCTION

Thermochemical capture of solar energy - Solchem - is an important concept in the development of economical, practical, alternate energy systems. Solchem provides an alternative to the more conventional method of transporting solar thermal energy to directly drive a heat engine power cycle. Solchem involves the use of the high temperature heat available from concentrating collectors to activate an energy absorbing chemical reaction which will then produce an energy rich chemical product. The thermochemical receivers involve no moving parts, only gas passages and catalytic surfaces.

Two gaseous chemical fluids have been considered for thermochemical energy conversion systems. These are $\text{SO}_3$ and a mixture of $\text{CH}_4$ and $\text{CO}_2$. The $\text{SO}_3$ dissociation reaction may be written as

$$\text{SO}_3 \rightarrow \text{SO}_2 + \frac{1}{2}\text{O}_2 + 22 \text{ kcal/mol.}$$

The forward reaction is used in a receiver to convert solar energy into chemical heat of reaction. This energy is then transported and is deposited in a storage system via the reverse reaction. The use of $\text{SO}_3$ as a working fluid offers the following advantages: (1) the lack of side chains in the chemical reaction; (2) the sensitivity of the reaction to control by catalysis; (3) the relatively low gas pressures.

involved; and (4) the temperature range of gas dissociation and recombination. The disadvantages are: (1) toxicity; (2) incompatibility with low cost materials; (3) corrosiveness at receiver temperature; and (4) poor thermal conductivity.

The CO₂-CH₄ reforming-methanation reactions and the accompanying shift reaction can be written as

\[ \text{CO}_2 + \text{CH}_4 \rightarrow 2\text{H}_2 + 2\text{CO} + 61 \text{ kcal/mol.} \]

\[ \text{CO}_2 + \text{H}_2 \rightarrow \text{H}_2\text{O} + \text{CO}. \]

The reforming chemical cycle offers the following advantages: (1) the relative nontoxicity of the components of the working fluids; (2) the low pressure and temperature requirements for both feedstock and synthesis gases; (3) the compatibility of the working fluids with low cost materials; and (4) no serious corrosion problems at receiver temperature. The disadvantages are: (1) complicated chemistry with several possible side reactions; and (2) carbon deposition can be a problem.

The work performed under this contract falls into three categories. These are: (1) SO₃ dissociation experiments; (2) CO₂-CH₄ reforming experiments; and (3) methanation and closed-loop experiments.

II. SO₃ DISSOCIATION EXPERIMENTS

Although SO₃ is a toxic chemical, there are well established rules for handling the material which have led to production rates of \( 2 \times 10^{10} \) kg/yr. In addition, SO₃ is the end product of SO₂, which is introduced into the atmosphere in enormous quantities during the burning of coal.
Its use in a closed loop energy conversion and transport process thus imposes negligible ecological problems. The use of SO\textsubscript{3} does, however, involve significant material problems. At 100\textdegree C, these problems are not severe, although common material choices are largely restricted to Teflon, mild steel, stainless steel, glass, quartz and ceramics. At receiver temperature (1000\textdegree C), Teflon and steel are not usable. While there are indications that aluminium rich alloys, such as Kanthal A-1, may be usable, the materials of choice for SO\textsubscript{3} energy converters are fused silica and ceramics.

The objective of the present work is to measure the dissociation efficiency of several reactor-catalyst configurations. A facility was assembled at NMSU which permitted the testing of prototype Solchem SO\textsubscript{3} dissociation reactors under conditions similar to those which would be encountered in a full-scale solar power conversion plant. A typical reactor with its counter-current heat exchanger is configured as a flat helix whose outer diameter is 0.75 m and whose inner diameter is 0.30 m. This test article is positioned in a furnace in such a manner that only the innermost turn of the helix, the converter turn which contains the catalyst, is heated. Electrical heaters, simulating the focused solar beam, were located within a cavity which was surrounded by the reactor's inner (converter) turn. The test facility does not utilize closed-loop operation; gaseous feedstock, evaporated from liquid SO\textsubscript{3}, was introduced to the reactor at controlled temperature, pressure, and flow rate. The reactor effluent was completely scrubbed after being analyzed to prevent the release of SO\textsubscript{2} or SO\textsubscript{3} to the environment. From the molar fraction of SO\textsubscript{2} measured in the effluent gas the power converted to chemical heat of reaction was calculated.

