# Selected Technical and Economic Comparisons of Synfuel Options

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# **INCREASED AUTOMOBILE** FUEL EFFICIENCY AND SYNTHETIC FUELS

**Alternatives For Reducing Oil Imports** 

Background Paper #2

Selected Technical and Economic Comparisons of Synfuel Options

October 1982



CONGRESS OF THE UNITED STATES

# Preface

This volume contains papers written for OTA to assist in preparation of the report Increased Automobile Fuel Efficiency and Synthetic Fuels: Alternatives for Reducing Oil Imports. OTA does not endorse these papers. In several instances, the OTA report reaches somewhat different conclusions because of additional information which was obtained later. These papers, however, may prove valuable for readers needing more detailed or specific information than could be accommodated in the final assessment report, and are being made available for such purposes.

#### SELECTED TECHNICAL AND ECONOMIC COMPARISONS

### OF SYNFUEL OPTIONS

FINAL REPORT

April 1981

Prepared For The Office of Technology Assessment United States Congress

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#### EXECUTIVE SUMMARY

#### Study Scope and Content

This study is a comparative technical and economic assessment of selected synfuel technologies. It is a component part of a much larger study being conducted by the OTA on energy options. A key purpose of this study is to provide technical and economic comparisons among selected synfuel technologies which, to the extent possible, provides a background and basis which may assist the OTA in its policy deliberations. The synfuel technologies have been selected in consultation with and guidance from the OTA. They generically represent: oil shale production; direct and indirect coal liquefaction oil (including Mobil-M gasoline synthesis); and coal gasification (low, medium and high Btu) . The OTA Synfuels Advisory Board has been particularly helpful in providing for and reviewing information on these selected technologies, although by no means are they held accountable to or responsible for the study products.

The study effort built upon earlier work which attempted to the extent possible to standardize the engineering, planning, and estimating base of many processes. These efforts, as described in Chapters 2 through 4, were modified and extended to include additional concerns such as upgrading concerns and plant cost escalation concerns.

These standard or generic process units have been utilized, in conjunction with the assessment of site-specific planned/proposed synthetic fuel projects, to develop a set of alternate supply deployment scenarios. Two scenarios--a "business-as-usual" scenario, and an accelerated "pushingthe-limit" --have been developed in consultation with and direction from OTA staff.

Constraints and concerns affecting the scenario assessments have been discussed, as well as a discussion of the consequent labor needs. Supply site selection concerns, as well as end-use utilization concerns have also been identified.

As specified in the Introduction (Chapter 1), the study scope was confined to an assessment of the technical and economic comparisons of the selected synfuel technologies. On-line guidance and direction was provided by the OTA in making "mid-stream" technical and economic choices in the study effort. No assessment or interpretation of the policy implications was conducted as that was strictly considered outside of study scope, design, and performance. Such policy concerns were reserved for the OTA, with its well-established and defined review procedures.

## Study Findings

The study analysis investigated numerous technical and economic aspects of the selected synfuel technologies. Because of the very complex nature of this topic--as well as the need for cautious and critical qualification of the findings--each chapter attempts to summarize its findings in its own setting.

In brief, a snapshot of the study findings are as follows:

- 1. There is a fairly consistent relationship between the cost of the synthetic fuel product and the "quality specifications" of the product, as it is used in current end uses. Criteria used to measure product quality specifications include hydrogen content; octane number, aromaticity, lubricity, and a host of physical and chemical parameters (discussed in Chapters 4 and 5) that affect specific end use technology performance.
- 2. Although perhaps an oversimplification, there appears to be a high correlation, in a cost dimension, with the product "quality specificity" and the amount of hydrogen content and/or the average range of distillation of the product. Both coal and oil shale contain a lower fraction of hydrogen than petroleum. Natural gas and crude oil having the highest percent; oil shale next; and coal with the lowest fraction. Crude oil is a broad range material, with a broad range of quality, in this sense, as well. If we are to upgrade coal and oil shale to a series of products, of varying quality and specifications, comparable to the average yield of petroleum products, we can expect that, in addition to the greater extrac-tion cost of coal and shale, there will be a greater reforming, processing or upgrading cost. This cost reflects the necessary changes to be made in physical, chemical, and material properties of the primary synfuels products to make them equivalent in use to existing refined products. Going beyond that, the cost is proportional to the specific product yield quality or quality mix that is pursued. For example, it is much cheaper to produce a Low-Meal Btu gas from coal than a High Btu gas. Similarly, it is more costly to produce a low boiling, high hydrogen containing

fuels such as motor gasoline than to produce a higher boiling, lower hydrogen containing fuel such as home heating type fuel oil products.

3. Specific process differences disappear to a large degree under our analysis, and the cost of synthetic product is more dependent upon its composition than upon the particular process used. On the other hand, certain processes (or process approaches) may be more selective for certain type products or product-slates and hence more efficient and economical routes to those specific products.

Our review and analysis of approaches to the process of upgrading raw (direct) liquid fuels to stable combustion fuels and subsequently refining them to transportation quality fuels shows us the following:

- (a) At present, indirect liquefaction technologies such as Fischer-Tropsch are known technologies. As such, there is limited technical risk. At present, however, these processes are relatively expensive because of the chemical nature of breaking down hydrocarbons and later resynthesizing them. Most of the initial coal liquids projects will be indirect liquefaction processes.
- (b) Certain resources favor certain product slates. Coal ravors the production of highly aromatic products, like high octane gasoline pool maphthas. Shale oil favors middle distillate products. Sour Crude contains many times the (high boiling) residual content as these resources and favors the production of boiler fuels, although it is certainly less expensive, at present, to produce gasoline and middle distillates from heavy and sour crudes than from coal or shale.
- 4. Cost Comparisons (in 1980 \$)
  - (a) Fully (risk) discounted cost estimates of representative or generic coal-conversion processes vary from \$10.00 to \$16.00 per MMBTU of product.
  - (b) Future expectations of technology gains in the form of capital productivity may reduce these costs by over 30% (i.e., to about the \$12.00 per MMBTU0.

- (c) Upgraded costs add as much as \$2.00-\$2.50 per MMBTU (or \$10.00-\$15.00 per barrel) of product to the cost of oil shale liquids and-direct coal liquids.
- (d) Refined transportation fuels are expected to have the following cost ranges:
  - (i) oil shale liquids \$60-70/barrel
    (ii) indirect coal liquids \$70-80/barrel
    (iii) direct coal liquids \$80-90/barrel

### 5. Transportation Concerns

- (a) Transportation of synthetic liquids and gases are most likely to be transported by pipeline, with supplementary use of water borne carriers (where available), and unit tank train railcars.
- (b) Patterns of synfuel plant and refinery siting are expected to be influenced by both resouce location as well as existing infrastructure (existing pipeline capacity; existing refinery capacity).
- 6. Synfuel Deployment
  - (a) The development of reasonable scenarios of synfuel plant commercial deployment is extremely sensitive to the product role assigned to natural petroleum feedstocks, both domestic and imported.
  - (b) From the assessment of currently planned/ proposed commercial projects (described in Chapter 5), which provide the grassroot basis for our development of scenarios, we note that most commercial projects are directed toward the production of high grade fuels. Furthermore, due to the ever expanding cost of upgrading to meet increasingly stringent product user specifications, processes are being chosen to minimize these costs, and maximize high grade product yields. Oil shale, methanol, and Mobil-M gasoline are three examples of such product choices. A perceived outlook for natural crude supplies see higher volumes of lower grade crude oils available (sour crudes from Alaska and Saudi Arabia; heavy crudes with high viscosity from Venezuela and Bakersfield) . These crudes will require major refinery upgrading and consequent refinery investment, although this investment is considerably less than for synthetic fuels. Hence, redundant

investment in the synthetics area may occur if an integrated view is not taken. We believe that an integrated view will most likely be taken by companies engaged in regions considered, although these views may represent a regional and company specific optimization.

(c) Synfuel development will require the resolution of numerous technical, economic, and socioeconomic concerns. Key among these concerns are a provision of materials, as well as engineering and skilled labor requirements. Drag lines, air compressors, and large diameter reactor vessels are examples of material needs. Chemical engineers availability is an example of the latter needs. It is felt, however, that these needs can be met, even in the high scenario, with the <u>early</u> development of programmatic plans. Similarly, early planning can relieve or avoid potential socioeconomic and community disruption.

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# CHAPTER 1: INTRODUCTION

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#### CHAPTER 1: INTRODUCTION

### 1.1 Role of Synthetic Fuels

Energy consumption in the U.S. has become increasingly dependent upon foreign sources, especially in the liquid fuels area. Transportation energy usage is a dominant user of foreign petroleum. Motor gasoline alone accounts for over 35% of all petroleum products consumed in the Us. (Reference No. 1 ); petroleum itself accounting for over 43% of all the energy consumed in the U.S. (Reference No. 2 ).

Unfortunately, over the past 35 years, the ratio of U.S. oil reserves to total U.S. oil consumption has declined, even with Alaskan North Slope oil discoveries. On the other hand, oil imports have been increasingly filling the gap in petroleum supply-demand imbalances. From 1950 to 1977, domestic petroleum production fell from an average 85% of total domestic petroleum consumption to 47% in 1977 (Reference No. 3 )\* This trend has been somewhat slowed down recently by increased energy conservation measures --especially in the transportation sector--but it has not stopped. The impacts of this increasing dependence on foreign crude oil and refined products have been staggering. In addition to the increased and continual exposure to supply interruptions, and subsequent national security vulnerability, the direct costs of these imports have increased enormously (Reference No. 4 ). From a modest plateu of 1-2 billion/year in the 1958-68 time period, the direct costs have mushroomed to 25 billion in the embargo period (1973-74) , and are heading for 90-100 billion in 1980 The impacts of this capital drain (Reference No. 5). in domestic investments, subsequent jobs, and consumer inflation has been notable. In the third quarter of 1979 alone, domestic prices for energy, housing; food, and medical care rose at an annual 17.6% rate--with energy prices escalating at a 50.1% annual rate. Adverse impacts have not been confined to the U.S. domestic economy. Oil bills, being raised by OPEC faster than inflation--not only account for 25-50% of total inflation rates around the world, but also pose a global inflationary problem, apparently without end--unless alternate or substitute fuel supplies are found\developed in sufficient quantities and at competitive prices to put the lid on world  $\bar{\mbox{crude}}$ price escalation in a timely fashion.

#### Alternate Synthetic Fuels

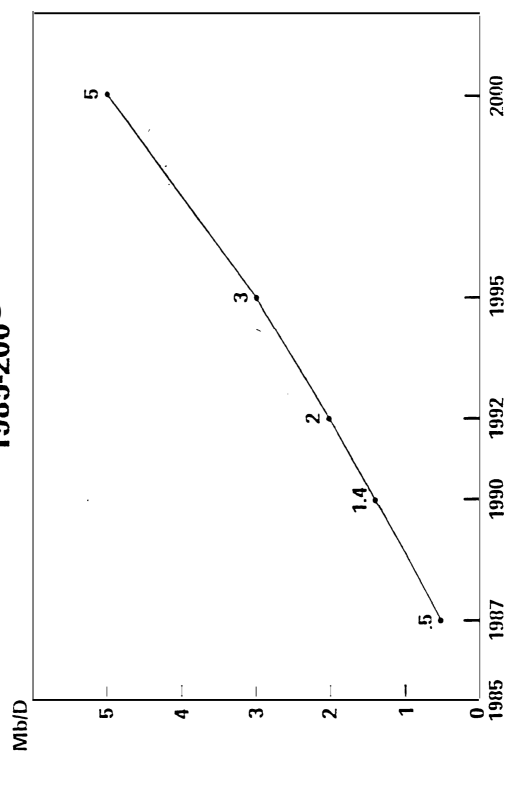
Many recent studies (Reference No. 6) have estimated the domestic energy and petroleum supply-demand imbalances. Most have credited conservation with decreasing petroleum demand from its historical rates of growth, and most have nonetheless projected a need for alternate domestic liquid fuels to fill the increasing domestic petroleum supplydemand imbalances.

Although most studies have agreed on the need for/and future role-of alternate domestic-fuels, they have differed in projecting their rate of growth in the marketplace, date of introduction, prospective cost, ease of usage, and "raw" resource availability -- as well as their potential environmental, health, and safety impacts. The U.S. Department of Energy has recently targeted synthetic production goals to reach 5 million barrels/day of crude oil equivalent from all synthetic sources by the year 2000 (Figure 1.1), and the recently passed Energy Security Act (6/30/80) has targeted goals of . 5 MMBD by 1987 and 2.0 MMBD by 2000. Although current forecasts vary, synthetics have generally been forecast to provide between 12-13% of total domestic energy by the year 2000, and even up to 30% of primary liquid fuel supplies. Although composition of those synthetic fuel targets and projections are varied (shale, unconventional gas, biomass, solar, ...), coal--as both a feedstock for synthetic fuels and as a direct combustion boiler fuel--generally has been projected to play a large and growing role. In many ways, this is a natural reflection of the abundant and regionally diverse U.S. coal resources and reserves. This is similarly true for shale as described in Chapter 2.

## 1.2 Scope of Study

The study design of this effort is, in a broad fashion, to provide for a technical and economic comparison of various selected synfuel technologies. As outlined in the contract study Scope of Work, the study team was directed to use existing published (and referenced) information and data. OTA staff and the Synfuels Advisory Group assisted in the acquisition of published data, as well as providing guidance and review. The study team was further directed to look solely at technical and economic aspects of selected synfuel technologies and specifically not at policy implications, interpretations, and concerns. These very





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real policy considerations are the stated prerogative of the OTA itself and its existing well-defined review procedures.

In consultation with the OTA staff, generic technology choices have been made (Chapters 3 and 4), and supply deployment scenarios developed (Chapter 5). Each chapter, and sub-section, specifically identifies the respective referenced sources and assumptions used. Where available in the literature, comparative estimates have been provided. Scope, timing, and budget greatly limited the degree of first-hand data verification. The recent ESCOE coal conversion study, as referenced in Chapter 4, was the scope directed starting point for the comparative economic analysis, with specific cost basis and assumptions provided in the addendum to Chapter 4.

The outline of the report is as follows:

Introduction to Role of Synthetic Fuels and Study Effort: chapter 1 Background on synthetic Fuel Processes Chapter 2 Discussion of Selected Synthetic Fuel Technologies: Chapter 3 Discussion and Comparison of Selected Synthetic Fuel Technologies Cost and Product Economics: Chapter 4 Supply Deployment Scenarios for Synthetic Fuels: Chapter 5 Appendices Glossary Bibliography

### Potential Next Steps

Potential next steps to the broad-based study effort could include site-specific, technology-specific detailed technical, economic, and socioeconomic evaluations. Sitespecific supply transportation and product distribution needs and costs; assessments of facility-specific integration of synfuel facilities with existing refinery capacity; and site and region-specific socioeconomic and labor/skill mix needs. Case study assessment are subexamples.

On the policy side, the OTA using this study, as well as other component study efforts, will be developing policy interpretations.

# CHAPTER 2: BACKGROUND

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#### Chapter 2: BACKGROUND

#### (I) <u>Description of the coal Conversion and Oil Shale Retorting</u> Fuel Cycles

### 2.1 Overview of the Coal and Oil Shale Fuel Cycles

In order to estimate investments from mine to end user (excluding automobiles), or to assess the rates of potential development of the synthetic fuels industry, it is necessary to consider all phases of the fuel cycles involved in the development. They include exploration for the resources, their mining, local transportation, beneficiation, transportation to conversion plants, conversion of the energy resources to fuels, and finally, their distribution to end users. Figures 2.1 and 2.2 (modified from Reference No. 7) describe in a schematic manner the energy systems involved in the case of coal conversion and of oil shale retorting.

#### 2.2 Coal and Oil Shale Resources

Coal and oil shale resources are defined as those deposits "that can be extracted and processed to yield products that can be marketed at a profit" (Reference  $N_0.8$ ). Estimates of resources are not limited by whether or not the deposits have been demonstrated, or whether they are extractable by existing technologies at competitive economic costs. If the resource has been demonstrated (i.e. its location, quality, and quantity have been determined by evidence supported by measuements) and its extraction is economically feasible, then it is classified as a reserve. Resources may become reserves as a result of changes in technical or economic development. The major coal and oil shale resources of the coterminous United States are shown in Figure 2.3 (Reference No. 9) and 2.4 (Reference No. 8) . The United States reserves and resources of coal are estimated as 178 and 1,285 billion metric tons (Reference No. 10). Other estimates vary widely, depending on economic and technical assumptions. For example a recent estimate of recoverable reserves of coal (Reference No. 11) places them at 38,000 guads<sup>2</sup> or equivalent to 150 billion metric tons of coal.

<sup>1</sup> A recoverability factor of 50 percent is assumed for resources.

2<sub>Quad</sub> isaunit of energy equivalent to 10<sup>15</sup> (quadrillion) Btu. It is approximately equivalent to 180 million barrels of oil or to 40 million metric tons of bitumious coal. On the average, one quad is enough to supply all the present energy requirements of about 3 million Americans for one year.

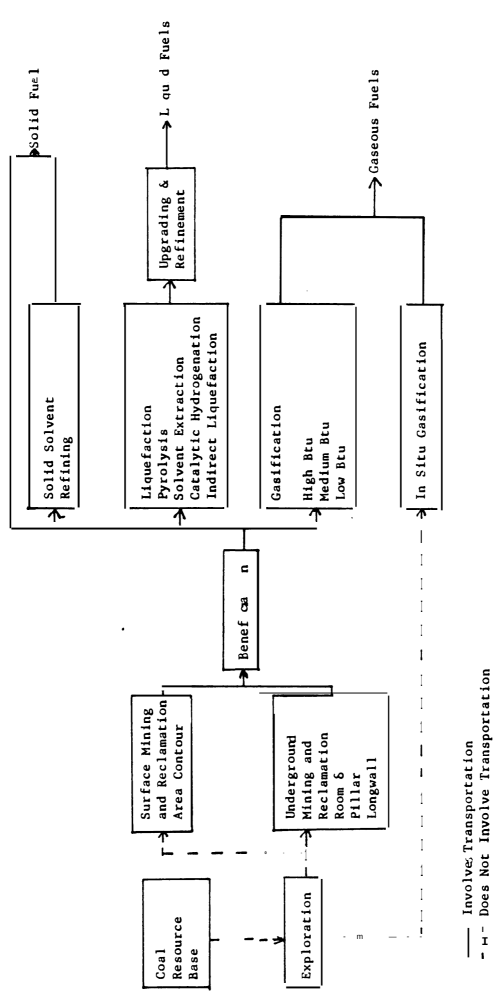
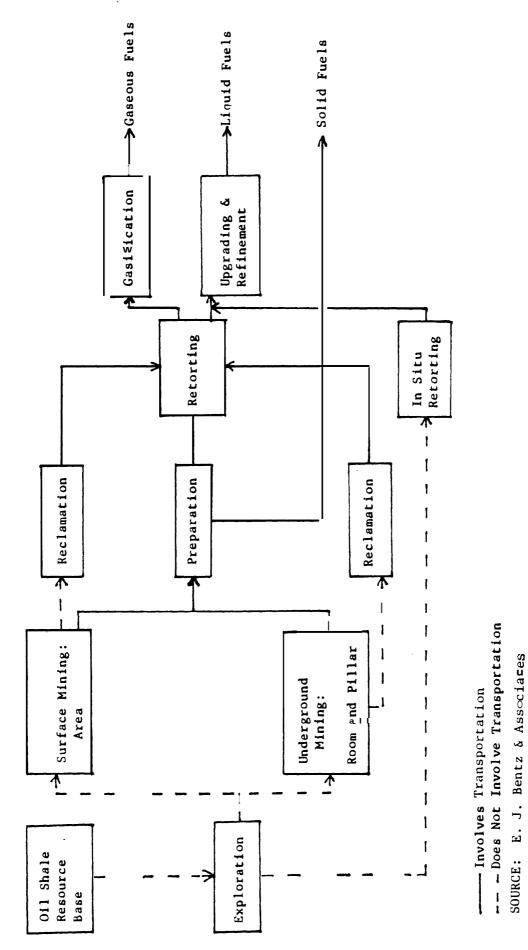


FIGURE 2.1 Coal Conversion System

E. J. Bent≥ & Associates

SOURCE :

2-2





2-3

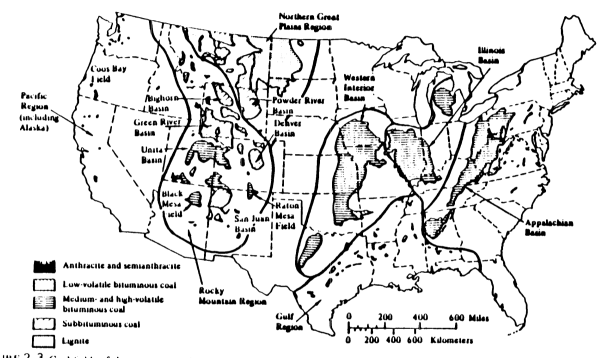
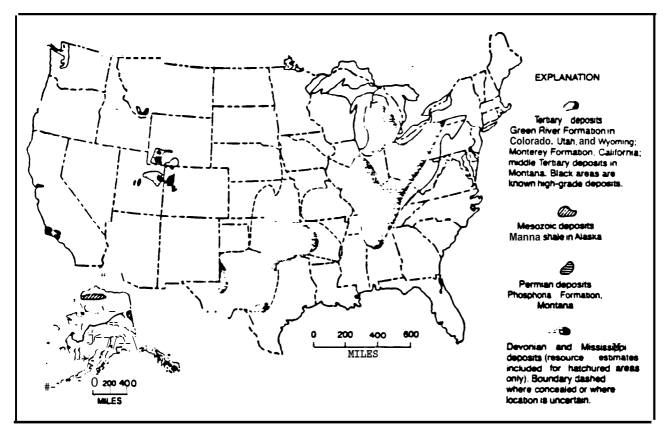


FIGURE 2, 3 Coal fields of the coterminous United States. Source: Adapted from Paul Averiti, Coal Resources of the United States, January 1, 1974, U.S. Department of the Interior, Geological Survey Bulletin 1412 (Washington, D.C.: U.S. Government Printing Office (Stock No. 024-001-02703), 1975), p. 5; and U.S. Department of Energy, Energy Information Administration, Coal Data (Washington, D.C.: U.S. Government Printing Office (DOE/E1A-0064), 1978), p. 1.



## Figuro24: Oil Shale Deposits of the United States

SOURCE D C Duncan and V E. Swanson, Organic-Rich Shales of the United States • nd World Land Areas, U.S Geological Survey Circular S23. 1965.

Estimates of recoverable reserves and resources of oil shale vary even more broadly than those of coal because of the poor existing state of knowledge. An estimate by the Committee on Nuclear and Alternative Energy Systems of the National Academy of Sciences places them at 3660 quads of recoverable oil at \$21.50 -\$27.50 in 1978 dollars (Reference No. 9) which is equivalent to about 1.50 billion metric tons of real. **Schur's estimates** of shale oil reserves are only 1100 quads which is equivalent to 44 billion metric tons of coal (Reference No. 11) . Oil shale resources have been estimated by OTA (1980) as equivalent to between 2,000 and 140,000 billion barrels of oil equivalent, or 440 to 3,090 billion tons of Coal.

The distribution of the coal and oil shale resources are given in Tables 2.1 (Reference No. 12) and 2.2 (Reference No. 8). There are large variations among the characteristics of coal and oil shale resources, as well as the characteristics of sites at which they are found. These characteristics affect the processes, economic rests and resource requirements of the development of a synthetic *fuels industry*. The important variables are the quantity and quality of the coal and oil shale in each site or province, ownership, relationships to markets and processing facilities, bed depth, seam thickness, availability of water resources, and competition for surface area usage. These are discussed at greater length in Appendices A and B (Reference No. 7).

#### 2.3 Exploration and Mining

Knowledge about coal and oil shale resources is usually obtained in stages. The steps begin with the assessment of geological and geophysical data and are followed by surface and areal photographic surveys and magnetic measurements. Finally, mapping and appraisal of regional deposits are done, based on seismic surveys and drilling. The steps are explained in greater detail in references below.

There are two basic methods of coal and oil shale mining, namely surface mining and underground mining. The choice between them depends mainly on the depth of burial and thickness of the seam. In the case of coal seams that are relatively close to the surface (i.e. less than 180 feet) surface mining is employed (Reference No. 13). In the case of oil shale, where the deposit is within a few hundred feet (200 to 300 feet) from the surface, it can be surface mined (Reference No. 14). However, higher quality oil shale is commonly located at depths of over 600 feet, so that it may be more efficient to apply underground processes of retorting rather than mine the shale (Reference No. 8).

A qualitative description of the mining methods and their impact are given references 7 and 8. Appendix A to Chapter 2 summarizes the the major components, resource requirements, costs and pollutants

## TABLE 2.1

## U.S. BY STATES RESOURCES-RESERVE BASE-PRODUCTION BITUMINOUS-SUBBITUMINOUS-LIGNITE COAL

# (Millions of Tons)

State	USGS Remaining Identified Resources January 1, 1974	Yearly Production 1977	Estimated Remaining Reserve Base January 1, 1978
Alabama Alaska Arizona Arkansas Colorado Georgia Illinois Indiana Iowa Kansas East Kentucky West Kentucky West Kentucky West Kentucky Louisiana Maryland Michigan Missouri Montana New Mexico North Carolina North Dakota Ohio Oklahoma Oregon Pennsylvania South Dakota Tennessee Texas Utah Virginia Washington West Virginia Washington West Virginia Wyoming Other States (Calif., Idaho, Nebr., Nevada)	$15,262 \\130,079 \\21,234 \\4,938 \\148,850 \\1+ \\146,001 \\32,868 \\6,505 \\18,668 \\22,226 \\36,120 \\1,000 \\1,152 \\205 \\31,184 \\291,639 \\61,387 \\110 \\350,602 \\41,116 \\7,117 \\334 \\63,940 \\2,185 \\2,530 \\139,000 \\23,359 \\9,216 \\6,194 \\100,150 \\136,891 \\688 \\$	$\begin{array}{c} 21 \\ <1 \\ 11 \\ 12 \\ <1 \\ 54 \\ 28 \\ <1 \\ 92 \\ 51 \\ 0 \\ 3 \\ 0 \\ 7 \\ 29 \\ 11 \\ 0 \\ 12 \\ 46 \\ 5 \\ 0 \\ 83 \\ 0 \\ 10 \\ 17 \\ 9 \\ 38 \\ 5 \\ 95 \\ 44 \\ 0 \end{array}$	1,823 $11,642$ $308$ $668$ $14,815$ $1+$ $65,286$ $10,495$ $2,882$ $1,385$ $12,360$ $35,788$ $800$ $1,027$ $118$ $9,457$ $108,282$ $4,344$ $32$ $15,954$ $20,736$ $1,276$ $57$ $23,335$ $428$ $932$ $3,210$ $3,982$ $29,225$ $1,932$ $38,822$ $53,182$ $447$

Table From: Solid Fuels for U.S. Industry, Cameron Engineers, 1979

## TABLE 2.2

#### POTENTIAL SHALE OIL IN PLACE IN THE OIL SHALE DEPOSITS OF THE UNITED STATES (billions of barrels)

	Range of shale oil yields, gallons per ton		
Location	5 - 10ª	10 - 25 <sup>⁼</sup> 25 - 100ª	
Colorado, Utah, and Wyoming (the Green River formation)	4,000	2,800	1,200
Central and Eastern States (includes Antrim, Chattanooga, Devonian, and other shales)	2,000	1,000	(?)
Alaska.****	Large	200	250
Other deposits.*.*.*** .******** .******9	134,000	22,500	(?)
Total.*e*9***0**** .******** .**9***	140,000+	26,000	2,000(?)

<sup>a</sup>Order of magnitude estimate includes known deposits, extrapolation and interpolation of known deposits, and anticipated deposits.

Data from: D.C. Duncan and V.E. Swanson, Organic-Rich Shales of the United States and World Land Areas, U.S. Geological Survey Circular 523, 1965.

associated with <u>generic</u> surface and underground coal and oil shale mining in East and West.

The following conclusions can be drawn about coal and oil shale, namely:

- 1. The Northern Great Plains and Rocky Mountain Provinces contain approximately 70 percent of the coal resources in the United States and most of the nations low-sulfur coal (References No. 7 and 14a).
- Much of the coal likely to be developed in the near future can be surface mined. This estimate is based on existing trends of continued shift from underground to surface mined coal (References No. 1& and 14c), and on the abundant quantities of coal that can be mined by existing surface mining technologies (References 7 and 14<u>a)</u>.
- Competition for surface area usage is relatively low in those areas of coal mining (Reference No. 7).
- 4. The federal government controls the majority of the coal and oil shale lands (References No. 7, 8, and 1<u>4a)</u>.
- Water resources can become a constraint on coal development in the Rocky Mountain and Northern Great Plains Provinces (References No. 7, 9, and 14d).
- 6. The development of oil from oil shale resources involves tremendous quantities of materials that need to be mined and disposed. The production of 1 million bbl oil per day from oil shale would require the mining and disposal of about 1.3 million metric tons of shale per day (Refer-e No. 15).
- Most oil shale extraction is expected to be by underground mining, with only about 1.5 to 20 percent being extractable by surface methods (Reference No. 16). This proportion may change with technological developments.

## 2.4 Beneficiation

Coal and oil shale feedstocks require some preparation, called beneficiation, prior to their feeding into the conversion process.

The nature of the preparation depends on the characteristics of the feedstocks, and on the type of conversion process adopted. In some cases, mechanical upgrading is sufficient, and consists of any or all of *the* following processes:

- 1. Crushing and screening
- 2. Cleaning
- 3. Drying

In some cases of coal feedstocks, further processing is required. A portion of the ash and sulfur can be removed from the coal by simple **pro**cedures such as water washing, or magnetic separation of iron pyrites. Further upgrading of the coal can involve chemical processes, such as reacting the coal with various chemicals, or converting it to more desirable products. A further discussion of coal beneficiation is included in Reference No. 7 and Reference No. <u>13</u>.

Appendix Table 8 (Reference  $N_0.17$ ), summarizes the major components and resource requirements of coal beneficiation. Appendix A to Chapter 2 compares the costs of various chemical coal cleaning processes.

Oil shale beneficiation consists mainly of crushing and sizing. The process is further discussed in Reference No. 8.

#### 2.5 Transportation

Local transportation is mainly limited **to** the transfer of the coal or Shale between different parts of the mining area. Truck, belt conveyor, or rail transport are the most used means.

**Coal** *al so needs* to be transported beneficiation plants, and in large quantities and over large distances to coal conversion plants. There are a number of alternatives for transporting the coal, namely, railroads (both unit and conventional trains), slurry pipelines, and to a lesser extent, barges and trucks. The transportation of coal is further discussed in Reference 7. Appendix to Chapter 2, References 7 and 18, summarize the major components and resource requirements of transportation.

In the case of oil shale, siting of the conversion plan is near the mining area is envisaged. This is because of the tremendous quantities of shale involved.

#### 2.6 Conver~sion

The Conversion of coal and oil shale to other energy products is

covered in detail in Chapter 3 of this report.

### 2.7 Upgrading and Refining

Raw synfuels may be used directly in the market for **scme** applications without further upgrading; or they may require modifications before they can become substitutes for existing products. The need for different variations of upgrading will be detemined by the characteristics of the synfuels, and their uses. Liquid synfuels can be utilized in many end uses, the most important of which are the transportation, space heating, raising of steam in boilers and as chemical feedstocks. Substitution of coal or oil shale derived liquids for petroleum based fuels particularly in transportation, will create problems because of the differences between them. They differ mainly in the types and quantities of hydrocarbon species involved in the overall ratio of hydrogen to carbon atoms in the mixture, and to a lesser extent, an increased presence of ash, trace metals, and nitrogen compounds.<sup>3</sup> While the ratio of hydrogen to carbon is approximately 2 for petroleum, it drops in general to 1.9 for shale oil and in general 0.75 for coal derived liquids (Reference No. 19), although this depends on the specific product slate and operating conditions. The addition of substantial amounts of coal or shale derived fuels will mainly decease the hydrogen to carbon ratio, and increase the aromatic, nitrogen, and trace metal content of the refinery products.