From September 1979 through September 1980, twenty-five separate tests were conducted in the SO\textsubscript{3} laboratory on four separate reactors. The conditions attending these tests and the experimental results have been published\textsuperscript{5,6}. Conversion powers of 2kW with conversion efficiencies of greater than 50\% were observed in these single coil tests.
III. CO₂-CH₄ REFORMING EXPERIMENTS

A facility was assembled at NMSU in which reforming converter-heat exchangers were tested under conditions similar to those which would be encountered in a full-scale solar power conversion plant. A typical reformer was configured as a helical, tube-in-tube, counter-current heat exchanger surrounding a single turn converter in which the reforming reaction is catalyzed. Most of the receiver elements studies in this work were fabricated from type 316 stainless steel tubing. Other materials, including types 304 and 310 stainless steel and Kanthal A-1 tubing, have also been utilized. In nearly all of the (approx. 20) reactors tested, the outer diameter of the outer tube was 25.4 mm while that of the inner tube was 15.9 or 19.0 mm. The wall thickness of these tubes is typically 1.2 mm. Heat exchanger lengths have varied from 4 to 6 turns. In some of the reactors the converter turn was the inner turn of the helix, while in others a circular converter, manifolded to the heat exchanger, was used. A number of catalyst and catalyst configurations were tested. In the first reactor tested, the converter was loaded with small (6x6 mm) alumina pellets which were coated with NiO catalyst. Other converters studied during the first phase (1979) of the program contained NiO coated alumina Raschig rings. The number and distribution of these rings within the converter tube were varied and optimized early in the project.

Early in the project it was found that catalyst deterioration was an important problem. In an effort to circumvent this difficulty, different catalyst configurations and materials were studied. A short


sequence of tests was conducted with tubewall reactors. These converters contained Ni catalyst material deposited directly on the inner wall of the converter tubes. For the tubewall converters tested the conversion efficiency was unacceptably low. The most successful catalyst material used in the reforming work was ruthenium. The material was in the form of small, perforated stainless steel saddles, coated with a ruthenium bearing surface layer. The converter tube was tightly packed with the Ru coated stainless steel saddles. The Ru catalyzed reactors were found to have good long-term stability. During ten months of tests and experimentation there was no evidence of catalyst deterioration.

Two solar energy receivers were assembled and operated at the White Sands Solar Furnace in December 1979 and January 1981 respectively. At this facility, the full power of the furnace is concentrated within a conical envelope with a half angle of 45° and a focal spot of less than 10 cm dia. In December 1979 the full power capability of the furnace was 30 kW; in January 1981, 26 kW was the maximum power available.

From May 1979 through January 1981 approximately eighty separate tests were conducted in the reforming laboratory at NMSU and at the White Sands Solar Furnace. These tests involved some twenty reactors utilizing many different catalyst-converter configurations. The results of this work have been published 7-11.


IV. CLOSED-LOOP EXPERIMENTS

Relatively little work has been done on the energy delivery part of the Solchem system. An attempt was made to operate an ordinary re-forming receiver element as a methanator. The synthesis gas used as the feedstock in these tests consisted of mixture of 3 parts (by volume) CO to 3 parts H₂ to 2 parts CO₂. From the results of this work it was concluded that a good reformer is not necessarily a good methanator. The results, however, did indicate that methanation reactivity would be improved if the space velocity could be reduced by a large factor. A methanation reactor was then fabricated which was similar to the reformer receiver elements except that the converter turn was made from 5.08cm O.D. stainless steel tubing with 1.2mm thick walls. The converter was loaded with tightly packed NiO Raschig rings of the type described earlier. With space velocities in the range of 500 hr⁻¹, this reactor was found to perform well as a methanator, delivering about 1kW of heat while producing an effluent gas which contained 25% CH₄ and less than 5% CO from the 3:3:2 feedstock described above.

From April through October 1981 a sequence of experiments was performed in the NMSU Solchem reforming laboratory to demonstrate a low-power, laboratory model, closed-loop, energy delivery system. Figure 1 shows a chemical engineering flow chart for the system. Included in the chart are temperatures and pressures at various points in the loop.


Due to the fact that project funding was running low at this time, every effort was made to minimize spending in the construction and operation of the closed-loop test facility. Virtually all of the hardware comprising the loop were taken from the reforming and SO$_3$ test facilities described earlier. The most difficult item to procure was the blower. After a lengthy search, a $300 blower manufactured by a milking machine company was acquired and found to be technically adequate for short-term operations.

A total of sixteen test runs were made in the closed-loop facility. The first such run (Run 80) used a start-up (make-up) gas comprised of 77.5% H$_2$, 21.1% CO, and 1.4% CO$_2$ (volume percent). During this 2.75 hour test, the mass flow slowly dropped from 0.5 g/s to 0.1 g/s. The methane concentration in the reformer feedstock line was high (>40%) and the CO concentration in the synthesis gas line ran from 12 to 20% during most of the run. The [CO] in the feedstock line ran <1%. Approximately 500 watts of power was delivered to the methanation furnace. The test was terminated due to blower failure. Runs 81 and 82 were hampered by blower and catalyst problems.

On May 18, 1981 Run 83 was conducted with the same start-up gas composition as used in Runs 80-82. Figures 2-13 contain plots of the results from this test. The test was terminated after eight hours due to carbon build-up in the lines and on the mass-flow transducer. Except for the first and last hours of Run 83 the pressures, pressure drops, mass flow, feedstock and synthesis gas composition, and all temperatures exhibited very stable operation as power was delivered to the methanation reactor furnace. This demonstration of closed-loop operation was considered to be a success in spite of carbon build-up problems. After start-up in all of these runs the methanation furnace heaters were removed and methanator cooling was achieved by blowing compressed air through the furnace cavity.
Runs 84 and 85 utilized a slight increase in the hydrogen concentration of the start-up gas in an effort to decrease carbon deposits. The results from these runs were little different from Run 83 with line carbonization still a problem.

In August, September, and October 1981, Runs 86 through 96 were conducted with a start-up synthesis gas mixture of 80% H₂ and 20% CO₂. With the reformer operating at 900°C and the methanator at 550 to 600°C the equilibrium feedstock and synthesis gas compositions should be as follows:

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<th>Synthesis Gas</th>
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<tr>
<td>H₂</td>
<td>38.4%(vol.)</td>
<td>62.8% (vol.)</td>
</tr>
<tr>
<td>H₂O</td>
<td>35.2</td>
<td>17.1</td>
</tr>
<tr>
<td>CO</td>
<td>3.2</td>
<td>16.6</td>
</tr>
<tr>
<td>CO₂</td>
<td>7.2</td>
<td>3.3</td>
</tr>
<tr>
<td>CH₄</td>
<td>16.0</td>
<td>0.3</td>
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These numbers were calculated assuming 50% CO₂ reforming and 50% steam reforming at 4 atmospheres pressure.

During the course of the eleven runs made under these conditions, a total of 41 operating hours was put on the closed-loop system. The system generally performed well. There were some carbon fouling incidents, and considerable difficulty was experienced with the blower. Run 91 was typical of this series and graphs of its results are shown in Figures 14 through 25. The equilibrium values of the gas concentrations which were measured are consistent with those predicted above. Clearly, it would be desirable to measure the concentration of all constituents, but instrumentation for this purpose was not available.