The concerns and costs associated with <u>selective</u> upgrading are discussed in Chapter 4. The key concern is to match anticipated product demand slate specifications and tolerances with variable feedstock inputs (from West Texas crude to shale oil) at <u>least cost</u>. The factors that affect the cost are the kind of strategies that have to be developed to meet the challenge, and the decision whether to upgrade the synfuel at the conversion plant or at the refinery.

There are several strategies that can be used to adopt synfuels to product demand. One is to modify the engines using at present petroleum derived fuels to match the characteristics of synthetic fuels; another is to modify the synthetic fuels; a third is to develop an optimum combination of changes in both the supply and end use sections. Still, it should be pointed out, that many variations of upgrading can be conceived, not necessarily requiring conversion of the total raw synfuel streams to refined products Rather, some Synergistic effects can be used to incorporate synfuels upgrading into a variety of refining schemes, with significant improvement in *economics*.

 $<sup>^3</sup>$  Raw coal distillates contains 100 times the nitrogen of conventional petroleum (Reference No. 19) .

Early analysis (Reference No. 19) suggested that in the <u>short run</u>, <u>selective</u> synthetic fuel upgrading can alleviate fuel distribution concerns. More recent analyses (Reference No. 20) suggest that <u>transition solutions</u> will probably entail synthetic fuel finished <u>products - such as methanol in modified automotive fleet engines</u>. These matching concerns reflect the sensitivity of combustion engines to the hydrocarbon makeup of the fuel. Experience has \* that the combustion of fuels low in hydrogen content and rich in aromatics results in an increased formation of soot, in addition to various other in-field maintenance problems.

Existing and anticipated petroleum refining technology can upgrade synthetic oils to meet current engine and turbine specification. This is primarily done by the hydrogeneration of crudes. For most existing refineries, the development of such upgrading capabilities would require costly changes in the reactor vessels to withstand high pressures, and a further supply of hydrogen. Therefore, an economic evaluation needs to be carried out for each specific situation. It would determine whether product upgrading is more cost effective when conducted together with the primary coal hydroliquefaction step, thus forcing the conversion process to produce finished, more premium hydrocarbon liquids; or whether upgrading should be combined with refining.

Oil from shale with hydrogen to carbon ratio of 1.9 (vs 2 for petroleum) can be substituted for present fuels with some relative ease. Oil from coal conversion with hydrogen to carbon ratio of 0.75 requires more upgrading.

Preliminary studies indicate that the upgrading of the H/C ratio and reducing the aromatic and organic nitrogen contents of synthetic crudes is feasible but expensive in terms of costs and energy losses. chapter 4 discusses these cost comparisons<sup>4</sup>. Various estimates have been prepared for upgrading. Among them are references for coal conversion and shale . Estimates have been prepared by Chevron, U.S.A.

5 Other authors (Reference Nos. 21, 22, 23, 24 and 25) indicate and describe processes for upgrading shale oil but no comparable cost and energy estimates are given.

2-12

<sup>4 -</sup> uncertain still. surrounds the costs of alternate fuels for heat engines since absolute costs will not be established until fuel production plants are built and operated. However, for the purpose of initial screening of alternate fuels, relative rests can be established from published studies. Comparing these studies on a consistent basis in terms of total delivered costs and engine efficiencies is more important than the assessment of absolute product costs shown by **such** studies.

putting the cost of upgrading crude shale oil for a 100,000 bbl/d facility at 6.50, in first quarter 1978 dollars, equivalent to about 7.80 in 1980 dollars (Reference No. 8). Total cost of upgrading and refining crude synthetic fuels vary according to the capacity and location of the refinery, the nature of the crudes involved and the available facilities and options at the refinery. Estimates of the refining costs for crude shale oil ranged from 8.00 to 12.00/bbl (Reference No. 8). In the case of refineries modified for crude shale oil, estimates as low as 25.50 to 2.00/bbl are reported (Reference No. 8). Upgrading of these crudes may also result in energy losses as large as 25 to 50 percent of the original energy in the coal (Reference No. 19).

There are several studies underway to define capabilities of state-of-the-art petroleum refineries for syncrude upgrading and development of new refining methods specifically tailored towards syncrudes (Reference Nos. 22, 23, 26, 24 and 21). The preliminary conclusions that can be drawn are:

- 1. Syncrudes can be refined by conventional methods.
- 2. Products are interchangeable with petroleum derived products.
- 3. There are serious economic and energy penalties in upgrading (Reference No. 19), but research leading to improved refining processes to @@- the syncrudes and engine development to use then are expected to reduce the penalties.
- 4. Direct coal liquids may require more severe upgrading than shale oil (Reference No. 19).

The processing details of upgrading of various coal conversion and oil shale derived crudes as well as their properti<sup>\*</sup>es relative to petroleum males are given in references 22, 23, 26a and 26b.

Crude synthetic fuels can be upgraded either at the synfuel plant or at a refinery<sup>6</sup>. The upgrading process is similar in many respects to the refining of crude petroleun. Therefore, there may be economic and technical incentives to combine the two operations in one plant. Utilization of existing facilities, and the available options of existing refineries to mix syncrudes and petroleum crudes to ease the upgrading process are other advantages. Furthermore, upgrading requires water, so that the location of many potential synfuel conversion plants in dry areas may dictate the separation of the two. However,

<sup>6</sup> Most existing refineries will need to be modified before they can handle syncrudes.

a decision on whether to upgrade synfuels at the conversion plant or in the refinery should be based on a detailed analysis with consideration given to location and marketing factors.

### 2.8 Distribution to End Users

There is similarity between syncrudes and petroleum crude oil. It is therefore most likely that the mode of distribution will. be through the presently existing crude oil pipelines shown in Figure 2.5 (Reference No. 27). Some new pipeline additions or extensions will undoubtedly be built, depending on the location of the syncrude plants However, it is likely that the location of crude oil pipelines, as well as the availability of coal, water, etc., will be taken into account in siting the plants. Once the syncrude has entered the pipeline distribution system, it will probably be treated as another source of crude, as is presently done with syncrude from Canadian tar sands, and districted to refineries as a supplement to natural crude supplies (Reference No. 27).

#### (II) Synfuel Technologies Parameters

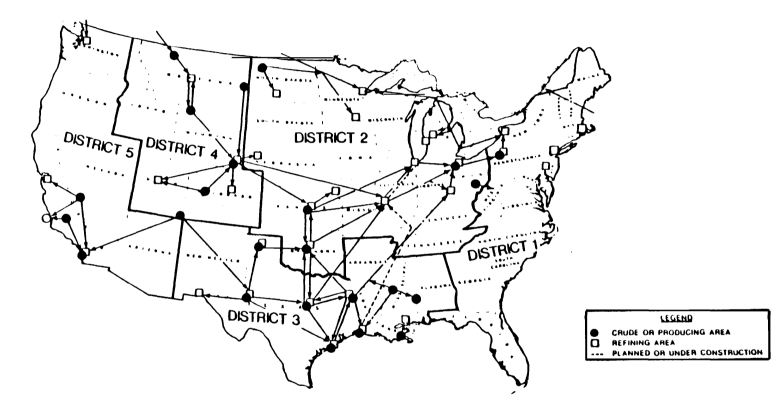
#### 2.9 Common Elements

There are about one hundred different processes for converting coal or oil shale to gaseous, liquid or solid fuels. Still there are important similarities among then. They relate to the physical size of the plants, their complexity costs, conversion efficiencies, and the requirements for resources such as manpower, feedstocks, land, water, and equipment. These factors are important for the understanding of the various conversion processes; the situation of constraints and time tables of implementation; and for understanding the uncertainties involved in projections.

### 2.10 Physical. Size

Larger sizes of synfuel plants do not cost propotionately more than smaller sizes. As a result there is an incentive to minimize costs by designing large capacity plants of the order of 50,000 to 100,000 barrels per day oil equivalent producti"on. Such large plants can provide the equivalent energy requirements of a city of about a quarter to half a million people. The investment required for a 50,000 barrels per day synfuel plant is estimated between \$2 and \$4 billion (Reference No. 28) . The constucti"on of the plant will be a major engineering endeavor and require about five to eight years. The size of the plant is measured in square miles<sup>7</sup>. The amounts of material that have to be handled by

<sup>7</sup> Two squares miles of land are required for a 125 million CFD gasification plant producing the equivalent of 22,000 barrels of oil per day (Reference No. 28).



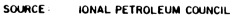


FIGURE 2.5: CRUDE OIL PIPELINE NETWORK

the conveion plants are also very large. In the case of a 50,000 barrel per day sRC II coal liquefaction plant, coal feed is estimated at 32,000 tons per day and solid wastes at 36,000 tons per day on a dry basis (Reference No. 25). This is equivalent to one railroad car every three minutes. In the case of a similar capacity oil shale retorting plant, the quantities are double those for the coal liquefaction plant (Reference No. 28).

### 2.11 Complexity

All of the synfuel processes reviewed in this paper have varying degrees of overall complexity<sup>®</sup> (Reference NOS. 28, 29, and 30). This is because many individual steps are required. However, although some complexity is unavoidable, redundant complexity is costly in terms of such factors as investment and operation costs, efficiency, lag time between initiation and finalization of projects, and reliability of operation. Rogers (1979) divided complexity into the following categories, which can apply to all coal and oil shale conversion processes, and comments on their implications:

- . "Reaction complexity. A process which requires several consecutive reactions is less desirable than a process involving fewer reactions. The sensitivity of any one reacti"on to changes in any of the important variables, such as temperature, concentration, etc., may have strong effect on quality control and reliability. Coal is a heterogenous material and imposition from a given mine often varies with time. This further aggravates the reaction blem.
- Operational complexity. A process with many steps which entails multiple handling of solids and fluid streams will be prone to more equipment failures and consequently greater downtime. As detailed in the section on reactor complexity, the methods used for gas/solid contacting and catalytic conversion can also greatly increase process complexity. As a general rule, solids cause more problems than fluids, and liquids are more troublesome than gases.
- Operating regime. The chemistry of coal conversion processes normally involves high operating temperatures and pressures. Very high pressures or temperatures involve more difficulties. Special materials and equipment such as high pressure solids feeders and non-standard items must be built and maintained

<sup>8</sup> Shale oil retorting and upgrading systems may not be as complex as some coal. conversion systems.

with much higher standards than required for simpler conditions.

• Auxiliary facilities. The number of *required support* facilities such as catalyst reclaiming, by-product recovery plants and special utility services will make the process complex. Each auxiliary service brings with it its own complexity factor with an influence on cost and reliability."

Comprehensive tables comparing the process complexity of various coal conversion processes have been published (Reference No. 29). It should be noted that many of the coal conversion and oil shale retorting processes share many common unit operations. These include such steps as grinding, drying, preheating, reaction, ash separation, flashing, hydrotreating, distillation, storage, and many auxiliary operations such as hydrogen generation, removal of sulfur and nitrogen compounds, waste processing, electric power production and plant maintenance. Many of these unit operations are familiar and can be designed with confidence. There are, however, a few steps which are either difficult or impossible to accomplish with known technology. They are the ones that add Uncertainty to synfuel technology with respect to costs and time tables.

### 2.12 costs

Synfuel plants are capital intensive. As discussed in Chapter 4, capital cost ranged significantly as a function of product cost over the technologies. This makes the plant cost estimate very important in any economic study. However, existing cost estimates of synfuel processes have many uncertainties. They are primarily due to uncertainties associated with unproven technologies, changing inflation rates, and wide fluctuations of primary energy prices. There are therefore wide fluctuations among economists, particularly with respect to feedstock rests, price of products, the capital investment needed to build the facilities, and the rates of return on investment.

one can expect that capital investment in first-of-a-kind (pioneer) plants is going to be higher, in equivalent dollars, than later plants designed and built with the benefit of operating experience for the process involved. 9

<sup>9</sup> 

Learning experience cost reductions can be very significant. An example applying to a rapidly emerging industry is the chemical and allied products industry, where real non-energy rests declined by nearly 3% yearly for more than two decades.

Larger size plants cost less per unit of product than smaller , plants . The relationships between capital cost and plant size is **Given** (Reference No. 28) by the equation: x 10 Capital Cost =ks where: k is a constant

S is a plant production rate

The exponent, x, is generally somewhere between 0.4 and 0.9, although usually between 0.6 and 0.8. The exponential rule" only applies for process plants which are similar in all respects except size. It is generally not applicable for situations where sizes differ by more than a factor of ten.

There are no quantitative estimates of anticipated cost reductions due to experience in building synfuel conversion plants. On the one *hand*, the immature and undemonstrated nature of many of the synfuel precesses suggests cost reductions when the industry will reach maturity. On the other hand, experience has shown that cost overruns in major projects utilizing uncertain technologies are frequent - and perhaps unavoidable occurrence. Exhibit <u>4-16</u> depicts cost growth in pioneer energy process plants.

#### 2.13 Conversion Efficiency

High conversion efficieny is an important factor to be desired. It affects not only the product costs and the conservation of resources, but also reduces undesirable health, environmental and socio-economic " impacts which are related to the quantities of needed feedstocks that need extraction, transportation, and processing and to the size of the plant . Efficiency is often defined as the ratio of the useful energy leaving the plant in the form of products and by-products to the energy in the input streams, including feedstocks and ancillary energy. In designing conversion plants, optimum efficiency is selected to give the least costly synfuel production. The calculations are relatively simple when applied to balancing of investments with consideration of savings expected on more efficient processes or equipment versus the costs of the investments. However, the calculations become very complicated when they take into account the energy balance of the plant and assume credits for the sensible heats contained in the feedstocks or the products. Since the price of coal, oil shale and many byproducts is relatively low on the basis of energy content (relative

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This exponential rule does not apply to multiple train systems.

to other economic factors) , there are economic constraints to increase absolute efficiency. Also in tires of crash programs, the pursuit of the most efficient design and equipment may need to be compromized to reduce delivery times of equipment and services.

Table 2.3 (Reference No. 29), summarized process efficiencies for various coal conversion processes. They range between 65 and 70 percent for the coal gasification and direct coal liquefaction processes. They are estimated to be between about 50 to 60 percent for indirect coal liquefaction. When gasification and electric power production are combined, the efficiency drops to about 40 percent.

#### 2.14 Other Requirements and Concerns

In addition to the above mentioned factors that characterize synfuel technologies (size, complexity, costs, conversion efficiencies. ..), there are additional requirements and concerns that the development of a synthetic fuels industry have. Among them are:

- . Labor Require ments. Large labor requirements, both during construction and for operation of the synfuel plants are typical. They are also closely associated with potential socio-economic impacts due to the relatively sudden increases in demand for services and resources\* These impacts are mainly influenced by the size of the demand for labor relative to the size of the communities involved. For oil shale facility development in the West, these potential impacts can be large (Reference Nos. <u>31</u>, <u>32</u> and <u>33</u>).
- . Feedstock Requirements. As stated earlier, the amount of feedstocks that are required for a generic conversion plant producing 50,000 barrels of oil per day equivalent are very large. About 30,000 -40,000 tons of coal per day are needed by a typical coal conversion plant and double that amount of shale by an oil shale retorting plant. The demand on coal feedstocks alone has been projected by the EPA (Reference Nos. 31, 32, 33) to increase from about .5% of projected U.S. coal output in 1985, to over 25% in 2000.
- . Land Requirements. As discussed earlier and specifically in Reference No. 31. the land requirements for synthetic fuel development includes not only the on-site land requirements of the physical plant, but also the land associated with extraction (mining) and with disposal.
- Water Requipments. As discussed earlier, and specifically in Reference Nos. 31, 32 t synthetic fuel plants require significant quantities of water. In the coal conversion industry, water is

2-19

Process	Developer	Primary <b>Product</b>	Secondary Product	Feed Coal Type/Size	Year Beyun	Conf. Index	• Efficency11 Process/Product	Renarks
<b>SRC - I</b> 12	Southern Co. Services + EPRI + DOE + Guif	Solid Boiler Fuel	Naphthe	All types	1962	B-3	71,⁄70	
6RC - 11	Gulf + DOE	Liquid Boiler Fuel	Gas LPG Naphtha	All types with ash restrictions	1976	₽-4	70/70	
eds.	Down	Liquid Boiler Puel	LPG Naphtha Gas	All acceptabl	3966	C-3	64/64	
H-Coal Fuel Oil H-Coal Syncrude	RAT	Fuel 011 Syncrude	Naphtha Gas			C-3	<b>/74</b> ∕69	
Fischer- Tropsch	Standard Tech- nology used M 50 years	Range of Hydrocarbons	LPG Alcohols NO. 2 011 Fuel 011 Gas	All coal is gasified	Before 1930	A-2	/48	Depends on gasifier efficiency
M-Gasolins	Mobil for Methanol to Gasoline Con- version	Premium Gasoline	LPG	Any gasifier to methanol process may be used		C-3	<b>22</b>	
Hethanol		Methanol		Depends on gasifier		A-2	/57	

Table 2.3: PROCESS SUMMARY

11 Efficiencies are highly dependent on product mix.

12 The SRC I process as presently designed for the SRC I precommercial demonstration plant would be a two-stage process which produces liquid fuels, as well as SRC solid. SRC II and EDS produce a distillate syncrude.

SOURCE: Reference 29

•

Procéss	Developer	Pristary Product	Secondary Product	Feed Coal Type/Size	Year Regun	Conf. Index	1 Efficiency Process/Product	Remarks
002 Acceptor	Conoco Coal Development Co.	High Btu Gas Low Btu Gas	None	Lignite or Sub-Bituminous /8x100 mesh	1968	B-2	<b>69/67</b>	Claimed to be the only fluid- ized bed process that can handle lignites
Hygaa	lgr	High Btu Gas	Haphtha	All with pretreat	1954	C-3	78/66	-
Bigas	Bituninous Coal Research	High Btu Gas	None	A11	1963	C-4	64/60	
Synthane	Builines PERC Lumnus	High Bru Gas	Char	All/20 to max 200-200	1961	C-3	65/63	-
Œ	Combustion Engineering	Low Btu Gas Electric Power	None	111	1974	C-4	/40	
Westing- house	Westinghouse	Electric Power	None	A11	1972	C-4	/38.4	Efficiency ahown with induction
		Medium Btu Gas	None			C-4	/83	
Lurgi	Lurgi	High Btu Gas	Tar, Oils Char Naphtha Coal Pines	Non-to low caking	Before 1930	A-3	72/59	Based on 1975 FPC application by ANG-analyzed by Stearns-Roger

Table 2.3: PROCESS SUMMARY (continued)

mainly used for hydrogen production, coding, waste disposal, and revegetation (Reference Nos. 31, 32, 33). In the case of the oil shale retorting industry, the main uses are for oil shale rein@, retorting, fuel upgrading, revegetation, and spent shale disposal (Reference Nos 8, 31, 33).

. Equipment Requirements. There are many kinds of equipment that will be required by the synthetic fuel industry. Among these items are:

For coal conversion: fabricated vessels, heat exchangers, rotating machinery, materials handling equipment, packaged plants, turbine generator sets, pollution control devices, piping, valves, and instruments and controls. The largest items are fabricated vessels, instruments and controls. They alone have been considered (Reference No. <u>34</u>) to amount for over 50% equipment needs.

For shale conversion: steel castings valves, air coolers, shell and tube exchangers, fired heaters, and boilers, preps, compressors, and pressure vessels, and tanks.

In the following chapters, we will look at these factors in more detail. Chapter 3 will discuss the individual process technologies; chap\* 4 will discuss the important assessment of costs; and Chapter 5 will discuss the projected deployment schedules of synfuel production. CHAPTER 3 : OVERVIEW OF SELECTED SYNTHETIC FUEL CONVERSION PROCESSES

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3.1	General Synfuel Processes	3-1
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#### Chapter 3: OVERVIEW OF SELECTED SYNTHETIC FUEL CONVERSION PROCESSES

3.1 General Synfuel Processes

The General term " synfuel processes" applies to the following:

- 1. Upgrading of coal. to gaseous, liquid or solid products with improved characteristics.
- 2. creversion of the kerogen in oil shale to gaseous or liquid fuels or products.
- 3. Recovery of petroleum crudes from non-conventional oil resources such as heavy oils and tar sands.

Upgrading of coal by subjecting it to a reaction with steam at high temperatures and pressures in the presence of air or oxygen, or to hydrogen, with or without a catalyst, is called conversion. The coal can be converted to gaseous (gasification) or liquid (liquefaction) hydrocarbons. The products have a much lower content of sulfur than the original coal. Oil shale can be retorted by subjecting it to high temperature and pressure, also producing gaseous or liquid hydrocarbons . Catalysts are used in synfuel processes when there is need to accelerate the reaction rates and affect the product state.

In this report, the following processes are included:

1. Coal gasification

-to medium Btu gas: generic - to high Btu gas: generic

2. Coal Liquefaction

by pyrolysis (none included)
 by solvent extraction: liquid solvent refined coal (SRC II)
 Exxon donor solvent (EDS)
 by catalytic liquefaction: H-coal
 by indirect liquefaction: Fischer-Tropsch (FT) Methanol

3. Oil shale retorting using:

true in situ retorting (none included)modified in situ: genericsurface retorting: generic

#### 3.2 coal Gasification

The process by which coal is gasified involves reactions of devolatization of coal with steam at elevated pressures and temperatures to produce CO and H<sub>2</sub>O. Gasification of coal involves basically the following reaction between steam and carbon:

c + Air or Oxygen +  $H_20$  co +  $H_2$  + Heat

There are many processes by which coal can be gasified producing low-, medium- or high-Btu gas. The definitions of the heat content of each of then are not rigorous. Low-Btu gas is a mixture of carbon monoxide, hydrogen and nitrogen It has a heating value of less than 300 Btu per standard cubic foot (Reference No. 25)=. This gas is of interest to industry either as a combustible fuel or as a raw material from which ammonia, methanol, and other compounds may be synthesized. Due to the low heating value, it cannot command high enough prices to justify long distance transport. Medium-Btu gas is a mixture of methane carbon monoxide hydrogen, and other gases. It has a heating value between 300 and 700 Btu per standard cubic foot (Reference No. 25). It is suitable as a fuel for industrial consumers, but because of its low heating value, is not economic to transport over great distances. High-Btu gas consists essentially of methane. It has a heating value of appro approximately 1000 Btu per standard cubic foot, and is compatible with natural gas in that it can be substituted for natural gas in existing pipeline systems.

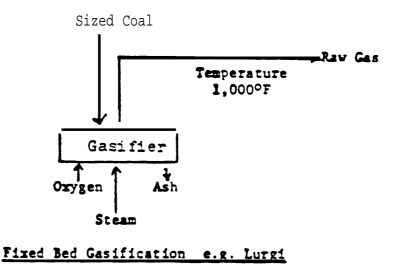
Coal gasification processes can be divided into three major process types\_according mainly to the way in which the feedstock coal, steam, and the product **gases** are **contacted**. They are:

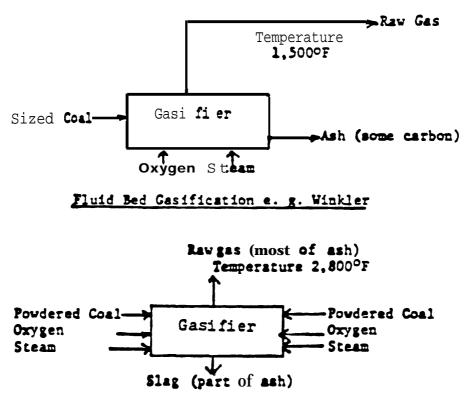
- 1. Fixed bed gasification in which the crushed, sized coal is fed from the top of the reactor vessel. Steam, air or *oxygen are* blown upwardly.
- Fluidized bed gasification in which the finely sized coal particles are "fluidized" by the steam, air or oxygen, which are piped through them.
- 3. Entrained bed gasification: in which the even finer coal particles are blown into the reacting gas stream prior to entry into the reactor. The coal particles are suspended in the gas phase, and are filtered and recycled until a product gas with a suitable heating value is produced.

Figure 3.1 (Reference No. <u>31</u>) describes the main features of these three **processes**.

1

Usually, low-Btu gas has a heating value below 200 Btu per SCf; and medium-Btu gas ranges in heating value between 300 - 350 Btu per Scf.





Entrained Gas ificat ione. R. Koppers-Tot zek

SOURCE : Reference 31

-.

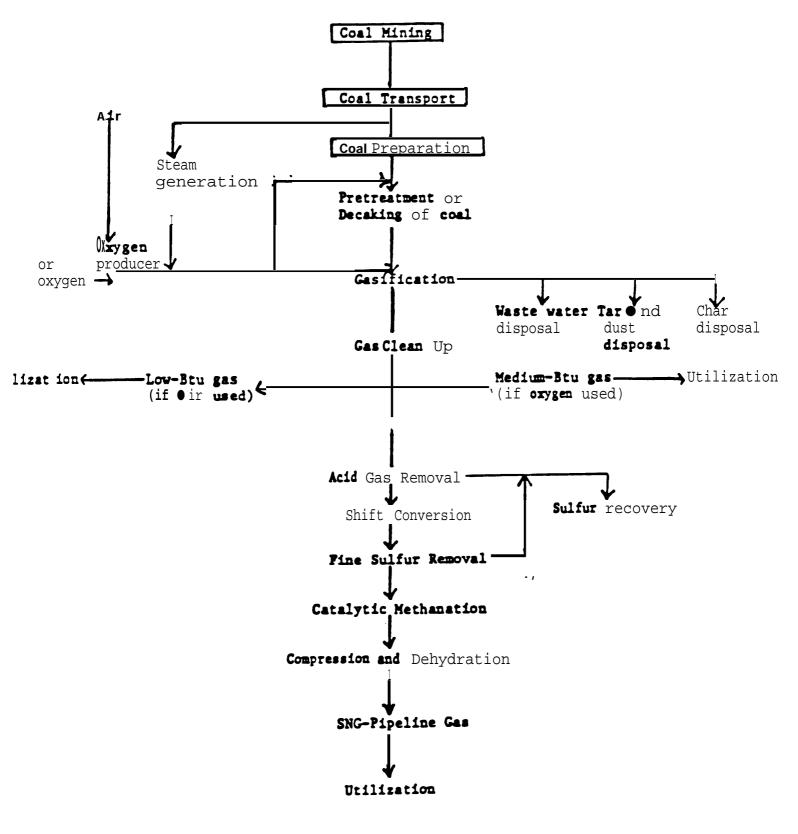
Figure 3.2 (Reference No. 31) is a schematic diagram of coal gasification. It represents the whole coal gasification fuel cycle, including the production of low-, medium- or high-Btu gas. All of these gasification processes share a number of process steps. If high-Btu, pipeline-quality gas is desired, essentially all of the following process steps are required. In some cases, some of them may be omitted, depending on the type of coal being processed and the type of gas product desired. The process steps are as follows (Reference No. 25).

- 1. Pretreatment of coal<sup>2</sup> (if sizing or caking are problems).
- 2. Primary gasification of coal.
- 3. Secondary gasification of carbonaceous residue from primary gasifier.
- 4. Removal of CO<sub>2</sub>, H<sub>2</sub>S, and other acid gases.
- 5. shift conversion for adjustment of the carbon monoxide/ hydrogen mole ratio to the desired 1:3.
- 6. Catalytic methanation of the carbon monoxide/hydrogen mixture to form methane.

#### Pretreatment

The coal received at the plant must be further cleaned and crushed or ground before it can enter the gasifier. Extaneous materials such as shale, rocks, metal, etc. are removed by conventional cleaning methods. For fluidized or entrained gasification processes, the coal needs to be finely ground. Crushing and sizing may also be required for other processes. In the case of certain bituminous coals called caking coals, agglomeration of the material is observed when they are heated. Treatment is needed if they are to be gasified by fluidized or Moving bed processes, or even in fixed bed reaction. The caking characteristics are destroyed when the coal is heated to low temperatures in the presence of air or oxygen.

<sup>2</sup> Pretreatment of coal by partial oxidation with air or oxygen is not in general a cost-effective approach to destroying the caking characteristics of certain coals, such as Eastern kiminous coals, because of the loss of Btu values of the coal in producing  $OO_2 \& H_2O$ . The caking problem is a serious problem in the processing of such coals and limits the applicability of current commercial gasifiers such as the dry-bottom Lurgi to Western Subbituminous coals and lignite.



#### Primary Gasification

This is the heart of the process, and is basically a pyrolysis process of the raw coal. The coal feed is contacted with synthesis gas (carbon monoxide and hydrogen). The coal is devolatized according to the following general reaction (Reference No. 25 ).

COAL + HEAT (Pyrolysis) + Methane, water, tars, phenols, hydrogen sulfide, hydrogen, carbon dioxide, char, etc.

The pressures used for gasification range from atmospheric pressure to 1000 psi. The heat required to maintain the endothermic gasification reaction is supplied from burning coal. Air or oxygen are also needed to support the combustion reaction. If air is used, the product is low Btu gas ranging from essentially a carbon monoxide/hydrogen mixture (Koppers-Totzek process) to mixtures containing various proportions of carbon monoxide, carbon dioxide, hydrogen, water, methane, hydrogen sulfide, nitrogen, and typical products of pyrolysis such as tar, oils, phenols, etc. If oxygen is used, medium Btu gas results.

The bulk of the original coal is transformed into a solid char. Certain coals are more "reactive" to gasification than others. Thus the type of coal being processed determines to a large "extent the amount of char produced, and the analysis of the gaseous products. The char is usually gasified by additional processing steps, or is marketed.

#### Secondary Gasification

Secondary gasification involves the gasification of char from the primary gasifier. This is usually done by reacting the hot char with water vapor to produce carbon monoxide and hydrogen.

If the desired final product is either low- or medium-Btu gas, secondary gasification is usually followed by scrubbing and cleaning. Carbon dioxide and sulfur compounds are partially removed, and the resulting gas is used directly. If high-Btu gas is desired, shift conversion and methanation are further required.

### Shift Conversion

In most gasification processes, a shift reaction is employed prior to methanation. Its-purpose is to react a portion of the carbon monoxide with steam to form more hydrogen.

 $co + H_{,O} + CO_{,2} + H_{,2}$ 

By this exothermic reaction the ratio of carbon monoxide to hydrogen may be increased to 1:3 mole ratio needed to produce methane. Otherwise, deactivation of the catalyst used in the methanation takes place.

The catalytic shift conversion reaction is a wellknown process, but it has not been applied on the large scale required for commercial coal gasification. For coal gas shifting, conventional iron-chromium catalysts may be used; however, the coal gas stream must be purified prior to shifting (Reference No. <u>25</u>).

### Methanation

If carbon monoxide and hydrogen are present in the mole ratio of 1:3, the coal gas can be reacted in the presence of a catalyst to produce methane. Group VII transition elements such as iron, cobalt, nickel, ruthenium, rhodium, palladium, osmium, iridium, and platinum have been found to be effective catalysts. The following exothermic reactions occur simultaneously within the methanation unit (Reference No. 25 ).

$CO + 3H_2$	<b>→</b>	$CH_4 + H_2O$
$CO_2 + 4H$	2 →	CH <sub>4</sub> + 2H <sub>2</sub> O
со + н <sub>2</sub> о	<b>→</b>	co <u></u> + ,2
2C0	+	<b>co, +</b> c

Special care must be taken to prevent deactivation of the catalyst by temperatures above 750°F. It can also be Poisoned by carbon deposition. These can be circumvented by ensuring that the mixture of carbon monoxide and hydrogen shall be fed to the methanator in the ratio of 1:3. Scrubbing of sulfur from the synthesis gas feed is employed to alleviate sulfur poisoning of the catalyst.

The final step to prepare high Btu gas for marketing is to remove water to specified levels. The product gas usually undergoes compression prior to storage *or* marketing.

## 3.3 Coal Liquefaction

### 303.1 General

Coal liquefaction processes are conversion processes in which liquids are the primary products. Some gases and solid char may also be produced.

There are two basic routes to coal liquefaction, namely direct and indirect liquefaction. In direct processes, slurried crushed coal is reacted directly with hydrogen at high temperature and pressure conditions to produce liquid hydrocarbons. In indirect liquefaction, coal is first gasified to produce a hydrogen-and carbon monoxide mixture. Further recombination with the aid of a catalyst produces liquid products.

Direct liquefaction is further broken down into three generic processes, namely: pyrolysis, solvent extraction, and catalytic liquefaction. The yields and physical properties of the produced liquid products depend directly on the reactor conditions and degree of hydrogenation.

### Pyrolysis

In pyrolysis processes, coal is heated to temperatures above 750°F. It is converted into gases, liquids, and char. The latter accounts for more than 50 percent of the weight of the feed coal and requires hydrogenation. Some amount of solids remain in the raw gas and liquid products. They consist of unreacted coal and ash, and can be relatively easily removed from the gas stream. But the liquid requires filtration, distillation, or some other treatment to remove the solids.

### Solvent Extraction

This process makes use of coal derived liquids known as "donor" solvents to increase the fraction of the coal that goes into solution. The "donor" solvents act as source of hydrogen to the coal products, and are reacted together at temperatures up to 950°F. Hydrogen may be supplied under pressure in the extraction step, or it may be used to hydrogenate the solvent prior to recycle. In some processes the unreacted coal is used to generate the necessary hydrogen. In other processes, the hydrogen is generated from by-product gases or from additional raw coal.

#### Catalytic Liquefaction

In this process, pulverized coal is mixed with 1-1.5 parts of recycle solvent. A suitable catalyst is used to add hydrogen. Most precesses of this type operate in the liquid phase with catalyst dispersed throughout or in a fixed bed. Some processes now in the development stage involve the injection of catalyst-impregnated coal into a stream of hot hydrogen at about 950° F for a very short time (Reference No. 25).

#### Indirect Liquefaction

Two stage conversion of coal typifies indirect liquefaction processes. Coal is first reacted with steam and oxygen to produce a gas composed primarily of carbon monoxide and hydrogen. This gas stream is subsequently purified to *remove sulfur*, nitrogen, and ash. The product gas is then catalytically reacted to yield liquid hydrocarbon products.

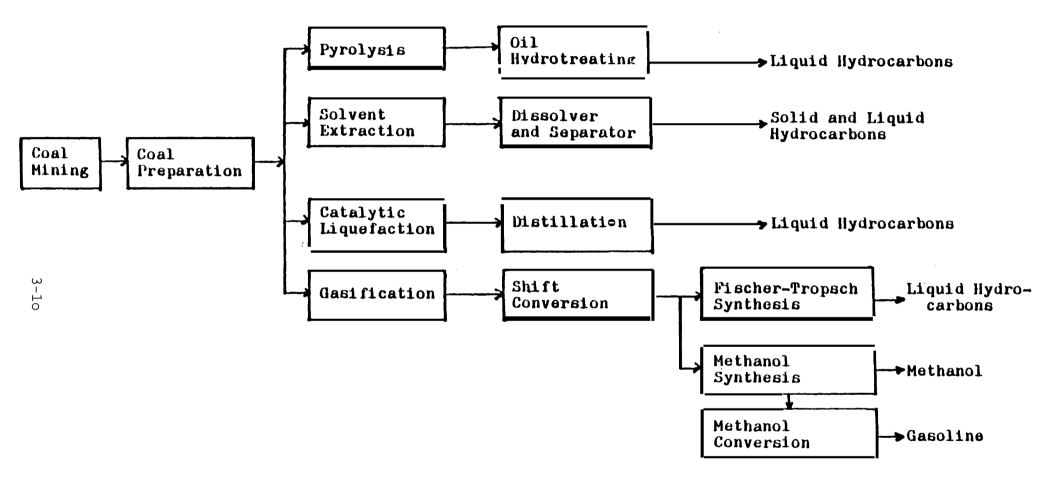
Figure 3.3 (Reference No. 31) presents a schematic diagram of the basic liquefaction processes. Each of them produces several types of products and same gas, which may be used within the plant.

Removal of solids from coal liquids is a critical step in most of these liquefaction processes. Although there is currently a trend toward elimination of the solid-liquid separation step by the recovery of a solids-laden vacuum bottoms stream for gasification, most existing plant designs call for some type of physical/chemical solids removal systen. 3 The three processes receiving the most current interest are critical solvent deashing, antisolvent deashing, and pressure filtration (Reference No. 25).

Separation of ash and unreacted coal particulate from coal liquids is difficult because of the small size and large quantity of the solid particles, the snail density difference between solids and the liquid, and the high viscosity and melting point of the liquids. The Kerr McGee Corporation has been developing a separation technique which utilizes solvents such as benzene, toluene, xylene, pyridene, and cresols near their critical temperature and pressure, hence the term solvent deashing (Reference No-. 25) .

3

Solid\liquid separation is a critical step only in direct liquefaction process. Most modern coal hydroliquefaction processes in the pilot plant stage of development, such as SRCII, EDS, H-Coal (syncrude rode) do not require a solid/liquid separation stage.



\* Only major products are shown. The last process represents indirect liquefaction. SOURCE: Reference 31

## 3.3.2 Liquid Solvent Refined Coal (SRCII)

The SRCI process was developed to convert high-sulfur, high-ash coals to low-sulfur and ash solid fuels. The SRCII is the same kind of process, except the product is a liquid rather than a solid. This is achieved by adding more hydrogen through the following steps:

- 1. Recycling of a portion of the product slurry as solvent for the feed coal.
- 2. Higher residence time in dissolver.
- 3. Higher pressure.
- 4. Use, of vacuum distillation to separate solids from liquid, rather than the troublesome filtration step employed in SRCI .

Figure 3.4 is a schematic diagram of the SRCII process (Reference No. 35) . Table 3.1 summarizes the components, resource requirements, and potential impacts of this process (Reference No. 17). The feed coal is first pulverized to less than 1/8" size, dried and mixed with process derived solvent in a slurry mix tank (Reference No. 35) . Feed coal is limited to those containing certain trace mineral elements which may be required to act as catalysts for the breaking of solids to liquids in the liquefaction reaction<sup>4</sup> (Reference No. 291. However, in cases where the problem is concentration rather than the presence of specific trace elements, a recycle of residue may broaden the allowable coal feeds (Reference No. 29) . The coal slurry is then mixed with hydrogen generated by gasification of the vacuum bottoms from the liquefaction step and reacting with steam and oxygen in a gasifier-converter. The slurry is pumped through a preheater (700 to  $750^{\circ}$ F) and passed through a dissolver (2000 psi, 820 to  $870^{\circ}F$ ) to dissolve about 90 percent of the coal (Reference No. 35) . The following additional reactions take Place in the dissolver (Reference No. 35) .

- 1. The coal is depolymerized and hydrogenated.
- 2. The solvent is hydrocracked to form lower molecular weight hydrocarbons, ranging from light oil to methane.
- 3. Much of the organic sulfur is removed in the form of hydrogen sulfide.

The *sultry stream* from the dissolver is split into two. One is recycled to provide solvent for coal slurry mixing. The other is fractionated to recover the primary

<sup>&</sup>lt;sup>4</sup> Opinions differ about the role of the trace minerals as catalysts. The primary "catalyst" in the SRCII process may well be the pyritic mineral matter contained in the coal and not "trace mineral elements. "

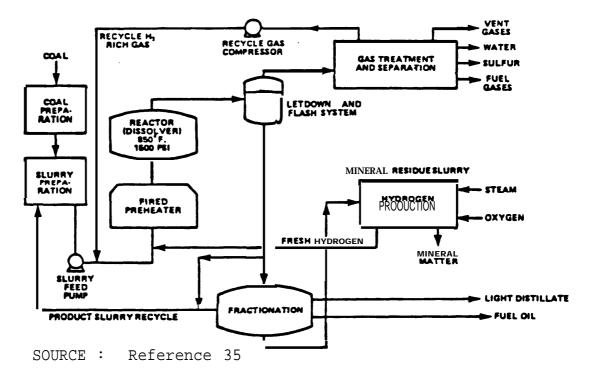


Figure 3.4

A SCHEMATIC DIAGRAM OF THE SOLVENT REFINED COAL (SRC-II) PROCESS

TA3.E 3.1					Solvent Refined Coal II
(400-1 \$13110) 	RESNURCES USED: (Por 10 <sup>13</sup> Bin Produced)		COMPARENT ON STORES		
112E1 • 0.000 bal/day • 135 = 1012 Buv/day (2) • 01 Plant availability (2) • 01 Plant availability (2)		Nput 40,919 tunn 12,821 ftu/lb	AIR PULLITANTS <sup>(1)</sup> particulaten 502	• • • • •	
011:022 ± 1014 Bir/yoor(2) 011:022 toos page PDM cosi(0) • eiticiory 42(0)	COAL ANALYSIS(DAF) <sup>(a)</sup> antature	L . by welght)	MAT Ny Serie ar buna 10	• e " " "	
• plant 116 30 yange(0) • 6.7 ± 104 biu/bal(3)	volatile matter fined cathon cultur	15.09 01.12 01.12	ar sealc radalua assiste		
peschiftion • Call la crushed to about 200 meah and	1	1.1	t i nc c'hr celue		
dried to three percent melature and electric during elevent. A 70 to 53	LAND <sup>(C)</sup> plant facilities	Acres 1.5	ni.hel lead		
Portent Nydrogen Bisture in Addad. Nie Bisture in firet prehened, then Vydromonied in the firet of the			polyguelear organic material		
temperature 14 415 to 455 C; preserve 68 to 136 atmospheres . Liquefaction	WATER <sup>(c)</sup> cooling towers	Acro-Fr.	WATER POLLUTANTS no direct discharge to		
produces a minture of maser, vapor. liquids, and polids. The mases and	plant use steam	0.11	any watercourse		
version are experted from the liquide and solids and the verse condensed.	alacel laneoua total	0.9 5.5	SOLID WASTE <sup>1,1</sup> dry tom equivalent	1010 (11)	
the figures to experised from the mailed by filtration (BAC 11).	00515 construction	48	EN MAY PHONKE(2) Avectude (581-11)	<u>Equivolont Barrolo</u> 149,254	
	eperal lon	4	(+nergy +ontent = 6.) x 10 <sup>6</sup> Btu/bbi)	•	
CCCPCONENTS • CTURNOTS	PERSUMMEL construction	<b>N</b> A			
e BEC roector e dering weit	operation & minimumer	4			
e pre-bastere e gas recevery plant					
filtering w					
LIV (AD-WEATAL LOWCENS - a strondations - a strondations - a cold water - a cold - a co					
"The fait presented are based as a conceptual dealy.	a of a commercial facility.				
evaluate. The data should not be used directly for comparison with achor coal liquifaction processes. (1)After coal proparation tectudes solid varis from also burial and tailings pond. (2)Aspressiative values subscred for analysis purposes.	er competions with other con mine buriel and teilings po ores:	l liquefaction proc			
800K (a) Hittena Associates, Jac., Standarda of Practice Namual for Sulvent Belland Coal Liguelaction Procesa, 1918. b) Hittena Associates, Jac., <u>Fritonental Antennent of Coal Liguelaction</u> , 1918. (c) Two, <u>Chartestationa and Deta Ia the Area of Guil Neur Environmential Data Book</u> , Volume IV, 1918. (d) Plant and Pryor, <u>Boiler Fuel and Building Roces</u> , 1978.	<u>rectice Namual for Sulvant B</u> Ancenent of Gal Ligueleci Lee of fuel Reve Environment A floods, 1918.	eflace Coal Liquefa Lon, 1918. el Data Book, Volum	<u>cilum Fracewa</u> , 1918. + 1V, 1918.		
SoURCE: Reference 17					

products which consist of naphtha, low sulfur fuel oil, and a vacuum residue which is separated from the solution in a filtration unit. The residue consists of heavy oil, ash and undissolved organic material from the coal (Reference No. 25).

The gases from the dissolver are treated to remove hydrogen sulfide and carbon dioxide. Liquid petroleum gases and pipeline gas are separated in a cryogenic separation unit. Unreacted hydrogen is recovered and recycled.

Recent developments have resulted in increased efficiency of the SRCII process. A combination of solid and liquid products are produced. A wide range of products can be obtained depending on the severity of recycling. Table 3.2 (Reference No. 25 ) shows the properties of a typical mix of products.

### **3.3.3.** Exxon Donor Solvent (EDS)

The process is similar to SRCII, except that the major portion of the hydrogen supplied as part of the solvent is chemically combined rather than in the form of a free dissolved gas (Reference No. 29). A schematic diagram of the process is illustrated in Figure 35 (Reference No. 35 ). Crushed coal is liquefied in a reactor at 800-880°F and 1500 - 2000 psig (Reference No. 25 ). The reaction is non-catalytic, in the presence of molecular hydrogen and the hydrogen-donor solvent, which transfers hydrogen to the coal. The product from the liquefaction reactor is separated into two portions. One part is sent to the solvent hydrogenation unit to produce donor solvent. It is a catalytically hydrogenated recycle stream which is fractionated from the middle boiling range of the liquid product, and has a boiling range of 400 - 850°F (Reference 25 ). After hydrogenation, the solvent is mixed with No. fresh coal feed, heated in a furnace, and pumped into the liquefaction reactor.

The other portion from the product liquefaction reactor is a slurry. It is separated by distillation into gas, naphtha, middle distillate, and a bottom product that contains heavy liquid, untreated coal and mineral matter. The vacuum bottoms slurry is cooked to produce additional liquids.

The major advantages of the EDS process are:

1. High yields of low sulfur liquids are obtained from bituminous and sub-bituminous coals or lignites (Reference No. 25 ). A yield

## TYPICAL PROPERTIES OF SRC FUELS USING RECYCLE SRC II PROCESS

	<u>Solid Fuel</u>	<u>Distillate Fuel</u>
Gravity: `API	-18.3	5.0
Approximate Boiling Range: `F	800+	400-800
Fusion Point: 'F	350	
Flash Point: `F		168
Viscosity: SUS at 100°F		50
Sulfur*: Percent	0.8	0.3
Nitrogen*: Percent	2.0	0.9
Heating Value: Btu/lb.	16,000	17,300

\* Assuming Western Kentucky coal feed with 4% Sulfur and 2% Nitrogen. SOURCE: Reference 15

3-15

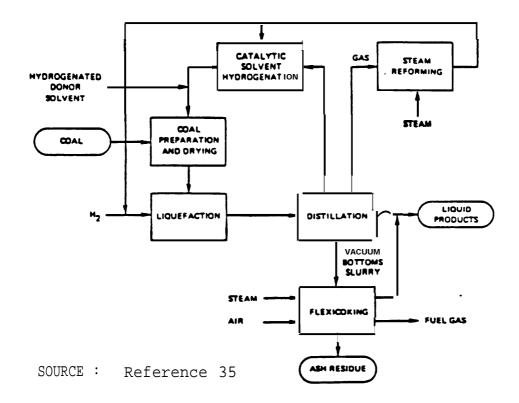


Figure 3.5 A Schematic Diagram of the Donor So I vent Liquefaction Process

of 2.6 barrels of liquids per ton of dry coal is typical for an Illinois bituminous coal (Reference No. <u>25</u>).

- 2. The only by-products of significance are ammonia and elemental sulfur (Reference No. 25).
- 3. There is wide flexibility in product distribution by varying liquefaction conditions or adjusting solvent properties (Reference No. 25).

The typical properties of the products from the EDS process are shown in Table <u>3.3</u> (Reference No. <u>25</u>]. An estimated heat balance is given in Table <u>3.4</u> (Reference No. <u>35</u>).

#### 3.3.4 H-Coal

The H-coal process converts coal to hydrocarbon liquids by hydrogenation with a cobalt-molybdenum catalyst. An ebullated bed reactor is employed. The liquid products may range from a heavy boiler fuel to a synthetic crude product (Reference No. 25 ).

Figure 3.6 (Reference No. 35 ) is a schematic diagram of the H-coal process. Coal is first crushed to minus 60 mesh, dried, and then slurried with recycled oils at pressures of approximately 200 atmospheres (Reference No. 25) " Mixing of the slurry with compressed hydrogen follows, and the mixture is preheated. The material is pumped to the bottom of the ebullated bed reactor, with the-upward flow of slurry through the reactor maintaining the catalyst in a fluidized state (i.e. random motion). The catalyst needs periodic additions of fresh catalyst and withdrawals of spent portions. Typical temperatures of the slurry entering the reactor are 650 - 700°F (Reference No. 25 ). The finely divided coal and ash particles flowing through the ebullating bed are removed with liquid and vapor products.

The reactor effluent is separated into recycle and net product streams. Conventional processing equipment is used. The liquid stream is distilled to produce a mixture of light distillate and a heavy distillate product. Gaseous products composed of hydrocarbon gas, hydrogen sulfide and ammonia are separated. A portion of the heavy distillate is recycled as the slurrying medium.

The operating conditions of the H-Coal process can be altered to produce various types of primary products. For

## DONOR SOLVENT PRODUCT ANALYSES

	Heavy	Naphtha~	200°C	+ Fuel Oil
	Raw <u>Liquid</u>	Hydrotreated Liquid	Raw Liquid	Hydrotreated Liquid
Nominal Boiling Range, `c	70/200	70/200	200/540	200/540
Distillation, 15/5°C				
10 wt. % 50 wt. % 90 wt. %	106 180 199	92 157 182	247 368 433	239 347 412
Density $(g/cm^3)$	0.87	0.80	1.08	1.01
Elemental Analysis, Wt. %				
c H O N s	85.60 10.90 2.82 0.21 0.47	86.80 12.90 0.23 0.06 0.005	89.40 7.70 1.83 0.66 0.41	90.80 8.60 0.32 0.24 0.04
Higher Heating Value MJ/kg	42.6	44.9	39.8	42.1

<sup>1</sup>Excludes C<sub>6</sub>/70°C naphtha cut

SOURCE: Reference 25

### Tab I e 3.4

Estimated Heat Balance for a Commercial Scale EDS Plant

<pre>System Products Liquids Sulfur, ammonia System Losses Ash, combustibles and sensible heat Stack losses Energy losses via water and air Liquefaction and solvent hydrogenation (9.80%) Flexicoking (6.44%) Hydrogenation and recovery (6.72%) " By-product recovery, offsites, and miscellaneous (3.18%)</pre>	Btu/day (10 Btu's) 323,071 8,309	Percent of Total Energy Input 61.72 1.59
Liquids Sulfur, ammonia <u>System Losses</u> Ash, combustibles and sensible heat Stack losses Energy losses via water and air Liquefaction and solvent hydrogenation (9.80%) Flexicoking (6.44%) Hydrogenation and recovery (6.72%) " By-product recovery, offsites,		
Sulfur, ammonia <u>System Losses</u> Ash, combustibles and sensible heat Stack losses Energy losses via water and air Liquefaction and solvent hydrogenation (9.80%) Flexicoking (6.44%) Hydrogenation and recovery (6.72%) " By-product recovery, offsites,		
Ash, combustibles and sensible heat Stack losses Energy losses via water and air Liquefaction and solvent hydrogenation (9.80%) Flexicoking (6.44%) Hydrogenation and recovery (6.72%) " By-product recovery, offsites,		1.00
heat Stack losses Energy losses via water and air Liquefaction and solvent hydrogenation (9.80%) Flexicoking (6.44%) Hydrogenation and recovery (6.72%) " By-product recovery, offsites,		
Other miscellaneous	26,082 20,039 136,853 8,309	5.13 3.83 26.14 1.59
Energy Input	0,000	
Coal (cleaned)* Electrical power**	488,761 34,702	93.37 6.63

\* Coal - Illinois No. 6; 10,574 Btu/Ib as received prior to cleaning

\*\* Power based on 8,500 Btu/kwh to generate
 SOURCE: Reference 35

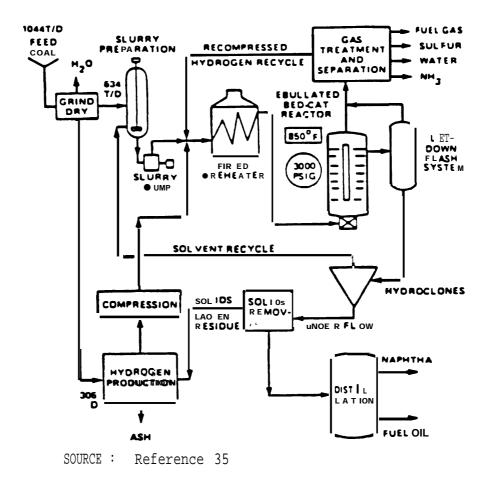


Figure 3.6

SCHEMATIC DIAGRAM OF THE H-COAL PROCESS example, relatively high temperatures and high hydrogen partial. pressures are used to produce a synthetic crude products. Vacuum distillation is used to separate the solids from the liquid phase. If gas and oil are desired, lower temperatures and pressures are used (Reference No. 25). Conversion and yield structure are determined by reactor conditions, catalyst replacement rate, and recycle slurry oil composition (Reference No. 29).

Table 3.5 (Reference No. 25) summarizes the properties of both the fuel oil syncrude products from H-coal.

Table 3.6 (Reference No. 17) summarizes the components , **resources** and potential impacts from H-coal process. It requires between 14,000 and 20,000 standard cubic feet of hydrogen for each ton of coal produced. Hydrogen consumption depends on the type of product produced, with less hydrogen required during the production of residual oil (Reference No. 25).

#### 3.3.5 Fischer-Tropsch Process

A commercial plant using a modification of this process is currently operaing in South Africa (Reference No. 36). This is the **only commercial sized** plant producing synfuels. Table 3.7 (Reference No. 35) is an overview of this plant.

In the Fischer-Tropsch process the coal is initially gasified (for description of gasification see section 3.2 of this report). The synthesis gas is then converted to largely aliphatic hydrocarbons using an iron or cobalt catalyst.

Figure 3.7 (Reference No. 35) is a schematic diagram of the SASOL I plant, which utilizes the Fischer-Tropsch process. Thirteen high pressure gasifiers convert coal in the presence of steam and oxygen to medium Btu gas containing mainly carbon monoxide, tars and oils. The product gas is then cleaned of carbon dioxide, hydrogen sulfide, organic sulfur, ammonia, and phenols. The cleaned gas is then subjected to the catalytic Fischer-Tropsch reaction which produces a mixture of gases, liquid hydrocarbons, and an aqueous chemical mixture that must be further processed to set the desired plant output .

The cleaned gas from the Lurgi gasifiers is partitioned into two streams. One stream is reacted in a fixed bed catalytic reactor to produce straight chain and medium boiling oils, diesel oil, LPG, and some alcohols. Operating conditions are 450°F and 360 psig (Reference No. 35). The other stream is combined with reformed product gas to increase the hydrogen to carbon ratio. It is reacted in a fluidized bed reactor

### PROPERTIES OF H-COAL DISTILLATES FROM ILLINOIS NO. 6 COAL LIQUIDS

	Fuel		Syncr	rude
Property	<203°C distillate	>203°C distillate	<197°c distillate	>1970C distillate
Specific gravity, 60°/60°F	0.864	0.979	0.838	1.025
Gravity, `API	32.3	13.0	37.4	6.6
Pour point, ASTM D-97, F	<5	<5	<5	<5
Color, ASTM D-1500 or (BuMines description)	NPA6	Brownish black	NPA4-1/2	Brownish black
Kinematic viscosity @ 100°F, ASTM D-455, C <sub>s</sub>	1.08	3.87	0.96	14.90
Saybolt viscosity, SUS, 100°F		39		77
Sulfur (Bomb) ASTM D-129, wt-pet	0.13	0.29	0.06	0.35
Nitrogen, Kjeldahl, Wt-pet	0.420	0.446	0.212	0.871
Carbon residue (Conradson) ASTM-524, Wt-pet	0	2.33	0	5.44

SOURCE: Reference 25

ENERGY SYSTEM	RESOURCES USED: (Per 10 <sup>84</sup> Btu Froduced)		RESIDUALS AND PRODUCTS: (Per 10 <sup>14</sup> Blu Produced)		
5126 • 39,000 bb1/day					
e .32 x 1012 Bcu/day	MIL.		AIR POLLUTANTS	Ione I.I	
e 105.12 m 1012 Bru/year	Coal: bituminews	57,756 tone	perticulates		
e 901 plant aveilability	energy content	11,900 Bcu/1b	\$02	20.2	
o operates 328.5 days/year	••		MO <sub>R</sub>	40.9	
e efficiency 701	COAL ANALYSIS	I (by weight)	hydrocarbone	0.6	
a plant 15fa 30 years	solsture	2.7	co	NA.	
	volatile matter	35.5	• reants	0.1	
DESCRIPTION	fixed carbon	51.7	cada lua	1.1 = 10-2	
e Coal to crushed to about 3/4 Inches	pulfur	3.0	BOTCUTY	6.2 u 10-3	
and then ground to stave 60 nesh and	dee	7.1	• Ibr	5.9 m 10-2	
mixed with recycled oil. The elurry			chronium	0.1	
to them prohested and fed to the	END(1)	Acres 3.9	Ich*i	3.3 a 10-3	
reactor and hydrogenated as it comes	plant facility		lead	1.1 m 10-2	
in contact with the abulatting bad of	solid wasts disposal	0.31	polynuclear SSMIC		
catalyst. Beaction temperature is			material	MA	
about 350°C. Games and vapors are	CONSUMPTIVE WATER USE	Acro-PL. 9.3			
them withdrawn from the top of the	process water		WATER POLLUTANTS		
reactor and peased through condensars	cooling water	101.6	no direct discharge late of	y welst source	
for product separation. The heavier	wante wilde	7.6			
product is withdrawn as a side-stream	potabla	.1	POLID WATE())	<u>10me</u> 3,203	
from the reactor and pumped to an	other (utility, queach, atc)	121.8	dry too equivalent	5,705	
stmosphere distillation unit.	Total	121.0			
		A-11 (1075)	BUBLCT PRODUCT	Berrele	
CONFORMERTS	<u>costo</u> (2)	Bullare (1973) 6,200,000	liquid fuel	154,250	
e crushere	CONSTRUCTION	3,700,000	(energy content Stu/bbl)	6.4 x 10 <sup>0</sup>	
e drying unit	operation.	1,700,000			
e hydrogen plant (ganifier)		Nachara			
a H-coal reactor	PENN SAMEL	<u>Morkere</u> NA			
e gas recovery plant	construction	4.1			
a distillation unit	opetation & maintenance	9.1			
BUTT BOUNDERTAL CONCERNE					

• • ir entestone

· solid veste

water pollution from runoff and leaching
 occupational basards and basith offects

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"The data presented are based on a conceptual design of a commercial facility. The data will be updated when more current data become available. The data should not be used directly for comparison with other coal liquefaction processes.

(1) This represents land committed to use ever the lifetime of the plant, divided by the <u>annual</u> output of the plant, expressed in trillion Btu., (405 acres = 105.12 = 3.9 acres) (2) This represents total cost of constructing the plant, divided by the <u>annual</u> output of the plant, appressed in trillion Btu., (4650 million = 105.12 = 56.2 million). (3) Does not include solid weats from aims burial and tailings pond.

BOURCES: Rittman Annocisten, Inc., <u>Environmental Annosoment of Coal Liguefaction</u>, 1978. TBM, <u>Characterisations and Data in the Area of Coal Neas Invironmental Data Book</u>, Volume IV, 1978. U.S. Energy Research and Devisionent Administration, <u>Synthetic Liguid Yueis Devisionent: Assessment of Critical Pectors</u>, BRDA 76-129/2. Fluer Engineers and Constructors, Inc., <u>N-Coal, Commercial Critical Pectors</u>, March 1976.

SOURCE: Reference 17

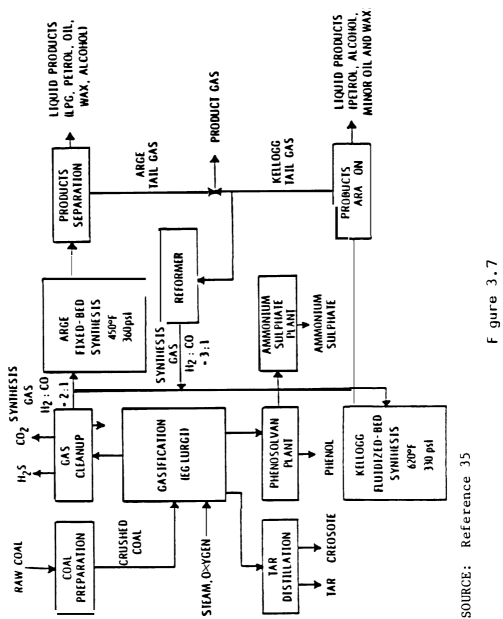
ω

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#### **H**-Coal

<b>Overvi</b> ews on	SASOL I and SASOL II, based on reference 8, follow:					
	SASOL 1					
LOCAT I ON:	Sasolburg, South Africa					
DESCR   PTION:	Gasification in Lurgi gasifiers					
	Two Fischer-Tropsch synthesis units;					
	<ol> <li>ARGE fixed-bed unit, temp. 230°C; press.23 atm.; catalyst, pelleted precipitated iron.</li> </ol>					
	<ol> <li>Kellogg SYNTHOL process, hlgh- velocity entrained-flow reaction using a doubly promoted iron catalyst.</li> </ol>					
SIZE:	10,000 bpd					
STATUS:	in commercial production since 1956					
YEARS OPERATION:	24					
COAL TYPE:	Subbituminous					
MAJOR PRODUCTS:	Liquid fuels, chemicals, and fuel gas.					
	SASOL II					
LOCATION:	Secunda, South Africa					
DESCRIPTION:	Gasification in Lurgi gasifiers,					
	Fischer-Tropsch synthesis unit using the Kellogg <b>SYNTHOL</b> process					
SIZE:	Nominal 40,000 bpd					
STATUS:	Anticipate ready for commissioning in 1980					
COAL TYPE:	Subbituminous					
MAJOR PRODUCTS:	Liquid fuels (gasoline is the major product).					
SOURCE: Reference	e 35					



Flscher-Tropsch Synthes s

3-25

at 620%' and 330 psig, (Reference No. 35). The main products are gasoline, fuel oil fractions, and various chemical products. The gasoline has a lower octane rating than the one derived from petroleum crude. The products produced do not fit well into existing markets. However, Mobil Oil Corporation has developed catalysts that improve the quantity and quality of gasoline (Reference No. 29).

### 3.3.6 Methanol Process

The production of methanol from synthesis gas is a specialized application of the Fischer-Tropsch reaction. Whereas the F-T process produces liquid fuels and chemical products, the Mobil methanol process produces gasolines . The schematic outline of this process is given in Figures 3.8 and 3.9 (Reference No. 35) . Table 3.8 '(Reference No. 35) presents a comparison of the thermal efficiencies of the Fischer-Tropsch and the Mobil methanol-to-gasoline process.

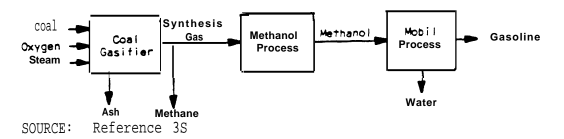
In the Mobil methanol liquefaction process, synthesis gas is produced from coal by any of the mediun-Btu coal-gasification processes. The synthesis gas is converted to methanol by a number of catalytic processes. The reaction is exothermic. The yield of methanol is optimized by using high pressures and low temperatures, optim um type and shape of catalysts, and of recycling of the unreacted gases.