It is believed that the carbon problem can be eliminated through the optimization of catalyst, catalyst configuration, pressure, temperatures, and gas composition. If this is not possible, the carbon
problem can be minimized and coped with by proper mechanical and chemical engineering design.

V. CONCLUSIONS

The work reported here has clearly demonstrated not only the feasibility but the practicality of the chemical conversion and transport of solar energy. The present field model Solchem receiver is capable of operating at the focus of a 6m paraboloidal collector with an efficiency of 60-65%. This receiver efficiency could be increased to perhaps 85% by improving the cavity design. If the spacing of receiver elements in the cavity were optimized, the difference between the temperatures of the front and back converter coils could be reduced from 350°C (Run 74) to less than 100°C. With all six coils operating in the 850-950°C temperature range and with 2 g/s of gas flowing through each, results show that the improved cavity receiver would be converting about 20kW of solar power. A major cause of energy loss was reflection of light from the back surface of the cavity out of the entrance aperture. Addition of a special receiver element designed to cover this surface would largely eliminate this source of energy loss. Finally, receiver heat losses could be reduced through the use of more efficient thermal insulation and through use of a concave cone shaped entrance window. Such an entrance cone would reduce both convective and radiative aperture losses.

Nickel catalyst deterioration was a serious problem early in the project. This problem, however, appears to have been resolved by the use of a ruthenium-on-stainless-steel catalyst. During hundreds of hours of operation of ruthenium catalyzed reformers no effects due to
catalyst degradation have been observed. Side reactions have presented minor problems. The shift reaction leads to the formation of small quantities of steam which are controlled by the steam reforming reaction and by allowing line temperatures to operate above their respective dew points.

A major problem area in closed-loop operation was found to be carbon deposition. It is strongly felt, however, that this problem can either be eliminated or tolerated through the optimization of chemical and mechanical design.
Start up synthesis gas composition

H₂  77.16% (vol)
CO  21.37 "
CO₂  1.47 "

May 18, 1981 Run No. 83 Closed Loop Solgas 0.6

SOLCHEM PRESSURE SET PRESSURE IN +

TIME IN MINUTES (×10¹)
FIGURE 5.

U, U, == 0.4 to 0.5, e^A

Flow transducer failed
FIGURE 8.
FIGURE 9.
May 18, 1981  Run No. 83  Closed Loop  Solgas 0.6

THERMOCOUPLE GRAPHS
THERMOCOUPLE 7  +
THERMOCOUPLE 8  X
Methanator Converter
FIGURE 13.

[Graph showing temperature profiles with annotations for thermocouples and temperature scale from 0.00 to 18.00°C.]
Start-up Synthesis Gas Composition

\[ H_2 \quad 79.7\% \text{ vol.} \]
\[ CO_2 \quad 20.3\% \]

Sept. 18, 1981  Run No 91  Closed Loop
FIGURE 15.

Sept. 18, 1981  Run No. 91  Closed Loop

SOLCHEM PRESSURE SET
PRES IN - PRES OUT +

DELTA P METH (PSI)

0.00
0.30
0.60
0.90
1.20
1.50

TIME IN MINUTES (x 10)

0.00  5.00  10.00  15.00  20.00  25.00  30.00  35.00  40.00
FIGURE 17.

Sept. 18, 1981 Run No. 91 Closed Loop

SOLCHEM MASSFLOW SET
MASS FLOW +

Time in Minutes

MASS FLOW (G/M SEC)
0.00 0.30 0.60 0.90 1.20 1.50
Sept. 18, 1981    Run No. 91    Closed Loop

THERMOCOUPLE GRAPHs
THERMOCOUPLE 1   +
THERMOCOUPLE 2   X
Reformer Converter

TIME IN MINUTES (x10^1)
Fig. 22: Thermocouple Graphs

Thermocouple 5: X
Thermocouple 6: O

Methanator Converter

Sept. 18, 1981
Run No. 91
Closed Loop
FIGURE 24.
<table>
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