The conversion of methanol to gasoline is a separate catalytic conversion process. The Mobil conversion process dehydrates methanol, then rearranges the carbon and hydrogen atoms. The zeolite catalysts employed in the process (called ZSM-5 class catalysts) have a unique channe The pore openings are of the right size to limit the size of the product molecules that can pass through then. the conversion proceeds to conventional high quality gasoline Reference No. 25 ).

Table 3.9 (Reference NO. 25) summarizes the overall material and energy balances of the methanol-to-gasoline conversion process.

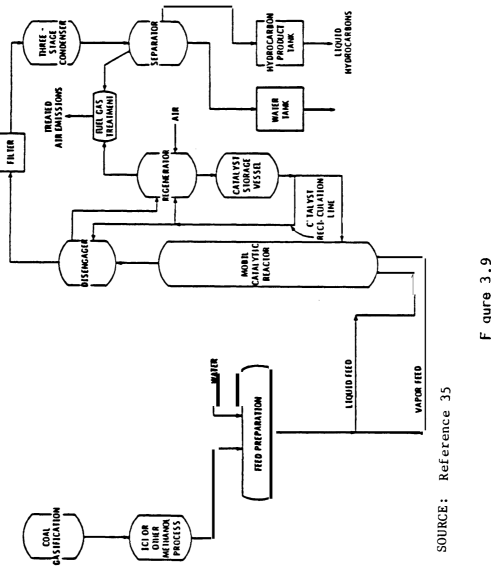
Table 3.9 (Reference No. 25) shows typical product yields produced from methanol by this conversion process.

<sup>5</sup> Even though no commercial demonstration plants of the "indirect" coalmethanol-gasoline process has been built as of this date, this route is considered by many authorities to be a very promising way to get gasoline from coal. There are several proposed studies and plants under instruction in the U.S. using this process (see Appendix chart). Also, New Zealand Liquid Fuels Trust Board (Report No. LF 5502, 10/31/79) has a large Mobil-M gasoline plant under construction (expected to become operational by 1983-5).









3-28

### Table 3.8

### Thermal Efficiencies

	<u>Methanol-t</u>	co-Gasoline <sup>7</sup>	Fische	$r-Tropsch^{7}$
	Btu/hour (1C <sup>°</sup> Btu)	Percent of Input	Btu/hour (106 Btu)	Percent of Input
Input				
coal <b>Coal</b> Fines (excess) Methanol Total Input	<b>19,383</b> (872) 18,511		19,708 <u>3</u> 19,711	
output				
<pre>SNG C3 LPG C4LPG 10 RVP Gasoline Diesel Fuel Heavy Fuel Oil subtotal</pre>	6,067 247 385 4,689 11,388	32.8 1.3 2.1 25.3 61.5	7,243 176 26 2,842 514 147 10,948	$36.8 \\ 0.9 \\ 0.1 \\ 14.4 \\ 2.6 \\ 0.7 \\ \overline{55.5}$
Alcohols sulfur Ammonia Power Total Output	19 83 18 11,508	$ \begin{array}{c} 0.1 \\ 0.5 \\ 0.1 \\ \overline{62.2} \end{array} $	290 19 83 <u>11</u> 11,351	$   \begin{array}{r}     1.5 \\     0.1 \\     0.4 \\     0.1 \\     \overline{57.6}   \end{array} $

<sup>6</sup> Thermal efficiencies are highly dependent on product mix.

7 The indirect liquefaction processes shown here may be Considered as gasification processes for SNG, with the major coproduct being galosine, e.g., for the "Fischer-Tropsch process" shown, the yield of SNG is 1.45 BOE/ton of coal, with a gasoline yield of 0.58 BOE/ton of coal. It is thus not representative of the SASOL-II process which emphasizes the production of liquid fuels.

<sup>8</sup> Direct thermal equivalent value (thermal efficiencies are highly dependent on product mix (see Section 7. 5).

SOURCE : Reference 35

### TABLE 3.9

### METHANOL-TO-GASOLINE BALANCES

	<u>Methano</u> l →	<u>Hydrocarbon</u> s +	Water
Material Balance	100 tons	44 tons	45 tons
Energy Balance:	100 Btu	95 Btu	0 Btu

### YIELDS FROM METHANOL

Average Bed Temperature,°F	775°F
Pressure, psig	25
Space Velocity (WHSV)	1.0
Yields, wt % of charge	
Methanol + Ether Hydrocarbons Water co, CO <sub>2</sub> Coke, Other	0.2 43.5 56.0 0.1 0.2 100.0
Hydrocarbon products, wt % Light gas Propane Propylene i-Butane n-Butane Butenes C <sub>s</sub> + Gasoline	5.6 5.9 5.0 14.5 1.7 7.3 60.0 100.0
Gasoline (including alkylates), wt, % (96 RON, 9 RVP)	88.0
LP Gas, wt %	6.4
Fuel Gas, wt %	<u>    5.6</u> 100.0
SOURCE: Reference 25	

### 3.4.1. General

**Oil** shale resources **vary** widely in their oil yields. High grade shale is normally defined as a deposit that averages 30 or more gallons of oil per ton of shale. Low grade shale averages 10 to 30 gallons per ton<sup>0</sup> (Reference No. 7). Several factors determine whether or not an oil shale deposit *is recoverable*. These include oil yield (usually equal or above 20 gallons per ton), zone thickness, overburden thickness, the presence of other materials *in the shale*, availability of needed resources such as water and services, and location relative to markets.

There are two major routes for converting oil shale to liquid or gaseous fuels. They are:

- Conventional mining followed by surface retorting (heating) , and
- 2. In situ (in place) retorting

In addition, there is modified in situ. In this process, the perme ability (i.e., void volume) of oil shale deposits is increased in order to enhance the in situ retorting by removing some of the shale. The methods of rein@ or increasing the permeability of the oil shale deposits are explained in reference &.

### 3.4.2. Surface Retorting

9

In surface retorting of oil shale, the heating takes place above ground. The shale is crushed to the right size, and fed into a retorting vessel. Heating the shale to between 800°F and 1000°′F removes abut 75 percent of the kerogen from the shale (Reference No. 8). Different retorting precesses apply heat to the shale in different ways. Gas or non combustible solids such as sand or ceramic balls can be used as heat carriers. The vapor produced during the heat@ is condensed to form crude shale oil. It can be further upgraded and refined to produce more marketable products.

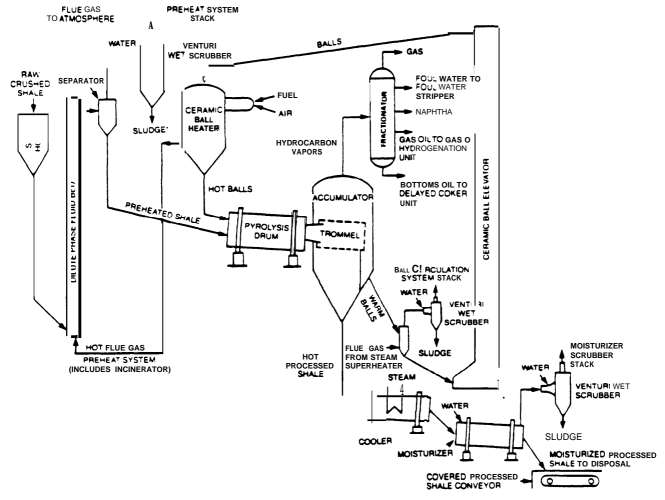
As a generic surface retorting process, TOSCO II is described. Its schematic diagram is given in Figure 3.10 (Reference No.\_8).

Ø

Shale deposits yielding less than 10 gallons of oil per ton are normally omitted from USGS resource estimates.

Raw oil shale is crushed to 1/2 inch and preheated to  $500^{\circ}$  F. It is mixed with hot ceramic balls 3/4 inch in diameter and at  $1200^{\circ}$ F in a retorting Pyrolysis drum (Reference No. 25). About two tons of balls mix with every ton of shale. The oil shale is heated to  $900^{\circ}$ F, releasing hydrocarbon vapors from the kerogen. The spent shale and the balls pass to the sealed accumulator vessel, in which the balls are separated from the shale by a heavy duty rotating cylinder with numerous holes. The balls are lifted by a bucket elevator to the gas fired ball heater, which heats the balls to  $1270^{\circ}$ F by direct contact heat exchanger. The spent shale goes through

### FIGURE 3.10



### The TOSCO II Oil Shale Retorting System

SOURCE Oil Shale Retorting Technology prepared for OTA by Cameron Engineers. Inc. .1978

a special heat exchanger which cools the shale for disposal and produces steam for plant use. Then the spent shale is quenched with water and moisturized to 14 percent, a level proper for disposal.

Hot flue gas from the ball heater is used to lift raw shale to a point at which it can subsequently flow by gravity into the pyrolysis drum. The flue gas also heats the raw shale to approximately 500°F.

Table 3.10 (Reference No. 25 ) summarizes the basic material balance for a TOSCO II retort module.

### TABLE 3.10

## BASIC MATERIAL BALANCE FOR A TOSCO II RETORT MODULE

Oil Shale

Feed ra	te, TPSD	10,700
Fischer	Assay, GPT	20

Pipelineable Shale Oil Product

production rate, BPSD	4,500	
Properties		
Gravity, *API	28.6	
Viscosity (SSU @ 30°F)	800	
Pour Point, 'F	30	

Table 3.11 (Reference No. 35 ) summarizes the energy balance for a plant producing 47,000 barrels per day. Table 3.12 (Reference No. 17 ) summarizes the components, resource requirements and potential impacts of surface oil shale retorting.

### Tab I e 3.11

### Estimated Energy Balance For a TOSCO II Plant producing 47,000 BPSD\* Upgraded Shale Oil From 35 Gallons Per Ton Oil Shale

	Btu/hour (lo Btu's)	Percent of Total Energy Input
Product Output		
Product oil LPG	10.30 0.70	58.00 3.94
Diesel fuel	0.11	0.62
System Losses		
Spent shale <b>and</b> moisture	1.78	10.02
Residual carbon (coke)	0.93	5.24
Ammonia	0.11	0.62
Sulfur	0.06	0.34
Cooling water	1.07	6.02
Water evaporat on on shale	0.25	1.41
Losses (includ <sup>ing</sup> flue gas heat)	2.45	13.79
Energy Input	17.76	100.0
Raw shale Steam	17.00 0.53	95.72 2.98
Electrical energy	0.23	1.30

\* BPSD = barrels per stream day

SOURCE: Reference 35

רו כ מזמאש				The start of the start of the second start of the start o
<pre>Milery Diff.m.i  IIII = 30.000 big/day of crude abula ell  10.13 = 30.000 big/day of crude abula ell  11.11 = 30.000 big/day  11.11 = 30.0000 big/day  11.11 = 30.00000 big/day</pre>	Miconcel Marning (nu loll bur Produced) FML(h) transmission ell contact ell contact composition must chain must chain must chain must chain must chain must chain must control on oblic must control on oblic must control on oblic must control on oblic transmission t	219,000 teas 11 al 11 a	HallBuilg AND PRONUCTSI (Pur 10 <sup>1</sup> Preduced) (Pur 10 <sup>1</sup> Preduced) porticicalistica porticicalistica come serve direct 41- charge to any using -conces <u>pontio builg</u> (b) <u>pontio builg</u> (b) refiland abilg oll buil content	11.1 11.1 11.1 11.1 11.1 11.1 11.1 11.
<ul> <li>and the restant provides the high provide the provided by the provided by the provide the provided by the provide</li></ul>		britera 3.1 11.5 11.5 11.5 11.5 11.5 11.5 11.5	bothered bistonance 1.5 1.1. 1.4 Vided by <u>samuel</u> production, manured in cril fon Btu. autput, anaurted in crilition Btu.	
<ul> <li>Nowcial (a) Environmental Provertion Aurory, Muniforthy Environmental Income of the Cort and Oil Analy Industries, 600/1-17-013, Parmers 1871.</li> <li>(b) University of Danver, Environmental Income Aurory Analysis, France And Analysis, France 1973.</li> <li>(c) University of Contame. Environmental Analysis, 1873.</li> <li>(d) Environmental Comparison, Environmental Environmental Analysis, 1873.</li> <li>(e) Environmental Comparison, Environmental Environmental Analysis, 1873.</li> <li>(e) Environmental Environmental Environmental Environmental Analysis, 1873.</li> <li>(e) Environmental Envited Environ</li></ul>	of Ind Environment of Industry of Medical Analysis (1994) Anal	ute Coal and Oll An Tric Deputs an Lie 1 1615.	<u>19   Industries</u> , 600/1-17-013, 7, 19400    <u>940 0 Freespo</u> , March II	

### 3.4.3 Modified In Situ Retorting

Occidental modified in situ oil shale retorting process is selected as representative. It involves the mining out of about 10 to 25 percent of the shale deposit. This mined portion would presumably be retorted by one of the surface retorting processes, or if its oil content is too low, will be treated as waste (Reference No. 37).

Figure 3.11 (Reference No. 8 ) represents in schematic form a generic modified in situ oil shale retorting process. 'Figure 3.12 (Reference No. 37 ) is a more detailed description of the Occidental modified in situ retorting process. As observed in Figure 3.12 , in steps A or the pre-detonation phase, drifts (chambers) are excavated at the top and bottom of the shale deposit, which is about 300 feet-thick. An interconnecting shaft is dug to connect the drifts. Rooms with a volume of about 15 to 20 percent of the eventual volume of the planned chamber are then mined. Shot holes are drilled to allow blasting of the shale oil to produce the desired fragmentation.

In the burn phase, the explosives in the shot holes are detonated. A rubble-filled chamber is created which can function as a batch retort. The percentage of void space and the particle size distribution of the rubble are a function of the explosive loading. Connections are made to air/gas recycle and air supply compressors. An outside heat source (e.g., off gas or oil from other retorts) is used for heating the rubble at the top of the retort. Oil shale and hydrocarbon gases are produced which move downward. Residual carbon is left on the spent shale.

The retorting reaction is terminated after a predetermined amount of the rubble has been retorted by halting the external heating supply. The residual carbon is utilized to continue the combusion process, which now does not need external heating. The flame front moves downwards, preceded by the liquid and gaseous products retorted from the shale by the hot, oxygen-deficient combusion gases. The liquid hydrocarbons collect in a sump, from which they are pumped to the surface. The gaseous by-products are used partially, with steam, as a recycle stream to control the oxygen content of the inlet gas. The four distinct zones that develop during the retorting are shown in Figure 3.11.

Table\_ 3.13 \_(Reference No. 17 ) summarizes the components, resource requirements, and potential impacts of modified in situ retorting.

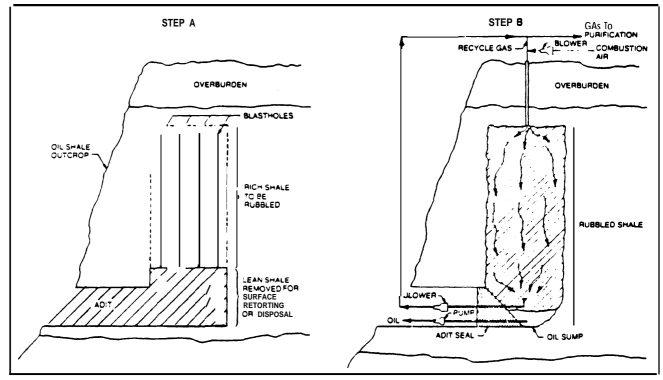
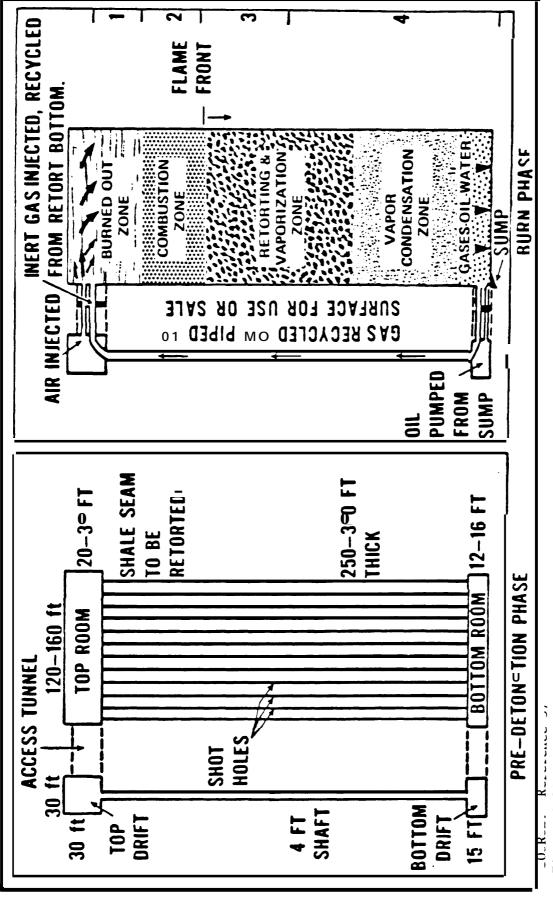


Figure 3.11: Modified in Situ Retorting

SCURCE TIA Sladek - Recent Trents in OH Snale - Part 2 Minung and Shale Oil Extraction Processes - Mineral Industries Bulletin, vol. 18. No. 1. January 1975 p. 18





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<b>ТАНГЕ 3 13</b>			Mod	Modified In-Situ Shale Retorting (Occidental)
<pre>PHECT STERM: PHECT STERM: 111 = 04,100 team of raw mhole misma/(a) + 101,400 team of raw mhole misma/(a) + 00,000 bill mu/year = 040,10 bill mu/year = 040,00 bill = 040,00 bil</pre>	resource usen: (Pre: 10 <sup>13</sup> Buu Produced) (Pre: 10 <sup>13</sup> Buu Produced) related anals united anals united anals united anals composition (a) content and anal content and and relation terrestion continue to continue to continue to production continue to continue to continue to continue to production	111, 900 tons 111, 900 tons 111, 900 tons 111, 900 tons 111, 900 tons 111, 900 tons 111, 400, 900 111, 4	RESIDUALY AND FROOUCTSI (Par 10 <sup>1</sup> Bu Froduced) ALE FOLLITATTS <sup>14</sup> . a) ALE FOLLITATTS <sup>14</sup> . a) Bot Bot Conco. There is assumed a second alliser is assumed a conco. Free is assumed a conco. Free is assumed a conco. Free is assumed a conco. Free is assumed a conco.	
600kCE1 (a) Commendationer, Incorporated, Synthetic Pael Manhoud, 1973. 10 1615 Base Group, Environmental Characterizations for Torney Technologies and End Uses, Bastation 2, 1914. (c) Basetamu of Largy, <u>Brait Environmental Impert Statement for the Conterpo OII Shale Leasing Program</u> , 1979. (d) Environmental Protection Agency, <u>A Preliminary American Fortunamental Imperts from OII Shale Leasing Program</u> , 1970. July 1977. (d) Antione OII, Juc., <u>Leases 6 Occidenty OII Shale</u> , Juc., Modifications to Detailed <u>Breeled Programs</u> , 0007-77-900, July 1971.	ic fuel mathematical and the fuel of the fuel of the fuel fuel for the fuel fuel of the fuel of the fuel of the fuel commutant of the fuel fuence to the fuel fuel fuel fuel fuel fuel fuel fue	ind End Usee, Roviel Pruurye 01 St ad Pruurye 01 St Ditailad Deve Jopper	an b, 1978. aie <u>taetan Progras</u> , 1979. <u>11 Shata Beveluamen</u> t, 600/7- 1 <u>12 Shata Dut Data Tract C-</u>	17-040, July 1977. 1. Fabruary, 1971.
SOURCE: Reference 17				

# 3.5 comparison of the Various Synfuel Systems With Respect to Resource Requirements $\frac{10}{10}$

In order to estimate the resource requirements of the coal and oil shale fuel cycles we need first to assess their energy utilization efficiencies. These are summarized in Table 3.14.

The *resource* requirements of coal and oil shale energy systems per 10<sup>6</sup> Btu of product delivered to end user are given in Tables 3.15 and 3.16. Tables 3.17 and 3.18 convert these requirements to energy systems producing 50,000 barrels of oil equivalent per day.

Manpower requirements for operating and maintenance labor of coal conversion plants are given in Reference 29.

They are:

Plant operatorsOperating supervisorsMaintenance laborMaintenance labor supervisors30Administration30Total355

These manpower requirements are for a basic (ESCOE) coal conversion plant that consumes 25,000 tons of coal per day with 22.4 million Btu/ton and produces 50,000 bbl/day liquids output.

Very considerable variations exist in the literature in respect to manpower requirements for the other phases of the fuel cycle. They depend on such variables as methods of mining, location of mine, kind of transportation system and extent of beneficiation. A table indicating the ranges of variables is given in the footnote in respect to the conversion plants.

10 Limitations of Data Sources: Evaluations carried out in this report are often sub ject to great uncertai nties because:

(1) The information available is only of preliminary nature. There are no full scale operating synfuel plants in the U.S. (subject to U.S. siting considerations), so that data needs to be extrapolated from pilot plants with many uncertainties of scale and dissimilarities associated with the extrapolation, as well as specific siting and f eedstock characteristics discussed below.

10 (cent'd)

(2) There are variations among sources which are often due to different assumptions or local influences. Changes in design account for some differences as the technology changes and the environmental regulations change. Many of the assumptions are not stated - or even referenced. Budget and time limitations, however, nessitate the need to use exist& data bases, rather than the development of new data.

Even estimating the range of uncertainties is often a value judgement **process**, **unless more**extensive on-site interviewing with site and process specific sources of information are developed.

3-40a

				(In Percent)		
	1 Coal Gas	l 2 Coal Gasification	3 4 Coal Liguefaction	4 efaction	5 6 <u>Oil Shale Retorting</u>	cting
	<u>Medium-Btu High-Btu</u>	<u>High-Btu</u>	Direct	Indirect	Surface Modifie	Modified in Situ
Beneficiation <sup>a</sup>	∃6.4-97.3	∃6.4-97.3 96.4-97.3	96.4-97.3 96.4-97.3	96.4-97.3	96.4-97.3	100
Transportation to Conversion Plant <sup>b</sup>	98.5	38.5	98.5	98.5	<b>5.</b> 66	100
Conversion to Fuel <sup>C</sup>	83	59	64-70	48-57	67	61
Upgrading and Refining <sup>d</sup>	N.A.e	N.A. e	75–95 <sup>f</sup>	95-100 <sup>f</sup>	977	6LL
Distribution to End User	96.94	97.1 <sup>i</sup>	98.8 <sup>j</sup>	98.8 <sup>j</sup>	98.8j	98 <b>.</b> 8j
Overall Energy Efficiencies	76.4-79.2	54.4-54.9	45.0-63.0	42.8-54.0	48.9-49.3	46.4
SOURCE: E. J. Bentz	E. J. Bentz & Associates	S				

Resource Utilization Efficiencies of Generic Synthetic Fuel Energy Systems

Table 3.4

E. J. Bentz & Associates SOURCE:

### Notes for Table 3.14

- a. Estimates of losses of coal and oil shale from beneficiation (in terms of Btu's) vary broadly among authors, depending on the assumed degree of upgrading and the kind of coal or oil shale used. Estimates vary from 0% (Reference 37a) ; 2.7-3.6% (Reference 7) ; and 12.5% for intensive beneficiation (Reference No. 17) .
- b. Average value of losses are 1.5% (time from Reference No. 7) . In the case of oil shale, where distances are shorter, 0 .5% is assumed.
- c. The @et efficiencies (rather than the process efficiencies) were used. The efficiencies for coal conversion processes are derived from Roger and Hill. (Reference 29) . In the case of H-Coal, the syncrude efficiency was used. In the case of oil shale retorting processes, the efficiencyes are derived from DOE (Reference No. 17) .
- d. Data on efficiencies of upgrading and refining syncrudes is very limited and unreliable (see Section 1.7) .
- e. N.A. means not applicable.
- Overall yields for SRC II of finished fuels range between 83 and 98 f. liquid volume percent of SRC II syncrude, depending on the product slate and how refinery fuel and hydrogen plant feed are supplied. An average of the net product yields ranging between 88 and 91 was assumed (Reference No. 22) . However, these values apparently do not include coal use for the\_production of hydrogen needs for the upgrading process. If coal-derived hydrogen is to be used (as against hydrogen from nuclear fission or from biosynthesis), then the upgrading and refining efficiencies for coal conversion products become 75 percent. However, in some cases it may be expected that all of the hydrogen and energy required for the Upgrading/refining process would be obtained from residuals, higher boiler fractions, and methane produced in the process or plant refinery(which may include the use of Petroleum • In the case of indirect liquefaction derived vacuum Processes, all the needed hydrogen is accounted for in the gasifier, and higher upgrading efficiencies can be achieved, depending on product slate .
- 9" Derived from Reference 26a. However, MIS oil is easier to upgrade, so that higher efficiency may be in order.
- h. Derived from Reference <u>17</u>.
- i. Derived from Reference 7.
- j\_ Derived from Reference 7 and 10.

		(In 10 <sup>-3</sup> ton of	fossil carl	on/106 Btu fi	(In 10 <sup>-3</sup> ton of fossil carbon/10 <sup>6</sup> Btu fuel delivered to end user)	o end user)
	l 2 Coal Gasification	2 ification	3 4 Coal Liquefaction	4 efaction	5 6 Oil Shale Retorting	6 etorting
	<u>Medium-Bt</u>	<u>Medium-Btu</u> <u>High-Btu</u>	Direct	Indirect	Surface Modi	<u>Modified in Situ</u>
Beneficiation	1.2-1.6	1.7-2.2	2.0-2.7	2.1-2.8	0.9-1.2	0
Transportation to Conversion Plant	0.7	6.0	1.1	1.2	0.2	0
Conversion to Fuel	0.7	25.1	22.2-26.7	33.5-40.5	11.4	18.7
Upgrading and Refining	I	I	3.7-18.5	0-3.9	7.9	11.0
Distribution to End User	1.4	1.8	6.0	6.0	0.4	0.6
cverall Consumption	9.1 <del>,</del> 10.3	9.1+10.3 27.6-27.9	27.4-40.7	35.8-44.≈	17.5-17.6	25.7

Table 3.15 Fossil Carbon Consumption of Generic Synthetic Fuel Energy Systems (J

SOURCE: E. J. Bentz & Associates

### Notes to Table 3.15

- a This table summarizes the consumption of fossil carbon contained in the feedstocks or products during the various phases of the various synfuel cycles.
- b The numbers in the table are based on the following assumptions:
  - (i) The resource utilization efficiencies are those developed in Table 3.14.
  - (ii) The carbon content of bitumimous coal averages 87.8%, lignites -72. 5% and sub-bituminous~ reals - 73. 5%. The carbon content of the kerogen (i. e., crude shale oil) averages 80. 5%. (Ref. 26b) . For convenience, an average figure of 80% for the carbon content of coals and kerogen is used.
  - (iii)The loss in fossil carbon is directly proportional to the loss in coal or kerogen.
  - (iv) The Btu content of a ton of coal is  $24 \times 10^6\,{\rm Btu}$  and of ton crude shale oil is  $36 \times 10^6\,{\rm Btu}.$
- c A sample calculation for medium Btu coal gasification is as follows:

A ton of feedstock bituminous coal has 24x.10° Btu, of which 18. 34x10° to 19. Olx10° Btu is delivered to the end users (74.4 to 79. 2% overall energy efficiency - see Table 3.14). Since a ton of feedstock coal. has 80% fossil carbon content, and 20.8% to 23.6% of it is consumed during the medium Btu coal gasification fuel cycle, (see Table 3.14), the total fossil carbon consumption of the cycle is between 0.1664-0.1888 tons per 18.34x10 to 19. Olx10 Btu delivered to end users.This translated to 0.009 to 0.010 tons of fossil carbon per 10° Btu.

Table 3.16 Water Consumption of Synthetic Fuel Energy Systems (Generic (In gallons per 10 <sup>6</sup> Btu product delivered to end user)
Table 3.16

5 6 Oil Shale Retorting Surface Modified in Situ	0.7-1.1	o	9-13	24	0
5 0il Shale Surface	0.7-1.1 0	0	9–32	24	0
3 4 Coal Liquefaction Direct <u>rudirect</u>	0.6-0.9 0.6-0.9 1 2 1.2	0	13-26	I	0
3 Coal Lig Direct	0.0-0.0	0	7-26	I	0
l 2 Coal Gasification Medium-Btu High-Btu	0.6-0.9	ч о Н	13-24	0	0
l 2 Coal Gasification Medium-Btu High-B	0.6-0.9	7. O	13-24	0	0
	Mining <sup>a</sup> ,b	Beneficiation <sup>C</sup> Transportation to Conversion Plant	Conversion <b>6</b> Fuel	Upgrading and Refining <sup>e</sup>	Distribution to End User

SOURCE: E. J. Bentz & Associates

e jb&a

### Notes to Table 3.16

- a The water required for mining and preparation of the coal or shale and for the disposal of ash or spent shale is a function of location, mainly through the amount of material that must be mined or disposed; and the degree of attested surface reclamation. Assuming 2/3 of coal is surface-mined and 1/3 is undergroundd mined, water consumption for surface mining ranges between 0.55 and 0.98 gallons per 10<sup>b</sup> Btu of product, and for underground mining - 0.75 gallons per 10<sup>b</sup> Btu of Product (Reference No. 17).
- b Assume 2/3 of oil shale is surface mined and 1/3 is underground mined. Water consumption or both kinds of operations range between 0.7 and 1.1 gallons per 10<sup>6</sup> Btu of **product (Reference No. 17)**.
- c Consumption of 1.2 gallons of water 10<sup>6</sup> Btu Of product is assuned for beneficiation of coal (Reference No. 17) and none for shale oil.
- d Consumption of water for the conversion of feedstock to fuels depends principally on the overall plant conversion efficiency, degree of water recycling, and the water content of the coal or shale. Consumption figures range from 13-24 gallons per 106 Btu of product for coal gasification; 7-26 for direct coal liquefaction; 13-26 for indirect coal liquefaction; 9-32 for surface shale retorting; and 9-13 for modified in situ shale retorting (Derived from References 17, 37b,c).
- e Water consumption for upgrading and refining is not available in the literature. The estimates presented for shale oil upgrading are based on private conversation with Mr. Bobby Hall and Ray Young of the American Petroleum Institute 3/81. For shale oil 100 gallons per barrel are needed to make the raw shale oil suitable for pumping, and 40 more gallons per barrel to convert it to transportation fuels. Polling of a large number of oil companies and API experts did not result in water consumption estimates for upgrading of coal liquids (namely: Robert Howell, Bonner and Moore, Fred Wilson Texaco, Patton, Nanny, Hall and Young of API 3/81).

	Situ							
torting	lified in S	N.A. <sup>7</sup>	N.A. <sup>7</sup>	N.A. <sup>7</sup>	24.0	18.5	18.3	
<u>Oil Shale Retorting</u>	Surface Modified in Situ	62.2-62.7	60.5-60.6	60.2-60.3	24.0	18.5	18.3	
Coal Liquefaction	Indirect	79.8 8.2-10.3	6.8-9.4 7.9-9.9	6.7-9.3 7.8-9.8	19'4-24.6 18.5-19.4	18 <b>.</b> -	18.3	
Coal Lig	Direct	7.0-9.8	6.8-9.4	6.7-9.3	19', 4–24.6	1a5	18.3	
Gasification	Medium-Btu High-Btu	3.0-8.1	7.8	۲.٦	18.8	18.8	18.3	
Coal Gasi	Medium-Bt	5.6-5.8	5.4-5.6	5.3-5.5	18.9	18.8	18.3	
		Mining <sup>5</sup>	Beneficiation <sup>5</sup>	Transportation to Conversion Plant <sup>5</sup>	Conversion to Fuel <sup>6</sup>	Upgrading and Refining <sup>6</sup>	Distribution to End User <sup>6</sup>	

Annual Feedstock Requirements for Generic Synthetic Fuel Energy Systems Producing 50,000 bbl Oil Equivalent per Day to End User (In millions of tons or barrels of oil) Table 3.17\*

SOURCE: E. J. Bentz & Associates

\* These are the quantities of  $\cos 1$  shale  $\Rightarrow$  equivalent oil leaving  $\sigma$ e indicated phase of the fuel cycle.

### Notes to Table 3.17

- 1. Same assumptions and references as those in Table 3.14.
- 2. Oil has energy content of 5.8 x  $10^6$  Btu/barrel.
- 3. Coal has energy content of 24 x  $10^6$  Btu/ton.
- 4. Oil shale has energy content of 3.45 x  $10^{^{\rm 6}}\,\text{Btu/ton}$  (based on 25 gallons of oil per ton) .

-.

- 5. Tons of coal or shale.
- 6. Barrels of oil equivalent.
- 7. N.A. is not applicable.

# Table 3.18\* Annual Water Consumption of Generic Synthetic Fuel EnergySystems Producing 50,000 bbl Oil Equivalent per Day to End User(In million gallons pervear)

	<u>Coal Gasi</u>	fication	Coal Lig	uefaction	Oil Sha	ale Retorting
	Medium-Btu	1 High-Btu	Direct	Indirect	Surfac	e <u>Modified</u> in Situ
Mining	64-95	64-95	64-95	64-95	74-120	74-120
Benef iciation	130	130	130 "	130	0	0
Transportation <b>to</b> Conversion Plant	0	0	0	0	0	0
Conversion <b>to</b> Fuel	1400- 2500	1400- 2500	740- 2800	1400- 2800	<b>950-</b> 3400	950- <b>1400</b>
Upgrading and refining	0	0			2500	2500
Distribution <b>to</b> End User	0	0	0	0	` 0	0

\* Sam assumptions and references as in Table 3.16.

SOURCE: E. J. Bentz & Associates

Table 1 Fo	Table 1 Footnote to Chapter 3: Ma <u>Pr</u>	Manpower Requirements of Generic Synfuel Plants Producing 50,000 Barrels of Oil Equivalent per Day	eric Synfuel Plants Dil Equivalent per Dav
	l Coal Gasification	2 Coal Liquefaction	3 4 <u>Oil Shale Retorting</u>
	Medium-Btu-High-Btu	Direct & Indirect	Surface Modified in Situ
Peak Construction (men)	l,500-4,800a	2,200-8,000 <sup>b</sup>	330 <sup>d</sup> 4,900 <sup>d</sup>
Construction			
(man-years)	3,400 - 10,800 <sup>a</sup>	7,500-25,∞0 <sup>b</sup>	1100 <sup>d</sup> 16,000 <sup>d</sup>
Operation and Maintenance (men)	320-500 <sup>a</sup>	355–3800C	1200 <sup>d</sup> –
	-	-	
<sup>a</sup> DOE, 1980, Comparative A	ssessment of Health and	<sup>a</sup> DOE, 1980, Comparative Assessment of Health and Safety Impacts of Coal Use. DOE/EV 0069.	. DOE/EV 0069.
b The lower value is deriv	ed from DOE/EV 0069; <sup>on</sup> e	<sup>b</sup> The lower value is derived from DOE/EV 0069; <sup>or</sup> e upper value - from Reference <u>34</u> .	nce <u>34</u> .
<sup>c</sup> The lower value is deriv	ed from Reference 29; <sup>u</sup>	<sup>C</sup> The lower value is derived from Reference $\underline{29}$ ; <sup><math>\sigma</math></sup> e upper value - from Reference $\underline{34}$ .	tence <u>34</u> .

SOURCE: E. J. Bentz & Associates

<sup>d</sup> Derived from Reference 17 and assuming 5 year construction of plant peaking at 30% of  $\circ \approx 1$  man-years labor requirements (Reference <u>34</u>).

3-50

CHAPTER 4 : SYNTHETIC FUEL PROCESSES CONVERSION COST AND PRODUCT ECONOMICS

### Section No.

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4.2	Scale of Production 4-7
4.3	Product Quality 4-8
4.4	Estimating Methods 4-11
4.5	Product Upgrading 4-22
4.6	Refining synthetic Liquids 4-28
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4-8	Addendum to Chapter 4: Basis for Cost Assumptions
	<ol> <li>Basic Conversion Plant* 4-43</li> <li>Assumptions of Product</li> </ol>
	<ol> <li>Assumptions of Product Upgrading</li></ol>

### 4.1 CONVERSION COSTS AND PRODUCT ECONOMICS

The following evaluation of a wide range of alternate fuels produced from coal attempts to build upon prior work in the field that has, successively, estimated the plant construction and operating costs for each process, standardized the bases of estimation (time of construction, size of plant, location, financing methods, etc.) and evaluated the quality of product produced.

Such work has been sponsored by the Department of Energy since the early 1970's. The most recent work was performed by the Engineering Societies Commission on Energy, Inc. (ESCOE).<sup>L</sup> That work collected prior analyses performed for DOE and others, made adjustments in each to account for differing assumptions regarding input prices, plant scale, financing methods and costs, and thus reevaluated them on a more common basis. The differences in product quality were factored for value based on current price relationship among natural petroleum products.

Our approach will differ in several regards:

First of all we shall use the baseline ESCOE plant models, capital costs and operating cost relationships, updated to a uniform 1980 dollar basis.

<u>Second</u> we shall scale all plants to a common <u>output</u> plant size in order to retain comparability at other, downstream stages of processing and use.

Third we shall deal with differences in product quality directly, and on a cost of product basis, by considering the additional costs required to upgrade lower quality products and make them comparable with the higher grade synfuels.

Fourth we shall then examine the methods and costs of further processing and transporting the generic synfu el products to make them available to end use markets.<sup>3</sup>

The ESCOE capital estimates were all adjusted to a 1980 dollar basis by the use of the Wholesale Price Index - Industrial Commodities Index. Others have frequently used the Chemical Engineering Plant Index, however we feel that no significant historical difference exists and the WPI Index basis is a more suitable bench mark for further forecasting since it is a component

<sup>1</sup>Coal Conversion Comparison, ESCOE Report FE-2468-51, July, 1979.

<sup>&</sup>lt;sup>2</sup>ESCOE scaled all plants to a common <u>input</u> size in order to simplify the costs - auxiliaries and off-sites are normalized.

<sup>&</sup>lt;sup>3</sup>We did not examine differences in end use efficiency that exist or are possible. This should be subsequently examined.

of Us. macro-economic forecasting models and the Chemical Construction Index is not.

Exhibit 4-1 displays the original capital cost estimates of ESCOE. Exhibit 4-2 updates these estimates to a uniform 1980 cost basis.

Operating costs are more complex. The major cost categories are:

• Coal

- Utilities Water Power
- . Catalysts and Chemicals
- . Labor
- Overhead
- . Maintenance

Coal prices are uniform to all processes - as are assumed costs of water, power and labor. The costs of overhead are a uniform fraction of operating and maintenance labor - they include administrative personnel costs as well as G&A expenses. The maintenance rule is made uniform among systems-although differences should exist on the basis of system approach.

The original ESCOE operating cost variables are shown on Exhibit 4-3. These unit prices provide the bases for updating the ESCOE costs to the values shown on Exhibit 4-4.4

The cost of producing hydrogen for product <u>upgrading</u> is partially imbedded in other estimates. The uniform condition is that hydrogen is demanded at a greater level then could be supplied from excess char, residue, or filtrate from the process plant. Therefore a hydrogen plant must be built at the upgrading plant site. This plant is designed to reform synthesis gas. The cost of hydrogen can then be based on the hydrogen plant's costs - including syngas feed at the estimated syngas product costs of our companion syngas plant. Alternately we could capitalize a coal gasification plant in this area, however that seems to be an even more unrealistic mode of system optimization.

In the long run, as product slate demand for synthetic coal liquids becomes clarified, the optimization of an integrated coalto-product plant can be designed in a much more sophisticated manner.

<sup>4</sup>The input costs were in certain instances drawn from original sources cited by ESCOE.

<sup>b</sup>Or reform synthetic fuel product - the cost is comparable \$6.25 - 6.75/MM BTU.

EXHIBIT 4-1

MAJOR ON-SITE PLANT COST IN MILLIONS OF MID 1978

MAJOR ON-STTE PLANT COST	STTE PLAN	VT COST	SNOITTIW NI		OF MID	1978	လ	
Category <u>Process</u>	<u>SRC-11</u>	EDS	H-FO	H-Syn	FT	ΣI	Lur.	West. Syn.
Coal Preparation	63	63	84	84	63	63	06	63
H <sub>2</sub> or Gasification	253	190	138	158	228	228	143	22
2 0, Plant	129	1	67	87	117	175	114	80
Gas Shift	I	I	о <b>£</b>	35	I	<b>4</b> 0	30	I
Acid Gas and Sulfur Plants	60	60	57	57	57	57	136	57
Reactor Section	195	180	140	210	55	106	0 <b>6</b>	I
Conversion	I	1	ı	I	100	75	20	42
Gas Plant	30	I	30	25	25	10	12	I
Flexicoker	I	160	I	I	I	I	I	1
Pollution Systems	44	44	40	40	40	40	55	24
Solvent Hydro. or Catalyst Prep.	I	82	I	I	e	I	i	I
Compression	1	I	I	I	I	1	200	1
Total less Int. Including Indirects	1262	1270	955	1134	1121	1212	1151	684
Notes: 1. M includes HF 2. Some EDS cost	F Alkylation. t included in		Flexicoker.	.er.				

All costs shown above are considered bare cost and have not been confirmed with process developer. э.

E. J. Bentz & Associates SOURCE:

	Reference: Tons of Coal/Day		20,938 22,584	22,242	20,695		31,095	20,833	10,263	26,174		23,000	כוכ בו	CTC'/T	N.A.
	Capital Cost/ Re MM BTU/ T Yr. Co		\$12.13 13 26					17.42				12.30	5	00.0	11.93
	Total BTU I El4/Yr.		1,081	1.115	1.048		1,112	8016	428	905		1.067 <sup>3</sup>	1 0673	100'T	.953 <sup>5</sup>
	Capital Cost/ Dailv BBL	•	\$26,210 28.440	25,040	19,620		34,600	27,930	12,170	42,650		(26,260) <sup>3</sup>	3005 LL	(UE1, LEV)	15,960
(Million 1980 Dollars)	50,000 BBL/ Day Output Basis		1310.8 1422	1252	980.9		1730	1396.4	608.4	2132.9		1313.1	000	C. 400	.198.
	ESCOE Basis <sup>1</sup> 1980 \$		\$1,565. 1.574.	1,407.	1,185.		1,391.	1,676.	1,482.7	2,225.4		1,427.	051	.100	.798.
	ESCOE Basis 1978 \$		\$1,262. 1.279.	1,134.	955.		1,121.	1,212.	1,195.	1,587.		1,151.	VOJ	. 100	700.4
	Coal Liquids	Direct Liquefaction	SRC-II EDS	H-Coal Syn. Oil	H-Coal F.O.	Indirect Liquefaction	Fischer/Tropsch	Mobil M' Lidom	Methanol	Methanol/SNG	Coal Gasses	<u>High BTU</u> Lurgi	Low BIU	ashrifitta saw	Shale Oil Surf. Retort.

50,000 BBL LIQUIDS/DAY PLANT BASIS

EXHIBIT 4-2

I

TOTAL CONVERSION PLANT INVESTMENT

2

<sup>L</sup>ESCOE - 25,000 tons coal/day input basis revised to reflect 20% contingency vs. 10% and 1980 dollars.

2 Mobil Research Center Basis - \$ 1977 - 27,300 ton coal input revised \$ 1980 and 1.738 markup of plant.

<sup>3</sup> Evaluated at average heating value of coal l quids 6.5 MM/BTU/bbl. x 50,000 bbl./day = 325 billion BTU/day (1.067 E14 BTU /yr.)

<sup>4</sup> OTA Basis - 3rd quarter 1979, 50, 🚥 bbl. basis.

<sup>5</sup>Evaluated at average daily value of 5.8 MM/BTU/bbl.

E. J. Bentz & Associates SOURCE:

	Total	7 W	104.8	0.611	121.0	95.0	0.111	$^{\circ}.601$	113.2	121.0	107.0	73.0	69.0	91.9	82.0	112.0	116.0	001.0	יר	/T.4	I
		1.000 1.100 1.100 1.0000 1.0000 1.000 1.000 1.000 1.000 1.000 1.000 1.000 1.000 1.000 1.00	55.	• <b>£</b> %	64.	48.	57.	56.	60.	65.	54.	47.	49.	50.	44.	58.	63.	53.	c yc	C.FC	I
	LANCE COST	Maintenance 2c	33.	38.2	38.5	29.	34.3	34.	34.	35.5	34.6	22.5	23.4	23.9	24.3	36.7	38.	32.	20 5	C.U2	I
EXHIBIT 4-3 COST = ATA ESCOE)	6 MAINTER	1.abor 2b	13.8	12.2	12.2	12.2	12.2	12.2	12.2	12.2	12.7	12.7	12.2	12.2	12.9	12.7	12.0	12.0	ς ς[	7.71	I
	OPERATING & MAINTENANCE COST	catalyst & Chem. 2a	3.0	<b>••</b> 9	<b>6.</b> 0	6.0	7.0	7.0	7.0	8.5	5.9	3.2	4.8	5.8	4.5	4.5	3.0	3.0	۲ ۲	) •	I
		Fuel 1	246.	246.	246.	246.	246.	246.	246.	246.	246.	246.	246.	246.	246.	246.	246.	246.	246	• • • • •	I
		Capital C	1092	1262	1270	955	1134	1121	1195	1212	1084	942	980	866	870	1151	1268	1066	684	<b>F</b> 00	E67
		Process	SRCI	SRC-11	EDS	H Coal: Fuel Oil	Syncrude	FrT	Methanol	M-Gasoline	$co_2$ Acceptor SNG	Syngas	HYGAS	BIGAS	Synthane	Lurgi	CE Power	West Power	Westinghouse	mfirla	Shale Oil

SOURCE: E. J. Bentz & Associates

4-5

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# ANNUAL OPERATING COST - 50,000 BBL LIQUIDS/DAY PLANT

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	Cost/ MM/BTU		6.47 7.06	7.24	5.51			9.61 11	7.75	11.57			6.74	4.74		8.31	
	Average Cost/ BBL Liq.		42.557 46.058	49.157	35.178			58,29 AA A60	20.189	(63.775)			.43.78)	(30,788)		48.20	
	Total Operating Cost		699 <b>.</b> 756.5	807.4	577.8			957.4	331.6	1047.5			1.617	505.7		791.7	
	Total Other		99.5 107.3	212.6	79.6			132.	48.	150.7			98.5	68.2		322.3	
	Taxes & Ins. 3%		39.3 42.6	37.5	29.4			51.9	18.3	64.		:	39.4	26.7		I	
ina Costs	Utilities 6 Supplies (50% of Chem.)		m m	3.5	m		1	້	י <b>●</b> m	Am		,	2	1.5		I	
Other Operating Costs	Maintenance 38 of Total Capital		39.3 42.7	37.6	29.4			51.9	18.3	64.0			39.4	26.7		1	
	Labor		11.9 13.	12.7	11.8			17.7	5.9	13.7			13.7	10.3		I	
	diems. (escalated 0 208)		و و	7	9				3.5	9			4	£		ı	
Feedstock	Coal (Shale) 0 \$30/Ton 00)		206.3 222.6	219.2	203.9			306.4	1.101	256.9			226.7	170.6		230.0	
	Capital Charges @ 30% of Capital		393.2 426.6	375.6	294.3		-	519. Ale e	182.5				393.9	266.9		239.4	
		Direct Liquefaction	SRC-11 EDS	H-Coal S	H-Coal F.O.	Indirect Liguefaction	Fischer-	Topsch	Methanol	Methanol/SNG	Synthetic Gas	Hİ BIU	Lurgi Med. BIU	Westinghouse	Shale 011	Surf. Retort.	

SOURCE: E. J. Bentz & Associates

### 4.2 SCALE OF PRODUCTION

Chemical process plant economics are highly sensitive to scale. Typical scaling factors or rules, are of the order of 60% -70%. This means that as plant size doubles the cost only increases by 60% - 70%. In the case of decreased scale - the factor works in the opposite direction, a decrease in scale to  $1\2$  plant scale leads to only about  $1\3$  decrease in cost, which in turn leads to almost 30% more capital being required <u>per unit</u> of output. In very capital intensive processes, the importance of this to product cost is great. Coal conversion processes typically have 1/2 of their costs derived from capital charges, therefore a doubling of scale could reduce total unit costs by as much as 15% - 20%.

For this reason the question of plant scale must be very carefully examined. ESCOE, in ordering the various estimates to the values shown in Exhibit .4-1 applied "typical chemical engineering scaling factors". It is beyond the scope of the present effort to audit that undertaking. However, it is incumbent upon us to avoid the distortion of fairly presented uniform cost data by another exponential adjustment of capital costs. We must rescale the liquids' plants since they have been standardized on an 'input' basis, whereas we must examine costs on a plant 'output' basis, since we are also examining downstream processes and costs, which in turn require uniform scale assumptions.

Several difficulties are present:

- 1. The optimal size of plant and vessels for various systems is not known, due to the fact that most processes are now being explored at 5 10% pilot plant scale.
- 2. In a shift from uniform input scale to a uniform output scale, the most efficient processes will suffer the greatest penalty for their <u>relative</u> downsizing. This is not realistic.
- 3. We are not aware of the relative changes that took place in the initial (ESCOE) standardization, hence are blind to the compound effect of a second scaling adjustment.

For these reasons, with the emphasis upon the above factors, in order of their ranking, we have chosen to restate costs on an output basis through a linear method of cost adjustment.

The principal justification for this apparently unsound procedure is found in the first factor above - there is no evidence of commercial scale economy available in the case of any processes, with the exception of gasification plants (or gasifier reactors). In that case, multiple train plants appear at subcommercial plant scale. In general, the bulk of the solid feed stock is so great, that initial reactor vessel sizes become limited by available fabricating (rolling, bending, heat-treating) facilities, as well as transportation constraints. Subsequent plant stage economics do not determine. The gasifier-reactor vessel size limitations are such that returns to scale may be limited at a relatively low level of output.

For this predominant reason, we have used a unitary cost scaling factor to shift from uniform input sized plants (25,000 tons of coal per day) to a uniform output basis - 50,000 bbl. per day. A normal procedure would otherwise unfairly penalize the most efficient processes. In the final analysis, efficiency will determine economic advantage.

#### **4.3** PRODUCT QUALITY (Reference No. 38)

The issue of product quality was resolved in a somewhat indirect manner by ESCOE. Their 'rating scale' value system (a measure of ordinal utility or value) which was based on present product price relationships is not a suitable method for long range economic analyses. During the long-run,. values change, end use patterns and conversion technology developments can create a surplus of a once premier product, or contrariwise, create a shortage of a previously unwanted by-product. Distillates and gasoline have traded places once and are perhaps posed to trade places again in their relative values.

The setting of widespread synthetic fuels production and use creates an entirely new framework for evaluating the 'normal refinery slate of petroleum derived products. We have created a slate of products that to some degree reflects the range of compounds present in crude oil and in some degree reflects the technology (now) available to separately produce these compounds. In some instances the products were specifically sought, in other cases markets were sought for by-products that were available.

When coal is introduced in lieu of crude oil to a substantial degree, the available range of products and by-products may be the same, but the proportions of availability will be quite different, as will be the cost of producing different fractions. "

The proportion of each fraction that can be derived from crude oils is highly variable depending upon the nature of the feedstock and the nature of the refining processes used. In general, increasing the lighter fraction (-350°F) involves more severe reforming, and higher cost. The use of a heavy, sour feedstock crude oil worsens this condition. The use of coal as the feedstock significantly exaggerates this condition in certain synthetic processes - such as direct liquefaction. Indirect liquefaction processes are specific for alcohols, gasolines and the light ends.

It is reasonable to visualize a population of crude oil and coal "refineries" with individually more specialized or limited product slates than are found in the universe of conventional refineries.

Broad slate coal synthetic liquids plants are unlikely to be widely deployed. This can be expected for several reasons:

- 1. Product upgrading is difficult and expensive once outside of the basic process.
- 2. A fair range of limited slate coal-conversion processes are becoming available, that more selectively produce various fractions.

The costs of achieving a given level of product quality increases in a slightly non-linear fashion as the percent hydrogen is increased or the boiling range is lowered. Exhibit 4-5 shows this relationship graphically. Benchmark products and costs are shown for several direct and indirect liquefaction processes. The indirect processes - which catalytically synthesize liquids from synthesis gas are specific for gasolines, alcohols and LPG. The direct catalytic hydrogenation processes tend to produce naphthenes and crude oil equivalent range compounds. The hydrogen solvent systems tend to produce a more limited range of product with a substantial (20 - 35%) naphtha fraction, the majority product in the distillate boiling range ( $350^{\circ}F - 750^{\circ}F$ ).

Increased yield of the higher quality products **can** be achieved by:

- . Increased coking of bottoms
- . Adding more hydrogen
  - . To process stream
  - . By hydrotreatment of products

The cost of the former is seen in the difference between SRC II and EDS on Exhibit 4-4. The Exxon donor solvent system cokes the bottoms (or heavy distillates) to yield more naphtha and LPG as follows:

	SRC II				EDS	
(8%)	6,400	bbl	Naphtha #2 Fuel Oil Distillate	(15%)	10,000	Naphtha LPG bbl Distillate
	72,300				75,400	

Similarly changing the H Coal process from a fuel oil to a synthoil mode increases cost as it lowers the average boiling range.

The distribution of product quality that is typical of each process is shown on the following page. (Exhibit 4-5).

		110	SYNT× ET <sup>±</sup> C		PLANT PRODUCT YIELDS	r YIELDS						
		- -	QUANTITY	I	BBLS/DAY (	OUTPUT						
			Direct Liquids	iquids		In	Indirect Liquids	iquids		Synthetic	stic	Shale
	Approx. API	SRC-II	SCE	H Coal (Syn.)	H Coal Fuel 0i1	Fischer/ Tropsch	Mobil 'M'	Meth- anol	Meth- anol SNG	Gas Iurgi H BTU I	West. Lo BTU	011 Shale Bit. (Surf.)
SNG (Low) MM BTU/Day											880	
Methanol (High) MM BTU/Day									140	8		
DALI	125 <sup>0</sup>	4,610				23,380	6,080					
Propane (C <sub>3</sub> )	148 <sup>0</sup>		2,950									
Butane (C <sub>4</sub> )	1100		3,160									
Methanol	NA							50,000	48,740			
$\omega$ soline (C <sub>5</sub> )	62 <sup>0</sup>					82,640	43,920					
Naphtha (C <sub>5+</sub> )	40 <sup>0</sup>	10,625	17,970	28,380	15,070	$1,490\frac{1}{2,490^2}$			1,260	2 <b>,</b> 025		
Fuel Oil (Heavy Distillate)	18 <sup>0</sup> +	35,000		21,620	34,930							50,000 <sup>4</sup>
Fuel or (Resid.	50		25,920									
<sup>1</sup> Light (Diesel) Fuel Oil API-56 <sup>0</sup> , <sup>2</sup> Heavy Fuel Oil API-41 <sup>0</sup> . 3 1.067 El4 BTU/yr. is equivalent to 50,000 BBL/day of typical Synthetic Liquids ≤ 6.5 Million BTU/bbl. 4 Synthetic light crude oil equivalent - approx. 20 <sup>0</sup> API.	Fuel Oil A r. is equi crude oil	API-56 <sup>0</sup> , <sup>2</sup> H Valent to equivaler	eavy Fuel 50,000 Bl nt - appr	Oil API-41 <sup>0</sup> BL/day of ty ox. 20 <sup>0</sup> API.	-41°. f typical API.	Synthetic	Liquids	≈ 6.5 Mi	llion BIU	J/bb1.		

SOURCE: E. J. Bentz & Associates

ejb&a

The range of quality is not entirely a function of the API gravity, the boiling range or hydrogen content, however, these related indices are sufficient for our purposes. We can relate the cost of producing a synthetic fuel to this scale. Exhibit 4-6 shows a graph of the production cost of the whole liquid product from various synthetic processes versus the average (50% distillation) boiling range of the synthetic product.

This chart shows the increase in average cost per million btu's as the average distillation range of the liquid is lowered. Thus gasoline costs more to produce via indirect processes such as Mobil 'M' or Fischer Tropsch, than naphthas, distillates and fuel-oils.

This scale, illustrates the relative costs of the ESCOE liquid fuel processes. It also contrasts the (1978) earlier ESCOE cost estimates with later estimates of shale oil costs developed by the Office of Technology Assessment (1980). The oil shale liquids, which reside in a higher boiling range than the coal liquids, appear significantly more expensive on this scale. In order to reconcile this discontinuity it is necessary to digress briefly.

#### 4.4 ESTIMATING METHODS

The accuracy of complex systems cost estimating has been the subject of several studies. These studies have been primarily behavioral rather than conceptual. As larger, more complex systems projects have been conceived, -the amount of unknown and untried system components have necessarily increased due to the great cost of large system prototypes. Pilot or process demonstration units and models are developed at extremely small scale for the same economic reasons; the subsequent scale-up is of a high order. Estimates drawn from bench or small scale pilot plants are subject to much greater estimating error.

Two overriding conclusions have been reached in this matter:

- 1. Cost estimates tend to decrease in variation from actual costs as the elapsed time between estimate and construction is shortened.
- 2. The accuracy of the estimate is related to the degree of detail of the design engineering.

Chemical process plants,  $^{\pmb{8}}$  public works,  $^{\pmb{9}}$  and weapons systems  $^{10}$  development and estimating histories have been analyzed, with

<sup>6</sup>Syngas (fuels) are not suitable related to boiling point measurement. <sup>7</sup>"An Assessment of Oil Shale Technologies", OTA - June 1980.

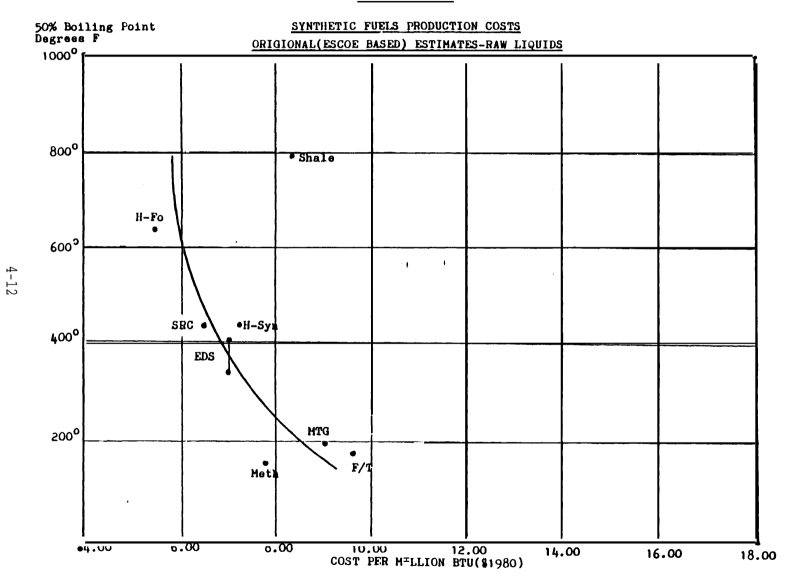
<sup>8</sup><u>A Review of Cost Estimates in New Technologies: Implications for</u> <u>Energy process Plants</u>, Rand Corp. for the Dept. of Energy July 1979.

<sup>9</sup> "Systematic Errors in Cost Estimates for Public Investment Projects ", Hufschmidt & Gerin, in <u>The Analysis of Public Output</u>, Columbia Univ. Press 1970.

<sup>10</sup> The Weapons Acquisition Process: An Economic Analyses, Peck & Scherer, Harvard Univ. 1962.

4 - 1 1

#### EXH ∃ T 4-6



SOURCE: E. J. Bentz & Associates, Springfield, Virginia

essentially the same range of variances found between early estimates and actual results - growth in costs have occurred of the order of 2-3 times the original estimate. The average of actual to estimated costs ( $\frac{Ca}{Ce}$ ), were found to be as follows:

System Type	Actual Cost/ Estimated Cost (Ratio)
Weapons System	1.40 - 1.89
Public Works	1.26 - 2.14
Major Construction	2.18
Energy Process Plants	2.53

The weapons system cost overruns were higher in the **1950's (1.89)** than in the 1960's (1.40) most likely, because of the greater degree of pioneering efforts and the greater lack of experience with large weapons systems at that time.

Exhibit 4-7 below shows the cost growth experience in pioneering energy systems as a function of the type of estimate employed (or available at that time). It can be seen that the preliminary estimates were nearly double that of the initial estimates - (84% above the first estimate) and the definitive estimates increased almost as much again from the preliminary estimates (134% above the first, or 50% above the preliminary estimate).

The ESCOE data were largely taken from preliminary estimates, based on Process Demonstration Unit (PDU) development experience, in one or two cases from pilot plant experience (at less than 1% scale) or from foreign commercial experience under different site and environmental conditions. The OTA shale oil values were derived from a very highly definitized engineering analysis. The degree of evolution which that estimate had undergone can be seen on Exhibit 4-E.

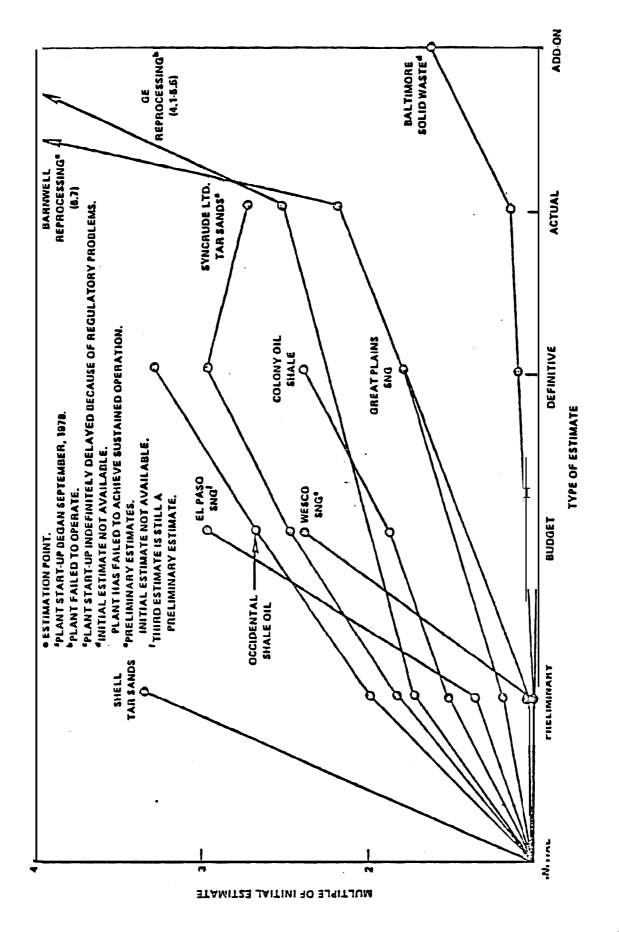
If the other ESCOE liquid synfuel plants were to increase by as much as have typically occurred between preliminary and definitive estimates, the costs would increase by about another 50%.<sup>12</sup> That would result in a shift of the cost line on Exhibit 4-6 as shown on Exhibit 4-9.

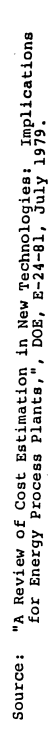
Such an interprelation of the quality of the ESCOE estimates would resolve the discrepancy between the ESCOE estimates and the OTA estimates (for oil, shale liquids) and produce a more continuous scale of synfuel cost relationships.

An alternative method of calibrating the various estimates for consistency with respect to the <u>status</u> of process <u>estimates</u> as well as the methods employed in the estimating process? would be to select

<sup>11</sup>Average increase from preliminary to definitive cost estimates for energy process plants.

 $12_{P3}$  - Reference 3.





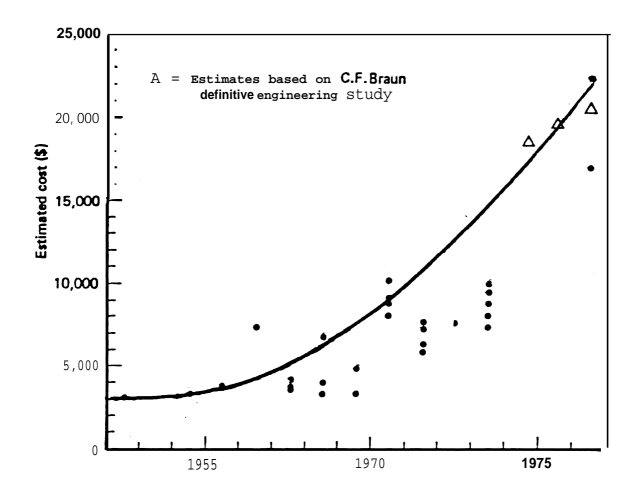
COST DROWTH IN PIONEER ENERGY PROCESS PLANTS (CONSTANT DOLLARS)

EXHIBIT 4-7

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4 - 74

#### HISTORY OF SHALE OIL CAPITAL COST ESTIMATES



Estimated surface shale oil facility contruction costs (capital costs/barrel/ca!endur day; constant \$ 1977)

a sub-set of processes that were developed on the basis of the same level of engineering definition or maturity - preferably the most advanced projects in this sense.

There have been more recent, updated design and estimating efforts undertaken in the case of:

1) Indirect Liquefaction - Mobil MTG. <sup>13</sup>

2) Methanol<sup>13</sup>

**3)** High BTU Gasification<sup>14</sup>

4) Direct Liquefaction - H-Coal<sup>15</sup>

These estimating efforts are essentially comparable with the (OTA) Oil Shale estimates in terms of the relative engineering and development maturity of the process plants involved.

Exhibit 4-9 also reflects the liquid fuel costs of 'generic" synfuel processes based on the selected "best estimates" noted above. These are not meant to be truly generalized processes (or generic processes), they are nonetheless representative, advanced members of each synthetic liquid product class.

The costs of these processes are shown in detail on Exhibit 4-10.

The effect of using the latest, or best estimates is approximately the same as was achieved by the use of the Rand Corp. (and others) cost estimating error factors. The original ESCOE values are increased by about 50% on average.

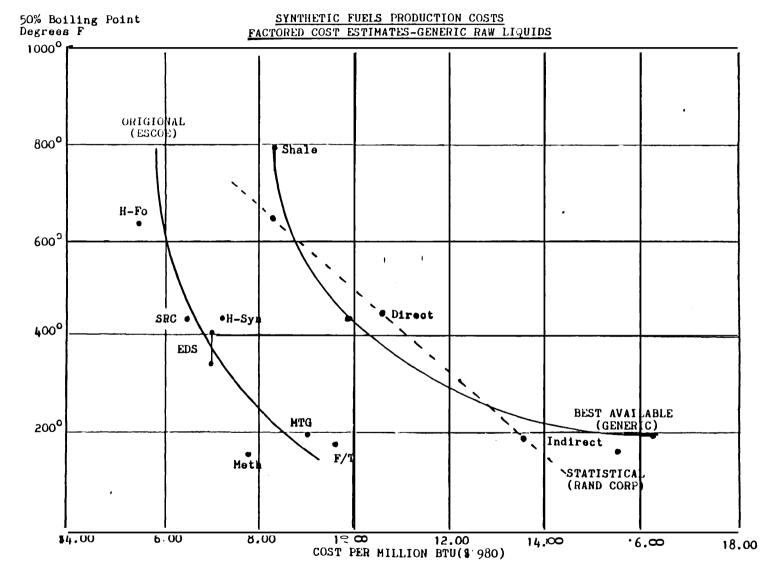
The satisfactory conjunction of factored cost estimates arrived at by the use of statistical variances derived from past estimating histories with the "generic" estimates taken from the most advanced projects, gives us an improved measure of confidence in the adjustment of ESCOE synfuel production costs to the higher levels displayed on Exhibits 4-9 and 4-10. The revised functional form of the liquid fuels is displaced to the right on Exhibit 4-9 by about \$3.00-\$4.00 per million BTU's. The relative costs are not appreciably affected considering the probable differences in residual (estimating) error contained in these estimates. It seems most reasonable, however, to presume that the majority of the estimating errors have been accounted for, and the values we are employing are normalized to the greatest practical degree possible at the present time: i.e., barring further engineering or demonstration plant design and construction experience.

13
Liquefaction Technology Assessment - Phase 1 ORNL-5664 Feb. 1981.
14
Unpublished Analyses
15

<sup>15</sup>Rand Corporation - Unpublished Analyses.

ejb&a

#### EXHIN T 4-9



SOURCE: E. J. Bentz & Associates, Springfield, Virginia

4-17

## BEST AVAILABLE ESTIMATES - TYPICAL SYNFUEL PROCESSES

	Revised Capital Est. \$ 1979	Revised Capital Est. \$ 1980	Capital Recovery 	Feedstock Cost (From 4-4)	Other* Oper. Costs (From 4-4)	Total Revised Estimate	Cost Per Barrel \$ 1980	Cost Per MM BTU \$ 1980	Cost 1 Gal. \$ 1980
Direct Liquids									
H Coal (Synfuel)	\$2 <b>,</b> 200	\$2,200	\$ 660	\$219.2	\$212.6	\$1,091.8	\$66.47	\$ 9.79	\$ 1.58
Indirect Liquids									
Mobil MTG (Lurgi)	2,685	3,054	916.2	205.3	159.8	1,281.3	78.01	16.18	1.86
SNG/Methanol (ICI- Lurgi)	1,849	21,035	631.1	256.9	160	1048 688. SNG 360.	11.88	15.53	1.00
High BTU Gas									
Lurgi (BGC)	1,600	1,820	546.	226.7	113.7	886.4		8.30	-

\*Adjusted f r capital cost changes.

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SOURCE: E. J. Bentz & Associates

The H-coal process (in the synfuel mode) has been used as a surrogate for direct coal liquids. Updated estimates of an unpublished nature were used that draw from the cumulative pilot plant histories and the most recent demonstration plant estimates. The Mobil Methanol-to-Gas (MTG) and methanol estimates were drawn from a recently published study by Fluor Corporation for Oak Ridge National Laboratory of indirect liquefaction processes. The study provided a (nearly) 100% gasoline option which virtually eliminates the by-product costing problems. The methanol estimates were joint production process schemes. The systems could Menthanol/SNG have been adjusted in keeping with the 100% gasoline MTG process scheme by eliminating the direct costs of methanol to gasoline stages. Alternately the by-product value of SNG could be directly priced by using the high BTU gas plant costs from the SNG estimate below. Both synthesis gas processes are Lurgi systems.

The SNG process estimate was taken from unpublished estimates drawn from advanced commercial design and estimating efforts. An advanced Lurgi gasifier - the British Gas Corporation slagging bed version - is used.

The costs of direct and indirect liquids - increase by about 50% - to remain in approximately the same relative cost relationship that the ESCOE based data displayed. The hi-BTU gas estimates only increased about 25% above the earlier ESCOE values. This appears to be reasonable considering the relatively more mature status of (Lurgi) gasification technology. The OTA oil shale liquids estimate of \$48.20\bbl reflects the precommercial stage of development. The level that we are attempting to standardize at, versus the development stage of the foregoing direct and indirect liquid systems.

#### Continuing Cost Escalation

The earlier analyses of Rand Corp. and others suggested that the potential cost increase from even a definitive estimate to the actual project costs of pioneer plants and major developmental systems is typically another <u>twenty percent</u> increase in cost. We can add that increment to arrive at an upper value for all systems.

There have been and continue to be other relevant post-commercial trends of commercial series production plants that were not considered by the authors of the cost escalation - studies cited above.

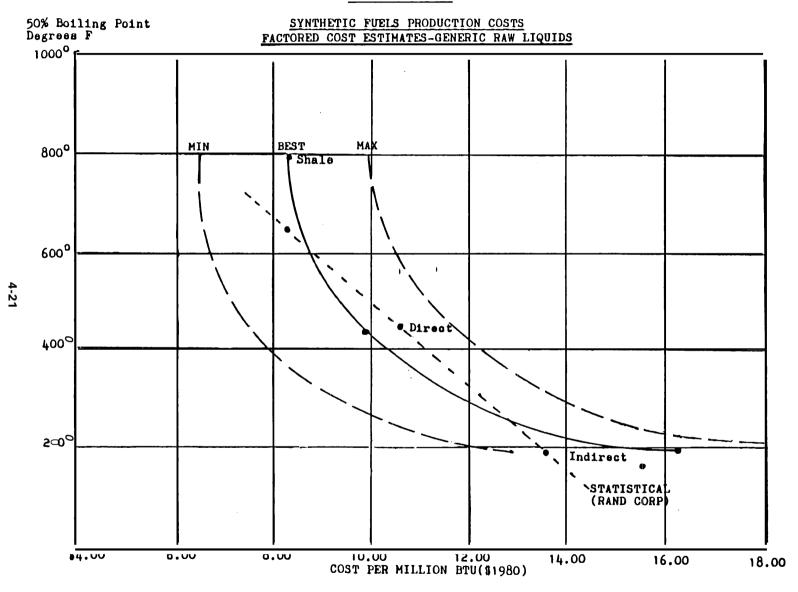
Historical data regarding the chemical process industry and petroleum refining industry demonstrates a strong pattern of capital productivity improvement or technology advance, during postdevelopment years. This can be demonstrated for the entire sector as well as in the micro-industrial setting of a single chemical industry segment.

A capital productivity rate of less then 2%/year can return the 20% (actual cost to definitive cost estimate potential increase during the first 10 years of commercial deployment. In 20 years at least a 35% redu ction in the capital outlay per barrel of product can be expected.<sup>16</sup>

These two viewpoints provide us with minimum and maximum estimates of the most probable range of expected production costs for synthetic fuels. Exhibit 4-11 illustrates the range of expected values for synfuel liquids based on these estimating limits.

This scale of values will be used to provide individual product (or by-product) costs. The presence of a significant amount of petroleum in the total supply equation, for as far as we can see, creates many cost and pricing complexities. We do not wish to complicate synthetic fuel supply economics with World Oil Price disruptions, or any free-market or administered market conditions. We will close our eyes to all of these dimensions and construct our cost schedule on the basis of coal based liquid, gas and solid fuel options or opportunity costs.

<sup>16</sup> This rate (1.4%) has been experienced by the <u>entire</u> chemical industry throughout the entire post war period (1949 to date). Specific industry sectors have experienced much greater rates of productivity improvement; viz, synthetic methanol experienced more than a 4% / year productivity gain for over 20 years.



SOURCE: E. J. Bentz & Associates, Springfield, Virginia

4.5 PRODUCT UPGRADING (References 39, 40)

The typical (direct liquefaction) coal liquids possess several characteristics that require upgrading in order to:

- . Provide product stability
- Permit mixture with conventional petroleum liquids . . . or
- Permit common use of pipelines and other infrastructure

The principal differences result from:

Lower levels of hydrogen - 9 - 10% versus 11 14% for petroleum and 11 - 12% for shale oils.

<u>Higher levels of heteroatoms</u> in both liquids and shale oil (nitrogen and oxygen compounds) than are found in petroleum feedstocks.

The lower hydrogen and higher heteroatom conditions are resolved together by hydrotreatment. Raising the hydrogen levels up above 10% results in the removal of most of the nitrogen and oxygen heteroatoms, and also decreases the aromaticity of the coal liquids and shale oils.

The high aromatic content of coal liquids makes the naphthas excellent high octane blending stock - however the high nitrogen and oxygen percent (2 - 3%) in the heavy naphtha range requires the use of fairly severe hydrotreatment to remove the diolefins and heteratoms - which are present in the form of phenols and cresols (oxygen).

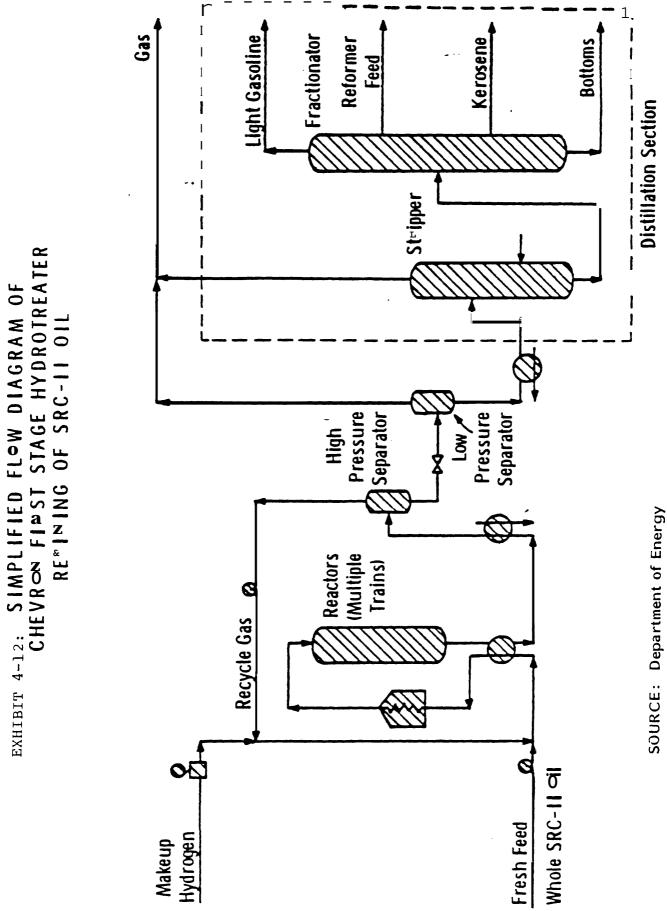
In the synfuel <u>distillates</u> the nitrogen level is higher and results in unstable compounds with rapid gum formation, making this **a very** unsatisfactory fuel unless upgraded.

There have been a succession of studies of synthetic liquids upgrading processes sponsored by DOE. They have been conducted on both shale oil and direct coal liquids.

The principal measures examined include:

- Hydrotreating (Exhibit 4-12)
- Hydrocracking
- . Fluid Catalytic Cracking

Catalytic reforming as well as hydrocracking are subsequently used to upgrade (naphthas) to finished transportation fuels. (See Exhibit 4-12 below) .



Consideration has also been given to variation in the hydrogen source for hydrotreaters - the partial oxidation of raw coal liquids, reforming of refinery products and overheads, or outside gasses.

An additional issue is the location of upgrading facilities; at the **coal** liquids (or shale oil - retort) plant, or at a conventional refinery, or both.

The factors which favor the synthetic oil plant location are:

- . available residue for hydrogen manufacturing
- . local upgrading permits common carrier transportation
- upgraded synthetic product can be blended with petroleum feedstock (in pipelines and at refineries)

The factors that favor a refinery location for upgrading are:

- . Superior prospects for system optimization
- . Availability of hydrogen from naphtha reformers
- . Uses available refinery capacity idled by lack of petroleum feedstock.

An alternative approach could be to perform a minimum amount of upgrading at the synfuels plant to facilitate transportation and storage, with product finishing and blending performed at a larger refinery site. The coal liquids <u>in general</u> do not require further cracking because they lie in the atmospheric gas-oil and naphtha range. <u>The shale oils require cracking to produce more usable</u> <u>product from the higher distillate</u> range such as jet fuel and dieseloils. The heavy distillates from coal liquids; if heavily hydrotreated (to 11% H bywt) can be used **as** a feedstock for a fluid catalytic cracker (FCC) where the product can be significantly upgraded.

Exhibit **4-13** illustrates the cost of upgrading various direct liquid process cuts.

The raw liquids versus the upgraded liquids are compared below in hydrogen content.

	Raw Liquid	Upgraded
SRC Naphtha	11.33%	11.6%
SRC Distill.	7.71	11.0
H Coal Distillate	10. 1	11.4
H Coal Fuel Oil	7.37	10.0

These cases cover the general conditions experienced by the range of most direct coal liquids - the samples being drawn from experimental laboratory investigations performed by Mobil Research and Development Corporation upon SRC light and heavy fractions and

		50 000 BBL) – 1980 \$			
		SRC-II	H Coal S	Svn. Crude or Fuel	Fuel Oil
	Naphtha	Heavy Distillate	Naphtha	<u>Distillate</u>	Fuel Oil
ч	<b>.</b> 0639 *1606	•0855 *5830	.0639 °1606	.086 ∍409	<b>.</b> 085 *593
Administration & Support G&A	.0394 .1828	• •	.0394	.077 .420	.100
	.4467	1.3434	.4467	.992	1.363
Fuel Utilities	.3517	.6691' .9716	.3517 .1089	.525	.154 .946
Cat. & Chem. Hydrogen	.0194	.9829 6.836	.0194 .5586	1.650 2.390	1.358 7,355
1	<b>1.0</b> 38€	9.4895	1.6386	5.142	9.813
Capital Recovery (30%)	2.577	8.484	2.577	5.449	7.933
Total Upgradiog Cost	4.062	19.32	<b>4</b> .o≤23	11.58	19.109
Product					
API H Content (wt %) BTU/lb.	37.5 11.6 18,500	24.5 11.0 18,780	Same as SRC II Naphtha	25.7 11.4 18,970	12.5 10.0 18,400
Plant <sup>±</sup> nvestment Total in Millior \$/BBL	\$141 \$8.59	\$464.5 \$28.28	\$141 \$8.59	\$298 \$18.16	\$434 \$26.44

DIRECT LIQUIDS UPGRADING COST

50 000 BBL) - 1980 \$

SoURCE: E. J. Bentz & Associates

H-coal distillate and fuel oil fractions. This pretty well covers the range of liquids produced by SRC and H-Coal (synfuel and fuel oil mode) and can be extrapolated to the EDS case.

Additional work performed by U.O.P., Chevron and Suntech confirm the general upgrading needs and the best approach - hydrotreatment.

The plant investment required varies from \$140 million dollars for the mild hydrotreatment required of the naphtha cuts ( $C_{\rm s}$  - 400°F) to as much as \$465 million for a hydrotreatment plant for the heavy distillate or residual SRC fraction and nearly that for the fuel oil fraction of H Coal fuel oil process plants.

The average upgrading cost is about \$2.00 per million BTU's varying from \$4.00-to <u>nearly \$20.00</u> per barrel. The latter figure represents an economic limit which suggests either a lower grade utilization of the heavier products or a different refining approach.

The direct liquids upgrading cost analysis can be compressed to a single representative-or "generic" upgraded coal liquid.

The general costs of upgrading are shown on Exhibit 4-14:

Naphtha's	<b>\$ 4.06</b>
L. Distillates	11.58
Heavy Distillates `- Fuel Oil	19.21 (19.11-19.32)

Individual processes such as EDS SRC-II and H-Coal (fuel oil mode) will differ in raw liquid base costs, but since the quality of product tends to vary in a reasonable relationship to their costs<sup>17</sup>, the costs of upgrading, which are <u>increasly related to</u> <u>quality</u>, lend to cause a clustering of upgraded direct llquid costs.

If we utilize the costs of H Coal production of raw liquids developed above as a base, the 'generic' costs for upgraded products would be as follows on Exhibit 4-14. The estimated costs of nearly \$75.00 per barrel or over \$12.00 per million btu's is for a product that is equivalent to a <u>high grade</u> refining crude oil feed.

The upgrading of shale oil to a suitable refinery syncrude has been estimated by Chevron to cost \$10.00 per barrel (in 1980 dollars) or \$1.72 per million btu. If this is added to the cost of raw shale-oil liquids at the retort, the total cost of shale oil "syncrude" is:

<sup>17</sup>See Exhibit 4-6 above.

# DIRECT LIQUIDS UPGRADED COSTS/BARREL

## (\$ 1980)

	Barrels/Day	Cost/BBL	Total Daily Cost
Naphtha	28,380	\$ 4.06	\$115,223
Distillate	21,620	11.58	<u>25° 36°</u>
	50,000		\$365,583 ⊨ \$7.31 Avg.
Raw Liquid Cost (per barrel)			66.47
Total Upgraded Fuel Cost Per Barrel			\$73.78
Total Upgraded Fuel Cost Per MM/BTU			\$12.30

4-27

SOURCE: E. J. Bentz & Associates

#### OIL SHALE LIQUIDS COST

(\$1980)

	Per Barrel	Per Million BTU
Retorted Shale Oil	\$48.20	\$ 8.31
Upgrading	10.00	1.72
	\$58.20	\$10.03

These compare favorably with upgraded direct liquefaction production in the 'syncrude' class as shown below:

SYNCRUDE PRODUCTION COSTS	
(\$1980)	
Per Barrel	Per Million BTU

Shale Oil	\$58.20	\$10.02
Direct Coal Liquids	21.12	18.5%
Shale Oil Advantage	12%	9%

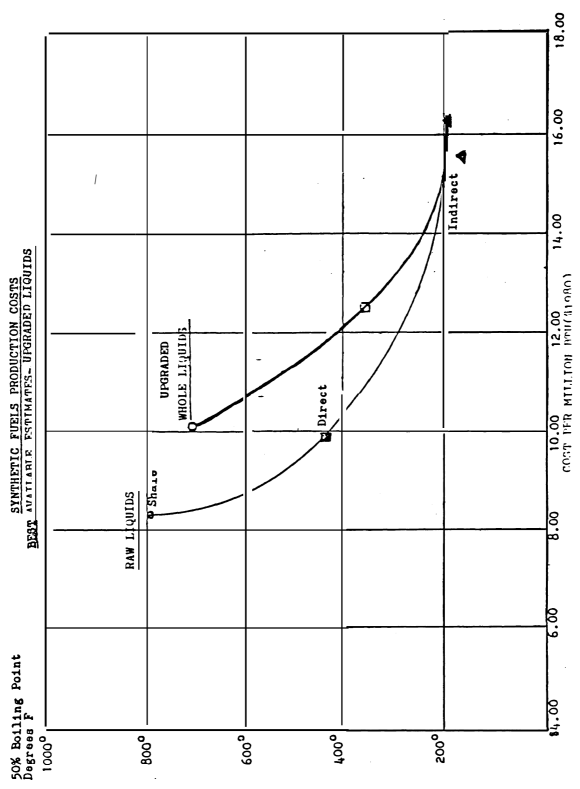
The shale oil has about a 21%-cost advantage as a refinery feed-Stock. This is reduced to less then a 20% cost advantage on a heating value basis. However heating values are not the principal criterion to be applied to refinery feedstocks - quite the opposite - the lighter crude demands a premium. In certain instances the coal liquid with higher aromatic content will be preferred, at other refineries the shale oil, with a higher hydrogen content, and a greater yield of distillate product will be sought.

Exhibit 4-15 illustrates how the process of upgrading shifts the cost of oil shale and coal based synthetic crudes upward by \$1.75 - 2.50 per barrel.

#### 4.6 REFINING SYNTHETIC LIQUIDS

The direct liquefaction and oil shale synfuels have to be further upgraded to end-use product quality in order to be comparable with indirect liquid products such as methanol from coal or gasoline from methanol (from coal). In a wider sense, this is also desirable in order to achieve comparability with synthetic natural gas (SNG) which can be used for a wide range of end use applications in its 'raw' manufactured state.

The indirect processes produce refinery output (or intermediate) grade products, without the need for the "refining" of crude liquids. In order to compare direct liquids and shale liquids with indirect process liquids, we must bring the former EXHIB T 4-15



SOURCE: E. J. Bentz & Associates, Springfield, Virginia

into a state that is comparable. This requires the refining of the synthetic liquids to finished fuels.

Refining of shale oils and coal liquids will vary in cost depending upon the size, location and degree of integration of the refinery complex. We will assume that this is not done in an existing refinery (perhaps modified to better handle these feedstocks), but is performed at a new refinery integrated at the retort or conversion plant site. Such a refinery is under-scale (50,000 bbl/day) and remote from chemical complexes that might make better use of by-products and hence provide higher (by-product) credits or other similar economic benefits.

The costs of upgrading the raw coal and shale liquids to high grade (transportation) fuels is shown below:

#### REFINERY COSTS FOR SYNTHETIC (RAW) LIQUIDS

(\$1980)

	<u>Cost Per Barrel</u>	Cost Per Million BTU
<u>Shale Oil</u> (Hydrotreat & Hydrocrack)	\$18.50	\$3.19
Coal Liquids (Hydrotreat)	\$18.29	\$4.02

The costs of refining synthetic liquids cannot truly be determined without specifying the product slate produced. The costs of refining a particular feedstock can vary depending upon the product cuts sought. The basis used above is not strictly comparable between the processes. It tends to slant the refinery approach to the type of slate that is favored by the feedstock - Light distillates in the case of shale oil, and gasolines and distillates in the case of coal liquids.

Exhibit 4-16 illustrates the potential variation.

These costs can be seen to vary dramatically if different product slates are sought. If the highest grade transportation fuels are maximized, to provide the highest degree of comparability with indirect liquids. The costs are as follows:

#### REFINERY SYNTHETIC UNITS TO 100% TRANSPORTATION FUEL

		(\$ 1980)			
		\$/BBL \$/MM BTU		<u>Coal</u> \$/BBL <u>\$</u> /MM BTU	
Raw Liq	uid	\$48.20	\$ 8.31	\$66.47	\$ 9.79
Upgradir	ıg	18.50	N.A.	18.28	N.A.
	Total	\$66.70	- \$11.50	\$84.75 _	- \$14.61
Average BBL	Heat Content\	5.8 Million BTU		5.8 Mi	llion BTU
		4-30	)		ejb&a

## PROCESS AND SLATE

# (1980 \$)

Feedstock	Hydrotreat & Hydrocrack	Severe Hydrotreat	Moderate Hydrotreat
Coal Liquids* Product	Motor Gasoline	Motor Gasoline Plus Jet Fuel	Motor Gasoline Plus #2 Fuel Oil
Slate	(100%)	(1/3 - 2/3)	(1/3 - 2/3)
Cost	\$20.70	\$18.29	\$12.55
Shale Liquids	Hydrotreat & Hydrocrack	Hydrotreat-FCC	Coking Hydrotreat
	3/4 - 1/4		(4/5 - 1/5)
Product Slate	Motor Gasoline Plus Jet Fuel	Jet Fuel Plus Motor Gasoline	Jet Fuel
Cost/BBL	\$18.50	\$17.00	\$16.00

4-31

\*SRC-II

SOURCE: E. J. Bentz & Associates

By comparison, indirect liquid (methanol to gasoline) costs are about \$78.00 per barrel; approximately in the middle of this range. The cost per million BTU's is lower for shale and coal liquids, refined to a transportation slate consisting of gasoline and distillate fuels (jet fuel and diesel oil). If direct liquids are refined to a 100% gasoline slate the costs would increase to \$87.17 per barrel or above \$19.00 per million BTU's.

Exhibit 4-17 graphically displays the finished fuels in a framework which relates the product quality to the finished fuel cost.

Exhibit 4-18 calculates the total cost of refining coal liquids. A 50,000 barrel per day refinery for coal liquids would cost between \$420 million and \$690 million. The lower case represents a moderate hydrotreatment plant producing #2 fuel oil and gasoline, the upper case represents a hydrotreatment and hydrocracking plant that produces 100% gasoline.

Instead of using other indirect measures of product value, <sup>18</sup> we can use a cost based scale. The lighter fractions cost more to produce from both coal and shale, whether by direct or indirect means. By-product credits do not have to be assigned to determine the cost of a single cut liquid. Upgrading plant has been assigned to individual fractions so that the full cost of the beneficiated product cut is known. The costs of fully refining the product are developed incrementally by determining the cost of creating a 100% gasoline yield, and two subsequently lower grade mixtures.

The alternate product slate refinery costs of Exhibit 4-18 can be used to develop a measurement of the direct costs of products in a multi-product refinery run. The principal cost differences result from the increased capital (per unit of product yielded) and the increased consumption of hydrogen associated with higher grade product slates.

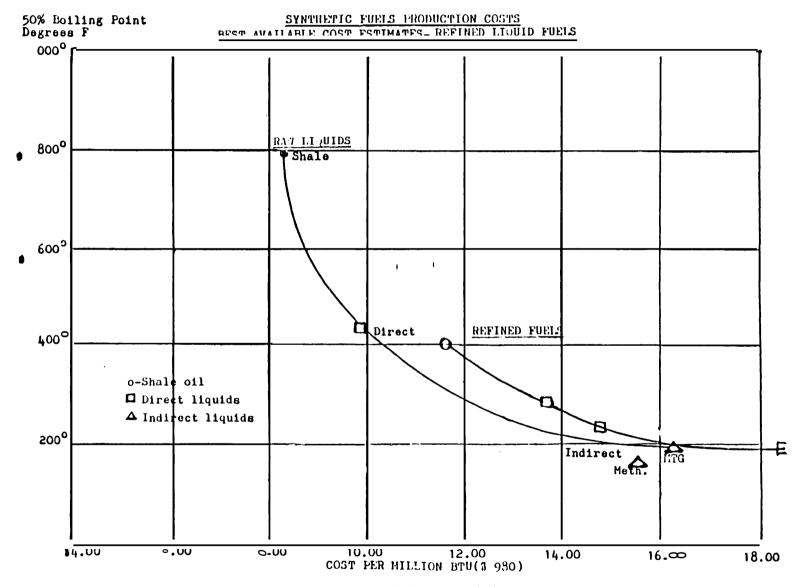
If we take the per barrel cost of producing a 100% gasoline slate. and assign it to the gasoline fraction of a mixed slate as the appropriate cost of that portion of the output, the remainder of the total cost divided by the number of barrels of the other product (jet fuel or #2 fuel oil) will give us the unit cost of the "secondary product".

Exhibit **4-19** shows this costing procedure for the slates presented for direct liquids refining in Exhibit 4-17.

By using this method, we are not artificially lowering the cost of gasoline production by assuming a market equilibrium price

4-32

<sup>&</sup>lt;sup>18</sup>Product value ratios are commonly used. They are of absolutely no meaning in a long-term and discontinuous supply context. The use of such ratios is a major violation of the most elementary laws or principles of economics as a measure of utility.



SOURCE: E. J. Bentz & Associates, Springfield, Virginia

4-33

#### DIRECT LIQUIDS (SRC-II) REFINING

(50,000 BBL/Day) 1980 \$ per BBL.

	· · · · -	-	
\$/BBL	Motor Gasoline (Hydrotreat Plus Hydrocrack)	Motor Gasoline <u>Plus Jet Fuel</u> (severe Hydrotreating)	Motor Gasoline <u>Plus #2 Oil</u> (Moderate Hydrotreat)
Operating Labor	.244	.183	.183
Maintenance	.791	.669	.487
G&A	.852 1.887	.670 1.522	.487 1.157
Fuel	.183	.304	.122
Utilities	.183	.122	.122
Cat. & Chem.	.304	.365	.244
Hydrogen	5.540 6.210	5.750 6.541	3.230 3.718
Capital Recovery	<u>12.603</u>	<u>10.228</u>	<u>7.67</u>
@ 30%/Yr.	\$20.70	\$18.291	\$12.545

SOURCE: E. J. Bentz & Associates

# SRC II REFINED TO PRODUCT COSTS

	H	Barrels/Dav		Cost/BBL*	Total Daily Cost
CASE <sup>±</sup>	Motor Gasoline	50,000	6	\$87.17	\$4,358,500
CASE <sup>⊥</sup> <sup>⊥</sup>	Motor Gasoline	15,395	6	87.17	1,341,982
	Jet Fuel	34,605	6	(83.69 **	( <u>2,896,018</u> )**
	Total	50,000	, <b>6</b>	\$84.76	\$4,238,000
CASE III	Motor Gasoline	16,995	6	\$87.17	\$1,481,454
	#2 Oil	33,005	6	( <u>74.74</u> ) **	( <u>2,466,796</u> )**
	Total	50,000	6	\$78.965	\$3,948,250
	Product Costs	Notor Co	- 1		
		Jet Fuel	sorrue		4.95) \$17.61/M BTU 5.67) \$14.36/MM BTU
		#2 Oil			.825) \$12.83/MM BTU
*Cost from	**Values average	in pa valu	renthesis infern Ne of motor gasol	red from weighted line and total product.	
SOURCE: E. J. Bentz & Associates					

,

for a lower grade (by) product. The method used is entirely an assignment of marginal cost to products. It would be more desirable to operate in a reverse manner, i.e., from the lowest product, assigning incremental costs to the higher product on a marginal basis. We, unfortunately, do not have a process estimate for a single slate of the lowest value product. The distillation range of all products is too broad to produce such an artificiality. Therefore we have begun with the marginal gasoline cost and assigned it as a by-product price to the lower value (mixed) slates, permitting us to infer the marginal cost of the lower grade products.

The results of this cost analysis are related to the costs of indirect liquefaction end products and shale products on Exhibit 4-20. The cost series increase as average distillation point is lowered. The average distillation point of most useful transportation fuels lies between 180° - 400 F, with the majority of the compounds contained lying within this range.

There is a persistence of the earlier noted relationship between product quality (as measured by average boiling point) and production costs of finished products. The relationship shows less than unitary cost increases per barrel, all greater then unitary cost increases per million BTU. The latter case is due to the generally lower heating value of the premier fuels that have increased hydrogen content. The increases in cost are about 7 1/2cents per barrel of liquids for every degree farenheit that the boiling range is lowered.

Exhibit 4-21 is a flow sheet of a process (examined by Chevron Research) for hydrotreating and hydrocracking of direct coal liquid (SRC-II) whole oil to produce 100% motor gasoline product. This is the first case on Exhibit 4-16. Exhibits 4-22 and 4-23 illustrate the refining process used to upgrade the whole liquid to ' gasoline and jet fuel by severe hydrotreating alone, and to a lower quality slate of gasoline and heating oil created by less severe hydrotreating of direct (SRC-II) liquids.

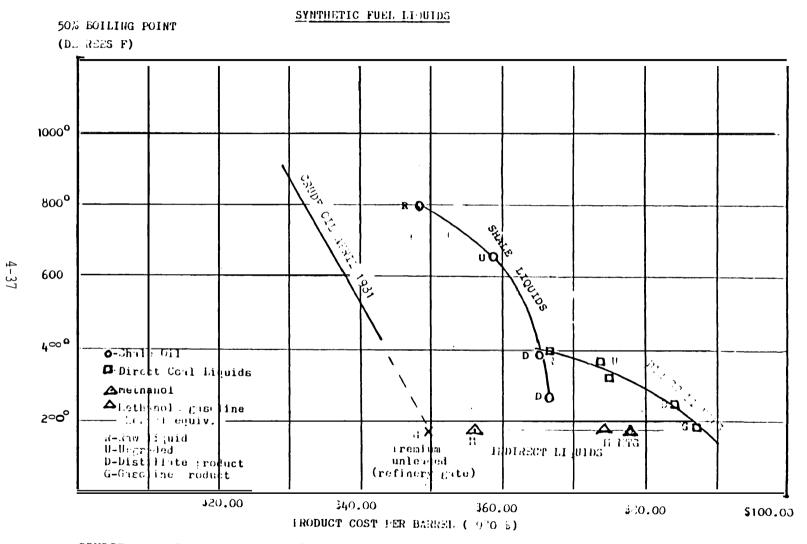
The latter case is more comparable to an upgrading process.

#### 4.7 TRANSPORTATION AND OTHER INFRASTRUCTURE COSTS (Reference 41)

Although we have differentiated between coal liquid's plant site upgrading facilities and finished product refineries, we have really not selected the site for refining. The upgrading must in most cases be done at the site of the coal liquids plant. The degree of upgrading we have embraced (Exhibit 4-15) is sufficient to permit the fuels to be used in as high a use as a combustion turbine, or transported without creating contamination or incompatible sediments.

Transportation costs are directly related to the distance involved, and indirectly related to the quantity moved or flow rate.



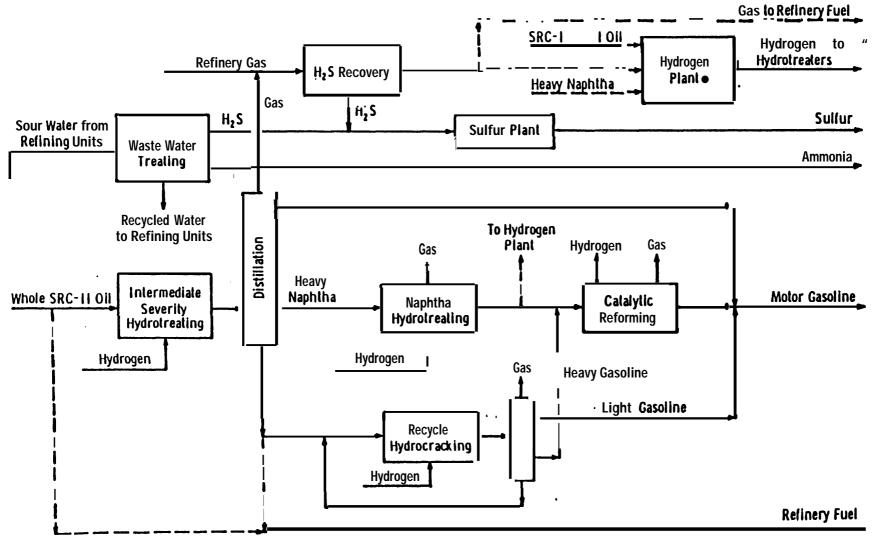


SOURCE: E. J. Bentz & Associates, Springfield, Virginia

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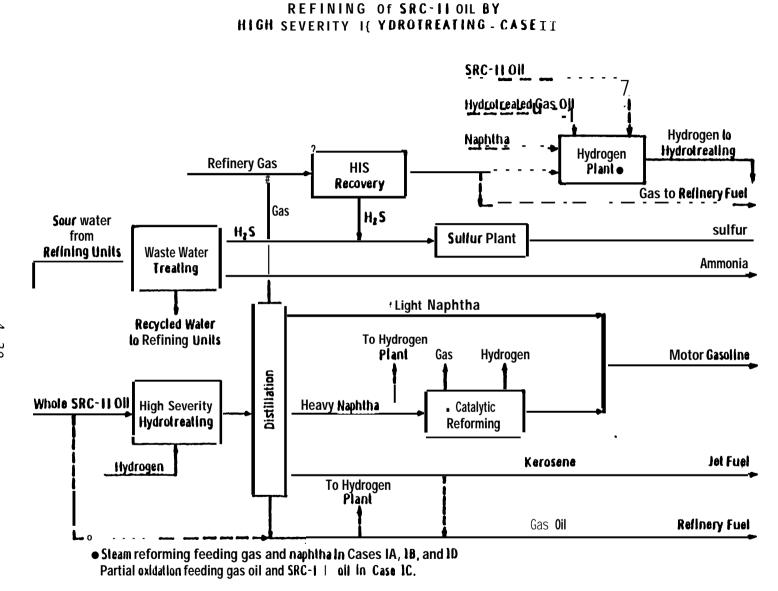
#### EXHIBIT 4-21: SCHEMATIC FLOW DIAGRAM REFINING OF SRC-11 OIL BY . HYDROTREATING AND HYDROCRACKING - CASE I

.



• Sleam reforming feeding gas and anphtha in Cases 4A, 4B, and 4D. Partial oxidation feeding SRC-II oil in Case 4C.

4-38



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SCHEMATIC F10W DIAGRAM

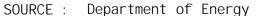
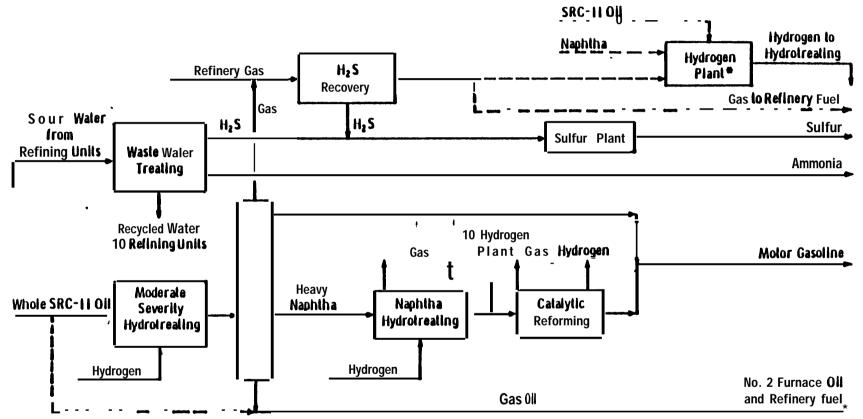


EXHIBIT 4-22:

4-39

ejb&a

EXHIBIT 4-23: SIMPLIFIED F10W DIAGRAM REFINING OF SRC-IIOIL BY MODERATE SEVERITY HYDROTREATING - CASE III



• Sleam reforming feeding gas and naphtha In Cases 5A, 5B, and SD. Partial oxidation feeding SRC-110111n Case SC.

SOURCE : Department of Energy

4-40

We cannot visualize any other form of transportation for these upgraded liquids, or for further refined products except by pipeline. The daily volume required to support a 6" or 8" pipeline is approximately the size of one or two 50,000 bbl/day plants. Considering the geographical concentration of coal and shale deposits it is not difficult to visualize a mining-conversion center adequate to support either:

. An upgraded liquids pipeline to a refining center

or

. A product pipeline to major pipeline junctions or product distribution terminals

The general location of all coal and shale resources is such that deep draft water transportation does not figure prominently in synfuels distribution patterns.

Without siting specific plants and conducting the refinery trade-offs - which would have to be done in context with both the balance of foreign and domestic petroleum supplies and the slate of (regional) demand for all liquids - we cannot develop very meaningful insights into either the operating (product) costs of transportation and distribution, or the capital requirements.

We will have to make some nominal assumptions and then establish unitary relationships. The future energy transportation patterns and infrastructure requirements are impossible to determine without a specific scenario. We shall briefly examine a \*cases:

- . Pipelining from Souther Illinois to Houston of syncrudes.
- . Pipelining from Wyoming to St. Louis
- . Pipelining from Western Colorado to L.A. of shale oil.

Southern Illinois to Houston

Raw Liquids<br/>(upgraded)33c/MM BTUWestern Colorado to L.A.<br/>Shale Liquids40\$/MM BTUWyomina to St. Louis8TURaw Liquids<br/>m ' \* ' 30 \$/MN! BTUMethanol68c/MM BTUMTG - Gasoline37\$/MM BTU

The additional capital investment required for synthetic fuel transportation is highly speculative to a greater degree. There

is a great deal of existing product and crude liquid pipeline as well as gas pipeline in place, that can equally serve the synthetic fuels industry. In all cases the pipelines are connected to either markets or distribution terminals at the delivery end. In most cases, the input end is originally either at a major refinery (and production) location or at a port location. The refinery connection argues for upgrading of liquids (coal and shale) at mine mouth conversion plant locations, and transportation to the existing refinery districts for product finishing. Such a general pattern would involve the construction of a minimum number of new "crude" synfuel pipelines from coal fields to refining districts.

We assume that the ultimate conditions would lead to the construction of several large diameter pipelines in such a pattern.

Methanol, which does not require refining, obviously will move in different patterns from coal field to the major terminals and markets.

Pipelines of that size (10-12") would cost an average of \$100,000 per mile, considering material, labor, and right of way and other expenses. Terrain would influence the cost, generally increasing construction costs but reducing right of way costs in some cases by an equivalent amount. 20" or greater diameter pipelines would cost \$250,000/mile.

A total construction budget of **50,000** miles of new pipeline of 12" diameter to 20" diameter would cost between \$5 billion and \$12 billion.

4.8 ADDENDUM TO CHAPTER 4: BASIS FOR COST ASSUMPTIONS

1) Basic Conversion Plant (ESCOE)

•Capital Costs

Year: Mid (June-July) 1979 dollars Scale: 25,000 tons of coal input Base Plant to installed battery limits: 1.63 Contingency: 10% Scaling exponential rule: C<sub>2</sub> = G ۲<sup>۲</sup>  $\lambda$  = .65 for vessel size  $\lambda$  = .9 with trains Outlay of Capital: instantaneous plant • Revisions to Capital Assumptions in This Report Year: Mid 1980 (June-July) Scale: 50,000 bbl/day liquids output Plant to Battery Limits: 1.73 Contingency: 20% Scaling: Linear Outlay of Capital: Instantaneous plant • Operating costs Coal Feedstock: \$30/ton (delivered) Coal: Illinois #6 Catalysts and Chemicals and Operating Supplies: at cost for amounts proscribed by process designer's material balance. •Labor Cost # Rate/Hr 120 \$ 10.00 Plant Operators Operating Supervisors 25 15.00 150 12.00 Maintenance Labor Maintenance Labor Supervisors 30 16.00 Administration 11.00 30 Total @ \$11.79/hr avg. 355 Fringes @ 35% -- changed to 40% = total labor rate of \$16.50/hr

Maintenance Cost (Materials & Contracts)

3% of total plant capital cost

<u>G & A</u>

Local taxes and insurance, 5% capital cost changed to total G&A - 5% capital cost

Capital Charge Rate

ESCOE basis not used. 30% of capital used as recovery rate (as per guidance of OTA staff) .

On-Stream Rate

90%--328.5 days/year

- 2. Assumptions for Product Upgrading
  - <u>Capital</u>

Basis -- Instantaneous Plant, mid-1980 dollars On-stream factor 90% 328.5 stream days.

• Hydrotreater

capitalized for each separate product stream.

• Hydrogen Feedstock Plant Capital

Not included, only cost feedstock "across the fence" from the plant complex.

- <u>Hydrogen Reformer</u> or manufacturing plant capital included
- Battery Limits

Includes hydrotreaters, waste water treatment, sulphur plants (commercial grade)

#### • Contingency

General -- 25% Battery Limits--15% Engineer---4% of investment capitalized Working Capital--45 days receivables; 30 day chemicals catalysts; 30 day feedstocks • Operating costs

<u>Hydrogen Feedstock</u> :	Syngas @ \$6.74/mmbtu raw gas liquids @ \$6=50/mmbtu includes recovery of production plant capital.
Hydroqen Pressure:	500 PSIG for SRC light (naptha) product2000 PSIG all other cases.
<u>Plant Size</u> :	<b>20,000</b> bbl/day upgraded to 50,000 bbl/day <u>for each product</u> <u>cut</u>

• Royalties

500 PSIG Hydrotreating-o-1500 PSIG Hydrotreating Fixed Bed\$30/bst feedSulphur plant-o-Waste Water\$75,000First 5,000 units\$14.70\unitNext 5000-25,000 units\$7.35/unitNext 25,000 + units\$5.25/unit

• <u>Sales Tax</u>

5% of equipment cost

• Maintenance

4% of depreciated capital/year

• Operating Labor

\$11.00/hr

• Labor Burden

45%

• Administrative and Support Labor

30% of operations and maintenance labor

• <u>G & A</u>

60% of operations and maintenance labor plus property-tax of 2-1/2% of plant investment

• <u>Utilities</u>

Fuel \$4/mmbut Steam \$3.50/1000 lbs Electricity 4c/kwh Water (make-up) 40c/1000 gal

• <u>Hydrogen Bleed</u> was assumed to be: 50 SCP/bbl @ 500 PSIG

100 SCP/bbl @ 2000 PSIG

By-product Credits

Ammonia (anhydrous) 100/tonHydrogen and Hydrocarbon off gasses (C<sub>1</sub>-C<sub>4</sub>) 4/mmbtu (\$1. **30/MSCF**)

- 3. Refining Cost Assumptions (Chevron Basis)
  - 1980 costs: Instantaneous plant (first quarter adjusted to June/July)
  - Mid-Continent Location
  - Cost correlations based on actual experience of Standard Oil of California, 1960-1970s adjusted for:

Lower field productivity Increased safety Improved efficiency and reliability Additional energy conservation Stricter environmental regulations

- 10% Contingency
- Utilities

Water 30c/1000 gal Boiler fuel, coal or refinery fuel power 3\$/kwh

• Maintenance

 $2\text{-}1/2\/yr$  of both on-plant and off-plant facility investment

• G&A

Property taxes @ 21/2% of both on-plant and off-plant/yr

• Labor

Operating-- \$110,000 per shift position/hr (\$18.30/hr including fringes) Support Labor (Administrative, security, technician) 65% of Direct Labor CHAPTER 5: SUPPLY DEPLOYMENT SCENARIOS FOR SYNTHETIC FUELS

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#### CHAPTER 5 : SUPPLY DEPLOYMENT SCENARIOS FOR SYNTHETIC FUELS

#### 5.1 Factors and Constraints Affecting Synfuel Development

In order for synthetic fuels to play a role in increasing domestic energy supplies, they must become available in sufficient quantities, at competitive prices, in a reasonable time frame. This is particularly true for transportation's needs for liquid fuels. With a relative lack of fuel switching capability, transportation more than other sectors (e.g., utility fuel switching to coal) must depend on increased conservation, expanded domestic crude production, and alternate liquid fuels.

The central <u>driving forces</u> that characterize the development of a synthetic fuel industry are (Reference No.  $4 \ 2$ ):

- (a) Depletion and cost escalation of conventional domestic energy supplies;
- (b) Shortages of environmentally acceptable fuels;
- (c) Constraints imposed on alternate energy systems;
- (d) The presence of existing, easily modified fuel distribution systems;
- (e) A seemingly chronic negative imbalance in foreign trade and payments accounts;
- (f) National security; and
- (g) Governmental incentives (such as those proposed under P.L. 96-126 and the National Energy Security Act) .

The central concerns are:

- (a) Technological and economic factors
  - product costs/markets (interfuel competition) Status of technology and technological risk Financial risk Capital availability
- (b) Environmental and social factors
  - Air quality
  - Water quality
  - Land reclamation
  - Social dislocation

- (c) Availability of resources
  - Energy resources
  - Water resources
  - Land/site availability
  - Skilled work force
- (d) National, State, and local policies, especially regulatory, taxation, and subsidy policies.

Key among the <u>requirements that characterize these concerns</u> are:

- (a) Technological needs
- (b) Significant lead times
- (c) Relative costs

In Chapter 3, we have looked at the technological needs; and in Chapter 4 we have looked at the relative In this chapter we will focus on the "staging" costs. over time of these technologies, so that we can appreciate the necessary lead times. In doing so we will attempt to develop realistic "bottoms-up" assessments for each generic fuel class. These scenario will be a 'business-as-usual" assessment, and a high "pushing-the-limit" assessment. b In developing these scenarios we have felt it crucial to build upon concrete actual data and engineering plans for each project class, rather than "top-down" estimates of aggregate growth." We also felt it necessary, as explained in the introduction, to limit our supply deployment scenarios to the year 2000, which reflects the upper limit of sound engineering judgment and actual/proposed plans. Post 2000 considerations are more dictated by an assessment of economic forces and prospective product markets rather than supply constraints.d The supply constraining forces of the "transition" period (1980-2000) reflect industrial 'build-up" times and constraints, rather than product demand shifts." Post 2000 considerations must consider demand shifts, end-use technology changes, and the introduction of other technologies (e.g., solar) .<sup>r</sup> This necessitates a macro-economic long-term forecast approach rather than a supply deployment scenario approach.

Because of the significance of "transition" period<sup>h</sup> constraints in realizing deployment schedules, it is useful to discuss these constraints prior to our development of the scenarios. In the following section we will discuss the key constraints. Following this discussion, we will present the actual assessments developed and compare them with other assessments referenced in the literature.

# 5.2 Constraining Factors in the Transition Period: 1980-2000

The construction of one 50,000 barrel per day synthetic fuel facility is a massive effort requiring huge dollar, manpower, and material inputs plus the management skills to integrate all these inputs into a workable system. Constructing a major synfuels industry multiplies the problems, introduces added complexity, and increases the probability that constraints of varying degrees will impact the schedule, cost or feasibility of success.

Any Us. proposed synfuels construction program will have to compete for manpower and other resources with related construction demands from the oil and chemical fields. U.S. refineries are undertaking a major upgrading program to enable existing refineries to handle lower grade high sulfur crude and to increase efficiency in producing full product slates with less energy waste. Fluor Corporation is predicting that U.S. refineries will initiate \$20 billion in construction programs in 1980, contrasted with a yearly average of only \$2 billion in the late 1970s. (Reference No. 43 ) Proceeding with the Alaskan Natural Gas Pipeline could require \$20 to \$25 billion in new construction costs. Similarly, the chemical industry is modifying its petrochemical plants in recognition of dramatically higher feedstock costs. The situation is further compounded by gigantic increases in construction programs abroad. For example, Saudi Arabia appears intent on pursuing a five year \$335 billion program of new refinery and petrochemical construction. These construction programs will use the same international construction companies, technical skills and equipment as will be required for U.S. liquid synfuels construction. (Reference No. 43).

The purpose of this section is to discuss the range of potential constraints to the development of a viable liquid (and gas) synthetic fuels industry in the U.S.

This discussion of constraints is organized into the following categories:

Equipment	availability supply constraints performance constraints
Critical Materials	
Manpower	technical laborforce construction laborforce
Coal Supply	
Water Supply	

ulletEnvironment,	Health	and Safety
		standards and requirements permits and licenses

- Siting physical location infrastructure problems
- Transportation
- Technology Uncertainties
- Financial/Capital Availability
- Economics operating costs product costs

Chapter 3 has already covered the <u>technologies</u>, and Chapter 4, the <u>economics</u>. Capital availability has not been discussed here in this report. Additional assumptions on monetary policy and macro-economic policy over the next 20 years will be needed to consider this topic.j

#### 5.2.1 Equipment Problems

Seven different types of equipment which might **cause** supply constraints have been identified as follows:

#### Availability - supply Constraints

- 1. Demand for pumps in synfuels plants will be Pumps: very large. However, for small pumps, less than 1000 hp, there should be an adequate supply since producers could expand to three shift operations and European and Japanese manufacturing is available (Reference No. )= Large reciprocating pumps would 44 be in very short supply assuming that existing baseline demand persists. The synfuels industry could require between 50% and 100% of current world production capacity (Reference No. 44 )
- 2. Heat Exchangers: Demand is expected to exceed 25% of total domestic and foreign production capacity (Reference No.45). However, the industries' ability to increase capacity is reasonably good. The limiting factors would be availability of welders and of heat-treated metal plate from primary suppliers (Reference No. 44). Without firm orders, the heat exchanger manufacturers are reluctant to expand productive capacity.

- 3. Compressors and Turbines: Like heat exchangers, demand for compressors and turbines by synfuels plants could exceed 25% of existing production capacity (Reference No. 45 ). Traditionally, there is a two year lead time for these equipments. Manufacturers have expressed confidence that they can meet peak demand in 1984. (Reference No. 44) However, failure to order well in advance of need could cause delays and escalate costs.
- 4. Pressure Vessels and Reactors: Although synfuels demand will exceed 25% of productive capacity, suppliers are confident that they can meet demand (Reference No. 45 ). There is slack in the system due to slow economic growth and the absence of demand for nuclear reactor vessels (Reference No.43 ).
- 5. Alloy and Stainless Steel Valves: Demand for specialized valves will exceed 25% of current productive capacity (Reference No. <u>45</u>). Manufacturers' ability to expand productive capacity hinges on:

 adequate lead planning time availability of chromium, molybdenum and cobalt availability of quality castings and forgings

availability of qualified machinists (Reference No<u>. 44</u>)

- 6. Draglines: Draglines, which are essential for coal surface mining operations, have a lead time of 2-2-1/2 years. However, no production constraints are likely if firm orders are placed in advance of need.
- 7. Air Separation (Oxygen) Equipment: Reference No. 46 identified air separation plant fabrication capacity as the "most severe single constraint. " The critical components identified were aluminum distillation towers which are currently shop fabricated and brazed aluminum heat exchangers used in these towers. Techniques for field fabrication (to maintain quality control) have not been perfected. Development of acceptable field fabrication could reduce this potential constraint. Added reliance on production in Western Europe and Japan could also help, assuming that transportation facilities were available.

8. Distillation Towers: A specially constructed facility.

The accompanying Exhibits 5.1 and 5.2 (Reference Nos. 44 ) summarize the equipment supply constraints for a 1 MMBD and a 3 MMBD scenario (2000);

<u>Performance Constraints</u>--the possible failure to perform to specifications at operating conditions.

Concerns with ability to meet specific performance standards have been expressed for five categories of equipment as follows:

- 1. Gasifiers
- 2. Extractors
- 3. Hydrotreaters
- 4. Oxygen compressors
- 5. Coal slurry heaters

The available operational data for these five categories of equipment are from useages in process environments which are significantly different from the coal conversions regimes in liquid synfuels facilities. Substantial development will be required to modify and/or scale up equipment currently in commercial use (Reference No. 47 ). Therefore, these five categories of equipment impose potential constraints to the synfuels industry which would result from equipment failure or substandard performance.

#### 5.2.2 Critical Materials

Materials critical to the synfuels program are cobalt, nickel, molybdenum and chromium. After two independent analyses, only chromium was identified as a potential constraint (Reference NO.44,46) . U.S. currently imports over 90 percent of its chromium use and will remain highly dependent on foreign supply. Demand for chromium by synfuels programs could reach 7% of total U.S. demand. Exhibits 5.1 and 5.2 depict this concern.

#### 5.2.3 Manpower

#### Technical Laborforce

Engineering design manhour requirements for construction of synfuels facilities are 1.5 to 3 times greater than those

## EXHIBIT 5.1 (Reference 44)

## POTENTIALLY CRITICAL MATERIALS AND EQUIPMENT REQUIREMENTS FOR COAL LIQUIDS PLANTS AND ASSOCIATED MINES

Category	Units	Peak Annual Requirements	Us. Production Capacity	Requirements Percent of Production
Chromium	tons	10,400	400,000 <sup>1</sup>	3
Valves, alloy and stainless steel	tons	5,900	70,000	8
Draglines	yd	2,200	2,500	88
Pumps and drivers (less than 1000 hp)	hp	830, 000	20,000,000	4
Centrifugal Compressors (less than 10,000 hp)	hp	1,990,000	11,000,000	18
Heat Ex- changers	ft²	36,800,000	50,000,000²	74
Pressure Vessels (1.5-4" Walls)	tons	82, 529	671, 000	12
Pressure Vessels (greater than 4" wall)	tons	30,785	240, 000	13

(3MMBPD Scenario)

 ${}^{1}{}_{\text{Current consumption}}$ 

 $^2\,{\rm Total}$  for surface condensers, shell and tube, and fin-type.

#### EXH B T 5.2 (Reference 44)

#### Table 1-8 SELECTED MATERIAL AND EQUIPMENT ITEMS REQUIRED TO MEET PROJECTED COAL LIQUIDS PLANTS (AND ASSOCIATED MINE) NEEDS

Category		* Peak	Annual Requ	uirement and	d Year	Production Cepacities			
Category		1 MN Scen			MBPD mario	US	Japan	Europe	Total
Stainless Steel	(tons)	81,733	(1986)	52,299	(1986)	1,954,000	1,988,000	N/A	3,942,000+
Aluminum	(tons)	2,705	(1985)	6,443	(1986)	4,800,000	N/A	N/A	8,735,000
Chromium	(tons)	4,364	(1985)	10,409	(1985)	400,000(1,2)	N/A	N/A	N/A
Nickel	(tons)	756	(1986)	1,805	(1986)	114,000	72,000(3)	35,000	221,000
Cast Iron	(tons)	23,195	(1986)	55,610	(1986)	18,200,000	979,000	10,648,000	27,827,000
Iron and Steel Forgings	(tons)	5,766	(1985)	14,323	(1986)	1,418,000(1)	867,000	2,681,000	4,964,000
Steel Plate > 1.5"	(tons)	26,647	(1986)	63,767	(1985)	1,900,000	14,000,000	1,540,000	17,440,000
Pipe, Alloy and Stainless Steel	(tons)	9,818	(1999)	24,545	(1999)	1,852,000(1)	3,000,000	2,176,000	7,028,000
Valves, Alloy & Stainless Stèel	(tons)	2,481	(1985)	6,892	(1984)	70,000	68,000	97,000	235,000
Reinforcing Bar	(tons)	23,503	(1985)	64,886	(1986)	6,187,000(1)	14,438,000	10,161,000	30,784,000
Draglines	(yd <sup>3</sup> )	810	(1987)	2,196	(1987)	2,500	N/A	N/A	N/A
Pumps & Drivers < 1,000 hp	(1,000 hp)	343	(1985)	830	(1984)	20,000	N/A	N/A	N/A
Centrifugal Compressors & Drivers > 10,000 hp	(1,000 hp)	194	(1985)	481	(1984)	15,000	N/A	N/A	N/A
Centrifugal Compressors & Drivers < 10,000 hp	(1,000 hp)	833	(1985)	1,989	(1984)	11,000	N/A	N/A	N/A
Heat Exchangers	$(1,000 \text{ ft}^2)$	15,260	(1986)	36,780	(1986)	50,000	82,000	N/A	132,000+
Non-Nuc Pressure Vessels 1.5-4" wall	(tons)	33,677	(1985)	82,529	(1984)	671,000	223,000	132,000	1,026,000
Non-Nuc Pressure Vessels > 4" wall	(tons)	12,314	(1995)	30,785	(1999)	240,000	254,000	99,000	604,000
Boilers	(MM lb/hr)	8	(1985)	19	(1985)	210	N/A	N/A	N/A

N/A = Not available

-

\* Peak refers to maximum aonua requirements

Current consumption
 US dependent upon foreign supplies

.

3) Includes other Asian countries

SOURCE: E. J. Bentz & Associates

Indirect synfuel proneeded for refinery construction. cesses are the most engineering intensive since they are, in effect, two separate systems, 'e.g., gasificatio<sup>®</sup> and synthesis. However, even the direct liquefaction process requires significant amounts of engineering design manpower (Reference No. 45). The need for chemical engineers would Under a scenario projecting be the area of greatest concern. 3 million B/D by the year **2000**, demand for chemical engineers increases significantly between now and 1985 (Reference No. 440. An additional 1300 chemical engineers representing a 35% increase in this specialty, i.e. , a 35% increase in the process engineering work force, as found in previous design and project work at present (in 1979: 3600 chemical engineers) in less than six years would be required for the synfuels program= Engineering schools can generate new inexperienced chemical engineers to meet this demand and qualified chemical engineers will remain a scarce and expensive commodity. Demand for will remain a scarce and expensive commodity. other engineering skills will also increase but at a more manageable rate. It should also be realized that potential growth in other sectors -- such as defense needs for engineering and construction skills -- may also place an added demand On skill availability.

#### Construction Laborforce

Skilled craftsmen such as welders, boilermakers, pipefitters and electricians are already in short supply. These shortages have been exacerbated over the last decade by increasing reluctance on the part of craftsmen to follow construction work and relocate. Since many of the synfuels development projects would be located in areas with existing overall manpower shortages and virtually no existing pool of skilled manpower, labor could become a significant constraint. Using the 3 million B/D scenario, this industry would require 73,000 construction employees in 1986, the peak year. This is approximately 2% of the entire construction employment force (Reference No. 44 ). More training programs and use of "nonjourneymen" or "helpers" to supplement the workforce could reduce potential shortages. Recruitment of women and minorities would help also. However, some of these steps might be opposed by labor unions. Labor unions are particularly concerned that open-shop (non-union) construction companies will gain a foothold in this program. The accompanying Exhibits 5.3, 5.4 and 5.5 (Reference No. 44), summarize the construction manpower requirements under the 1 MMBD and 3 MMBD scenarios.

#### 5.2.4 Coal Supply

Chapter 2 has discussed U.S. coal supplies. In brief, the U.S. coal industry currently has approximately 100 million tons of productive capacity which is not being used. In addition, the coal industry traditionally has

## EXHIBIT 5.3 (Reference 44)

#### TOTAL ENGINEERING MANPOWER REQUIREMENTS FOR COAL LIQUIDS PLANTS AND ASSOCIATED MINES

# 3 MMBPD SCENARIO (Persons)

Scenario	1984	1990	2000
All Engineering Disciplines			
Design and Construction	8,500	5,200	6,300
Operation and Maintenance		2,200	4,800
Total	8,500	7,400	11, 100
Chemical Engineering			
Design and Construction	1,300	740	920
Operation and Maintenance		1,050	2,250
Total	1,300	1,790	3,170

.

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## EXHIBIT 5.4 (Reference No. 44)

## PROJECTED PEAK CONSTRUCTION LABOR REQUIREMENTS

(Persons)

Craft	1 MMBPD Scenario (1987)	3 MMBPD Scenario (1986)
Pipefitters	7,170	16,920
Pipefitters-Welders	2,400	5,600
Electricians	3,020	7,190
Boilermakers	660	1,570
Boilermaker-welders	130	310
Iron Workers	1,760	4 , 2 5 0
Carpenters	2,700	6,400
Other	12,830	30,660
Total	30,670	72,900

## EXHIBIT 5.5 (Reference No. 44)

Craft	Current Union Craftsmen	Coal Liquids Peak Requir	
		<b>3</b> MMBPD Scenario	1 MMBPD Scenario
<u>Pipefitters</u> (including welders)			
East North Central and East South Central Regions	37,672	10,300	6,300
West North Central and Northern Mountain Regions	14,498	11,800	6,900
Boilermakers (including welders)			
East North Central and East South Central Regions	5,260	900	500
West North Central and Northern Mountain Regions	2,075	1, 100	600
<u>Electricians</u>			
East North Central and East South Central Regions	36,860	3,300	2,000
West North Central and Northern Mountain Regions	12,662	3,700	2,200

#### REGIONAL MANUAL LABOR FOR CONSTRUCTION AND MAINTENANCE FOR COAL LIQUIDS PUNTS AND ASSOCIATED MINES

<sup>1</sup>Source: Construction Labor Research Council

<sup>2</sup>Source: Obtained by computer run of **Bechtel** Corporation Energy Supply Planning Model, as described in reference 44.

surge capacity which is brought on line whenever the spot price of coal increases sharply. The lead time for opening up new mine capacity, both surface and deep, ranges between three and five years. Since the construction of major synfuels plants takes the same length of time, adequate new coal supply can be brought on line in a timely fashion. Finally, the U.S. coal resource is so large that it is very unlikely that there would be supply shortages over the next century. For all these reasons, coal supply poses no constraint to synfuels development.

## 5.2.5 Water Supply"

Chapter 2 has discussed water supply concerns. Also reference <u>31</u> discussed these in detail. In brief, while the U.S. has abundant water supplies in aggregate, there are certain specific geographic locations where water supply could become a constraint to development of a large synfuels program. This is particularly true in the semi-arid portions of the West where significant coal reserves are located.

". . sufficient water physically exists to support a significant-sized synfuel industry in the Upper Missouri and Upper Colorado River Basins, the primary western fuel resource areas." (Reference No. 33)

The problems with water supply in these areas are institutional and highly political and often emotion-laded. Thus far energy developers have been able to purchase water rights from farmers or Federal and State water impoundments. As long as a relatively full market exists for the transfer of water rights, energy developers can afford to bid away the required water supply. In addition, corporate planners will need to consider water supplies for the construction/operating laborforce, their families, and the communities which-will support them.

## 5.2.6 Environmental Health and Safety

### Standards and Requirements

The liquid synfuels technologies "appear to have no absolute environmental protection constraint that would universally limit or prohibit deployment." (Reference No.<u>33</u>) However, the direct liquefaction processes have some potential to expose workers or the public to toxic and carcinogenic materials. Such risks could be judged politically and socially unacceptable and could become a development constraint. The Prevention of Significant Deterioration program under the Clean Air Act could pose absolute limits to the number of plants able to locate in a specific geographic area since the allowable increments of ambient air quality could be fully utilized. In the case of oil shale where the resource base is concentrated in a specific area in and adjacent to Northwest Colorado, PSD limits are very likely to constrain the number of facilities permitted. These limits, still to be developed, have not yet been set. Ranges of capacity vary, however, on what is possible." In addition,

"Some yet-to-be-defined regulations, if promulgated in their stringent forms, appear capable of severely limiting a number of synfuel technologies. These regulations include air quality emission control measures for visibility, changes in the original prevention of significant deterioration (PSD) regulations, extension of PSD limiting increments to other pollutants, shortterm nitrogen oxide ambient standards, development of hazardous waste tests and regulations and special waste regulations, toxic product regulations, and occupational safety standards. " (Reference No. 33 )

A detailed assessment of the environmental, health, and socio-economic impacts is found in reference no. 31 .

#### Permits and Licenses

The permitting and licensing process is complicated and time consuming. However, it poses no direct constraint on the synthetic fuels deployment program. The process generates procedural delays and provides multiple access to various public interest groups opposed to specific projects, specific technologies, or specific sites. More importantly, the process can be used by local political jurisdictions to either force project relocation or extract concessions from the project developers. Permit considerations are specifically discussed in the project discussions to follow.

### 5.2.7 Siting

Siting constraints are discussed in detail by the author in reference 31. In brief, Physical availability of sites is not a constraint. However, optimal siting by industry using their objective function often conflicts with the goals of other interest groups. Since much of the synfuels development will occur in areas with low population density, "conflicts will arise between the rural social order which currently exists in the region and the new urbanized society which will accompany growth. Early planning is required to handle these impacts." (Reference No. 45 )

To overcome the "locate your plant anywhere but not here" syndrome, corporate planners will have to work closely with state and local officials as well as with numerous civic associations. This requires full consideration of the secondary effects of development on the infrastructure of the immediate and surrounding These by their very nature are site specific areas. analyses. What new roads, schools, services, homes and institutions will be required? How will these requirements be funded? Can the community be protected against the worst features of the "boom" scenario and from the downside risk of bust? What does happen if the project fails and is abandoned? These are reasonable questions which often do not have reasonable answers. References 31 and 32 have discussed these key problems°

#### 5.2.8 Transportation

Transportation constraints can be a key concern. They must be considered on a regional/site specific basis. Reference 18 has treated these concerns.

As discussed earlier in Chapter 4, transport costs can be a key part of delivered cost. As discussed later in this chapter, the availability of inexpensive bulk transportation is crucial to project development.

#### 5.2.9 Tradeoffs

Hence, energy supply deployment will be affected by many competing constraining factors. Any specific project consideration must provide for a best optimum solution. This is clearly seen in Exhibit 5.6 in the variation to which oil shale targets would be achieved subject to different goals (Reference No. 8 ).

We will now look at our development of alternate supply scenarios.

#### EXHIBIT 5-6

#### ALTERNATE SHALE OIL PRODUCTION TARGETS (reference 8)\*

			tion target. 400.000	
To position the industry for rapid development		1 "		
To maximize energy supplies		1′		
To minimize Federal promotion				
To maximize environmental information and protection				
To maximize the integrity of the social environment				
To achieve an efficient and cost-effective energy supply system				
Lowest <i>degree</i> of attainment Highe SOURCE Office of Technology Assessment.	stategree c	of attainme	nt	

#### -The Relative Degree to Which the Production Targets Would Attain the Objectives for Development

Shale oil product ion targets are affected by many technical, environment al, and socioeconomic factors. As described in reference 8, the OTA has assessed the variation of 1990 production targets with regard to many of these key factors.

#### 5.3 Development of Supply Deployment Scenarios and Comparisons With Other Estimates

### (A) Shale Oil

The oil shale industry\*is in an advanced stage of development compared to other synfuel processes such as direct coal liquids. Design and construction (not including permiting) for an oil shale facility is typically in the 3-5 year time frame. Permiting requirements vary with two years being a typical time period. Most proposed/being developed projects are located in the West in the Green River Formation in Colorado, Utah, and Wyoming (Piceance, Uinta, Green River, Fossil, Great Divide, Washakie, and Sand Wash Basins) . Eastern shale development using promising new technical advances, discussed in Chapter 3, are likely to come on later. As discussed in the opening section, constraints center about resolution of land lease issues (the federal government owns over 80% of oil shale lands), environmental and water availability issues, and availability of skilled labor, especially hard rock miners.

Table 5-1 lists the potential commercial scale projects, identifying their proposed location, process, estimated start up, and project scale (production). In addition, the Department of Energy is conducting above--ground and advanced retorting projects.\*\* At present, permiting has been obtained for: Colony (final EIS, and a conditional PSD for 50,000, BPD complex), Union (final EIS for a 10,000 BPD commercial demonstration module unit) , Occidental (conditional PSD), Superior (final EIS) , and Paraho (draft EIS) . Based on the above projects planned, as well as individual surveys, scenario build-up rates are shown in Table 5-2. Comparisons of these rates with other estimates are shown in Table 5-3. This information is current as of 12/80.<sup>P</sup>

Initial production of shale, expected in the West, is expected to be treated (upgraded/refined) in the Rocky Mountain region, and will utilize existing spare refinery capacity. The next anticipated sequential market area is the Midwestern refinery region utilizing current inplace pipeline capacity (to the extent that anticipated new crude finds in the Overthrust Belt will not absorb pipeline capacity). The key markets envisioned for shale oil is as refinery feedstocks producing a large middle distillate slate for anticipated growing middle distillate needs (such as diesel oil). Shale oil residuals have also been proposed for use in turbines (current tests being sponsored by EPRI at Long Island Lighting). Using a typical refinery product slate, estimated shale-derived products are depicted in Table 5-4.

Private communication, DOE 12/80.

<sup>\*</sup>I.e., the industrial interests (oil, chemical, as identified in table 5-1) that are comprising the newly created shale industry.

PROJECT	SITE	PROCESS	PROJECT SIZE (1 OOOB/D	EST START ) UP	APPROX. COST (B\$)
COLONY DEVELOPMENT (Exxon, Tosco) <u>STATUS</u> : \$75 million spent to-date; planning, detailed engineering design and cost- ing completed; construction suspended; Exxon recently bought 60% share with con- tingencies tied to 1985 start-up; Tosco may seek Federal loan guarantee to raise its share of capital	CO	Surf ace Retort	47	1985	1.7 (1980\$)
UNION OIL <u>STATUS</u> : All permits received to construct and operate 9000B\D experimental retort which will be done with pri- vate financing (and \$3 tax credit) ; 50,000B/D project depends on results of experi- mental retort.	CO	Surface Retort	50	1983 (9000B/D)	
TOSCO SAND WASH <u>STATUS:</u> \$2 million spent by end of 1978; planning ex- ploration, and environmental analysis; TOSCO could use technology developed for Colony project, but would have to raise capital for both projects.	UT	Surface Retort	47	1988	
RIO BLANCO (GULF, STANDARD OF INDIANA) <u>STATUS:</u> \$245 million spent to-date; shaft sinking & surface construction activit- ies; further action pending Federal incentive programs.	CO	Mod In Situ & Surface	76	1988	

## TABLE 5-1: POTENTIAL COMMERCIAL SCALE PROJECTS SHALE OIL

## TABLE 5-1 (Continued)

PROJECT	SITE	PROCESS	PROJECT SIZE (1000B/D)	EST START UP	APPROX. COST (B\$)
OCCIDENTAL-TENNECO <u>STATUS</u> : Site preparation & shaft sinking; detailed development plan.	CO	<b>OXY</b> Modified In-Situ	50	1986	
WHITE RIVER SHALE PROJECT (Phillips, Sun, Sohio) <u>STATUS</u> : Detailed development plan completed. Environmental monitoring continuing. \$86 million spent to-date. Title status cleared by Supreme Court decision.	UT	Surface Retort	50 to 100		
SUPERIOR OIL <u>STATUS</u> : Pilot studies com- pleted; environmental analy underway at BLM; feasibility studies underway; <u>pending land</u> <u>exchange appears to be con-</u> <u>trary to current DOI policy</u> .		Surface Retort	13 + minerals		
PARAHO DEVELOPMENT <u>STATUS:</u> Beginning feasi- bility study (DOE funded) .	UT	Surface Retort	30	1984	
GEOKINETICS <u>STATUS</u> : Beginning DOE funded feasibility study.	UT	Surface Retort	2 to 8	1985	
TRANSCO ENERGY <u>STATUS</u> : Beginning DOE funded feasibility study.	КY	IGT Hytort	50	1984	
CHEVRON <u>STATUS</u> : Recently announced initiation of feasibility study.	CO	Surface Retort	50		
SOURCE: E. J. Bentz & Associa	tes	5-19			

Scenario	1980	1985	1990	1995	2000
A Capacity added in period Total Capacity		. 5 . 5	7 . 5 8	. 5 9	0 9
B Capacity added in period		.5	9.5	8.5	.5
Total Capacity		.5	10	18.5	19

TABLE 5-2:	SHALE OIL	BUILD-UP	SCENARIOS*
	(in units OF CRUDE		

NOTE: Most shale plants are estimated to be sited in the Green River Formation (Colorado, Utah, Wyoming) .

\* Shale oil build-up scenarios were constructed using interviews and referenced literature as cited in table 5-1, text, and footnot p.

SOURCE: E. J. Bentz & Associates

т	ABLE 5	(t		s of bar	NT SCENAR rrels per		<u>1980-20</u> crude c	
Source	19	80	1985	1987	1990	1992	1995	2000
U.S. DOP (2/80)	E1 -	-	80	225	400	450	450	450
Scenario	A -	-	25		400		450	450
DR1 <sup>2</sup> (10/79)	-	-	185	350	700	800	925	950
National Energy P II (5/79)								900-1300
U.S. DOP (11/80)	2 <sup>5</sup> -	-	25	160	400-500	550-800		
Scenario	в -	-	25		500		925	950
OTA <sup>4</sup> (6/80)				400				
Shell				150				
* <u>NOTE</u> :					s are in and Wyor		t, in the	e Green River
	Interpr							
	•					n Actior	n Plan,	" Feb. 1980.
	h		rch Ins				_	
			<b>fuel</b> Cor 1, 11/80		n Plannin /80.	ng Task i	Force,	private
	<sup>4</sup> otaAr	Asses:	sment of	E Oil SI	hale Tech	nologies	, 6/80	
	<sup>5</sup> U.S. N Shell-	Jational -U.S. 1	Energy National	Outloo Energy	k 1980-19 Outlook,	990, Shel Feb. 19	l Oil 980.	Co., 2/80.
SOURCE:	E. J.	Bentz	& Assoc	iates				

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Scenario	Products	1980	1985	1990	1995	2000
2	Gasoline		4.25	68	77	77
A	Jet Fuel		5.0	80	90	90
	Diesel Fu	ıel -	13.5	216	243	243
	Residues		2.25	36	41	41
	Gasoline		4.25	85	157	162
В	Jet Fuel		5.0	100	185	190
	Diesel Fu	iel -	13.5	270	500	513
	Residues		2.25	45	83	86

TABLE <b>5-4:</b>	ESTIMATED	TYPICAL	SHALE	OIL	PRODUCT	SLATE:	*	1980-2000
	(thousand	s of bar	rels p	er d	<b>lay</b> of cru	ude oil	eg	[uivalent)

Table values derived using Table 5.2 values, and typical yield slates (Chevron Research, 1978 reference: "Refining and Upgrading of Synfuel From Coal and Oil Shale by Advanced Catalytic Processes") discussed in Chapter 4, Section 6.

Because of relatively higher hydrogen content and lower aromatic concentration (than in general to coal liquids) , a "natural" product slate from shale oil is a mixture of gasoline, diesel, and jet fuel.

SOURCE: E. J. Bentz & Associates

#### (B) Coal Gases

As shown in the accompanying project tables, there is a significant level of varied activity in the coal gases area. Key generic processes are low/medium Btu gas and pipeline quality H-Btu gas.<sup>4</sup>

#### Low/Medium Btu Gas

As discussed in Chapter 3 and in the Appendix, leading technologies include the regular and slagging Lurgi gasifier (especially in earlier years), Texaco, Westinghouse, Koppers, and Winkler gasifiers.

Since low/medium Btu gas offers industrial and utility users a relatively curtailment-free source of high quality fuel and chemical feedstock, it is expected that they will penetrate into the utility and chemical market. The Energy Security Act specifically exempts medium Btu gas from allocation and pricing regulations.

Low-Btu gas finds key market use as industrial fuels in such applications as kilns, small boilers, and chemical furnaces. At present it has been estimated that there are about 15-20 domestic facilities (Reference No. 48 that are beginning to use low Btu gas for these applications. These include chemical firms such as Dow Chemical as well as automotive giants such as General Motors.

The Glen Gery Corp. has itself four facilities gasifying coal to produce a fuel gas to fuel their brick kilns, while Caterpillar Tractor plant in York, Pennsylvania produces fuel gas for heat treating furnaces. NCA (8/80) estimates there are nine <u>commercial</u> plants (in operation, under construction, or in proposal/planning stage). It has been estimated (Reference No. 50 ) that low Btu gasifiers are feasible at approximately 3500 industrial plant sites. These plants are expected to be geographically located at coal/adjacent to available coal suppliers.

Medium Btu gas serves several markets. Among them are utilities and chemical feedstock markets. Medium-Btu gas could be used as a synthesis gas for producing chemical products (ammonia, fertilizers, plastics), as well as utility power. Similarly, steel industry uses fuel for blast furnaces and annealing operations.

A potential co-product, methanol, could also be used as a utility peak showing fuel in turbines, or as an automotive fuel (Reference No. 51 ). Medium-Btu gas can also be used in utility use in a combined cycle power generation mode. NCA (Reference No. <u>48</u>) estimates there are five commercial scale plants in the proposed/planning stage. Key demonstration plants at TVA, Memphis Industrial Fuel Use Plant, and Cool Water, California (Southern California Edison), are in advanced stages. It has been estimated (Reference No. <u>50</u>) that there are approximately 350 potential sites for single user or limited distribution medium Btu gasifiers. In addition, there are combined-cycle markets (Reference No. <u>51</u>). As shown on the accompanying tables (and NCA survey), likely locations for medium Btu facilities include Louisiana, Texas, Arkansas, Pennsylvania, New Mexico, California, Tennessee, Montana, Virginia, and Illinois. Table 5.5 lists the key proposed projects under way.

Table 5-7 gives the scenario deployments of medium Btu/L Btu gas. The rate build-up was estimated by review of the cited data tables, on-line surveys, and judgmental interpretation with alternate comparative estimates.

#### H-Btu Gas

As shown in the accompany table (Table 5-6), of proposed commercial scale projects most early H-Btu gas development will occur in the West, especially in the states of North Dakota, Wyoming, Utah, New Mexico, and Montana (Northern Great 'Plains Regions and Rocky Mountain Region). Construction is at present underway in North Dakota on the Great Plains Gasification project. As shown in the table, this plant could be producing by 1984, with a production of 138 mmscf/day, at which time a second plant would begin (an additional 138 mmscf/day). Later plants are expected to be deployed in the Southwest (Texas, Louisiana, Arkansas, Oklahoma), and in the East (Pennsylvania), and capture the use of existing transportation lines.

The predominant end use for H-Btu gas is space heating (industrial/commercial) . Industrial use of the gas will be in the chemical, utility, and steel, iron and glass products industries (i.e., large current users of natural gas) . Market penetration will be affected by the pricing treatment of gas (e.g., rolled-in pricing) over the estimation period (period of natural gas deregulation) . Table 5-7 gives the scenario deployments of H-Btu gas over the estimation period. It is based on judgmental interpretation of the plant-specific build-up data cited, and on-line survey results. Table 5-8 gives the comparison of the scenario estimates with those of other sources.

I. <u>THE FOLLOWING PROJECTS A</u> (1	2/80)	ENTLY UNDER	<u> CEVELOPMENT</u>	
PROJECT	SITE	PROCESS	PROJECT SIZE (1000BOE[D)	APPROX . COST (B\$)
REYNOLDS ALUMINUM CO.	VA			
APPLICATION: Power Generation for Aluminum Reduction				
can-/do*	PA			
APPLICATION: Industrial Gas				
MUNICIPAL UTILITIES BOARD	AL			
APPLICATION: Industrial Gas				
PANHANDLE EASTERN	TX		8	
APPLICATION: Industrial Gas				
MEMPHAS GAS*	TN			0.3
<u>APPLICATION</u> : Utility\Feedstock (construction begins in 1982)				
SAN DIEGO P & L	CA			
APPLICATION: Utility/Feedstock				
ILLINOIS POWER COMPANY	IL		2	0.1
APPLICATION: UtilityCombined Cycle (1982 target)				
SOUTHERN CALIFORNIA EDISON	CA		3	0.3
<u>APPLICATION:</u> UtilityCombined cycle				
HOUSTON NATURAL GAS	LA			
APPLICATION: Utility/Feedstock				
COOLWATER	CA		100MW	0.2
<u>APPLICATION:</u> Utility-Combined Cycle (1984 target)				

### TABLE 5-5 : POTENTIAL COMMERCIAL SCALE PROJECTS - LOW/MED BTU GAS

I. THE FOLLOWING PROJECTS ARE CURRENTLY UNDER DEVELOPMENT

\*These projectsarecurrently funded as part of the Fossil Energy Technology Demonstration Program. (12/80)

## TABLE 5-5: (I Continued)

PROJECT	SITE	PROCESS	PROJECT SIZE (10 00BOE/D)	APPROX. COST (B\$)
MID-WEST ENERGY COAL ALTERNATIVE, INC.	IL			
<u>APPLICATION</u> : Industrial Fuel/ Feedstock				
CARTER OIL A <u>PPLICATION</u> : Industrial Gas and Feedstock	TX			
ENERGY CONCEPTS <u>APPLICATION:</u> Electric Generation and/or Feedstock	ОН			

SOURCE: E. J. Bentz & Associates

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#### TABLE 5-5 (Continued)

#### II. THE FOLLOWING PROJECTS RECENTLY RECEIVED DOE FEASIBILITY GRANTS (PL-96-126)

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PROJECT	SITE	PROCESS	PROJECT SIZE (1000B/D)	EST START UP
UNION CARBIDE	TX	Texaco	21.550	1988
$\frac{\text{APPLICATION}}{\text{and feedstock}}$ : MBG+H <sub>2</sub> for fuel				
GENERAL REFRACTORIES	КY	Wellman-	1.034	1983
APPLICATION: LBG fuel to kiln for Pearlite Mfg.		Galusha		
CENTRAL MAINE	ME	Texaco	14.100	1987
APPLICATION: Combined cycle power- (new)				
FLORIDA POWER	FL	BGC-	7.458	1985
<u>APPLICATION:</u> Combined cycle repower		Lurgi		
TRANSCO	TX	Lurgi	21.550	1985
APPLICATION: MBG to existing power plants				
PHILADELPHIA GASWORKS	PA	TBD	3.448	1985
APPLICATION: MBG-fuel gas				
EG&G	MA	Texaco	28.500	1986
APPLICATION: Combined cycle power & methanol				

<u>NOTE:</u> Over 40 proposals were received in response to 3/79 Notice of Program Interest. About 50 proposals were received in response to Feasibility and Cooperative Agreement Solicitations under P.L. 96-126. pL 96-304 programs are not listed due-to the funding uncertainty associated with the current recission order.

PROJECT	SITE	PROCESS	PROJECT SIZE (1000BOE\D)	EST START UP	APPROX. COST (B\$)
GREAT PLAINS GASIFICATION <u>STATUS</u> : \$40 million spent for project design and en- vironmental work. All per- mits obtained but final FERC tariff to market the gas. DOE cooperative agreement & loan guarantee under P.L. 96- 126. Plant could be producing by <b>1984.</b> A second plant with additional 138 mmscf/day is contingent on the results of Phase 1.	ND	Lurgi	<b>25</b> (138mmscf/d)	<b>1984</b> earliest	1.5
WYCOAL GAS INC. <u>STATUS</u> : Recently received DOE cooperative agreement to develop definitive design, estimate costs, secure per- mits and approvals, obtain financing and identify long- lead delivery items; market is company owned pipeline to mid-West. Second phase would add a second 150 mmscf/d.	wy	Lurgi & Texaco	25 (150mmcf/d)		
EL PASO NATIONAL GAS <u>STATUS</u> : Initial 1972 appli- cation to FPC placed in abeyance. Coal commitment obtained; water lease ex- pected; FERC tariff required before construction.	NM	Lurgi	13 (72mmscf\d)	earliest 1986	.6
TEXAS EASTERN/TEXACO <u>STATUS</u> : Water and coal from Texaco's Lake Desmet Reservoir property. Recently announced privately financed feasibility study.	ΜΥ	Lurgi	50 (275mmscf/d)	could be operative by 1990	

## TABLE 5-6: POTENTIAL COMMERCIAL SCALE PROJECTS--HIGH BTU GAS

## TABLE 5-6 (continued)

PROJECT	SITE	PROCESS	PROJECT SIZE ( 1000 BOE/D)	EST START UP	APPROX . COST (B\$)
PANHANDLE EASTERN PIPELINE COMPANY <u>STATUS</u> : Coal and water commit- ments have been obtained. No filing yet before FERC. Second 135 mmscf/day stage if justi- fied by first stage results.	WY	Lurgi	25		2.
MOUNTAIN FUEL COMPANY <u>STATUS</u> : Feasibility study under way. No filing before FERC to date.	UT	Lurgi	50 (275mmscf/d)	1990	
NATURAL GAS PIPELINE CO. OF AMERICA <u>STATUS:</u> Preliminary engineer- ing design completed. No filing before FERC.	ND	Lurgi	50 (275mmscf/d)	late 1980s	
TEXAS EASTERN SYNFUELS <u>STATUS</u> : Beginning DOE funded feasibility study.	NM	Lurgi	43 (sng+MEOH)	late 1980s	
CROWE TRIBE OF INDIANS <u>STATUS</u> : Beginning DOE funded feasibility study.	MT	Lurgi	22	1987	

\* Refers only to PL 96-126 feasibility and cooperative agreements. PL 96-304 project programs are not listed due to funding uncertainty associated with the current budget recission order.

SOURCE: E. J. Bentz & Associates

SCE	SCENARIO		1985	5	1990	0	1995	20	2000
	Added Capacity in period (MMBD)	(# plants)	.025	(•5)	.155	3.1	.20 (4 )	.12	2.4
H-Btu Gas (A)	-		.025	.5)	.180	(3.6)	.380 (7.6)	.50	(10)
	Added Capacity	(# plants)	.025	.5	S-₽.	(7.5)	.35○ ≤7)	.250	(2)
H-Btu Gas		(50,000 BPD)							
(B)	Total Capacity	(# plants)	.025	(:2)	.400	(8)	.750 (15)	1.0	(20
Med/Low	Added Capacity	(E plants)	.06	(1.2)	.115	(2.3)	.125 (2.5)	.10	(2)
BCU GAS (A)	IN PETIOG (MMBD)	(50,000 BPD)							
	Total Capacity	(# plants	.06	(1.2)	.175	(3.5)	.30 (6	.40	(8)
Med/Low Btu Gas	Added Capacity in period (MMBD)	(# plants)	.06	(1.2)	.19	(3.8)	.15 (3)	.10	(2)
	Total Capacity	<u> </u>	.06	(1.2)	.25	(2)	.40 (8	.50	10)
Sum (A)	Total Capacity (MMBD)		.085	(1.7)	. 355	(1.1)	.680 (13.6)	06. (	(18)
SUM (B)	Total Capacity (MBD)		.085	(1.7)	.65	(13)	1.15 (23)	1.50	(30)
SOURCE:	E. J. Bentz & Asso literature cited i	z & Associates; scenarios cited in text, footnote	ו סי	onst at		sing ir .5 and	using interviews and 5.5 and 5.6.	1	referenced

H-Btu GAS AND MED/LOW Btu GAS SCENARIO DEPLOYMENT:

TABLE 5-7:

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TABLE	5-8	:	SYNTHETIC	COAL	GASES	COMPARISONS
-------	-----	---	-----------	------	-------	-------------

Source	1980	1985	1990	1992	1995	2000
National Energy Plan <sup>1</sup> (May 1979)						.8-1.0
Frost & Sullivan <sup>2</sup>			.8			2.2
Exxon <sup>3</sup>			.5			.7-1.5
u.s. doe <sup>4</sup>		.05	.36	. 6 3		
Shell <sup>5</sup>		.19	.49			
Scenario A		.085	.355		.680	.9
Scenario B		.085	.65		1.15 1	5

<sup>1</sup>U.S. National Energy Plan II.

<sup>2</sup>As reported in Synfuels, 2/80.

<sup>3</sup>Exxon Energy Outlook, 12/79.

<sup>4</sup>Private communication, DOE.

<sup>5</sup>Shell National Energy Outlook, 2/80.

SOURCE: E. J. Bentz & Associates

## (C) C<u>oal Liquids</u>

As discussed in earlier chapters, coal liquids consist of indirect liquefaction of coal (Fischer-Tropsch liquids, methanol, methanol-gasoline), and direct liquid processes (H-coal, EDS, SRCII) . As shown in the accompanying Table 5-9, all early (to 1990) commercial scale projects receiving current government support are in the indirect category, although several direct liquefaction proposals have been received. As such, indirect liquefaction liquids are expected to dominate coal liquids product in the later decades of the century. At present, the only commercially demonstrated coal liquefaction process is the Fischer-Tropsch process used in the SASOL plants in South Africa (described in Chapter 3) . This process technology, an indirect liquefaction technology, is being adopted and improved for use in the U.S. The other key indirect liquefaction processes are methanol production -- a well known commercial process technology, and Mobil-M methanol-to-gasoline process, which should be commercially demonstrated within several years. In addition to several U.S. funded domestic studies for Mgasoline (see Table 5-9), there is a pilot plant demonstration project in Germany (Reference-No. 48 ), and a natural gas-methanol-M-gasoline commercial project scheduled for operation -in New Zealand by mid-80's (Reference No. 49). At present, there are no "commercially available direct liquefaction processes. The government has jointly (with industry) funded an SRC 11 demonstration plant and an EDS, and H-coal pilot plants for operation in mid-80's. Including the government sponsored study projects, there have been a total of 13 commercial plants, 4 demonstration plants, and 4 pilot plants are proposed/or in operation in the U.S. (Reference No. 18).

The anticipated deployment, based on judgmental interpretation of individual planned projects, current survey work, and individual project reviews, is depicted in the accompanying Table 5-9. As expected, indirect liquefaction processes dominate throughout, with direct liquefaction processes coming on stream late in the century. Early deployment is expected in the Northern Great Plains and Southwest region to capture existing product pipeline capacity (and water transport) and to fill energy product demands. Direct liquefaction developments are projected to come on in the 90's, and focus their activities in the Appalachian and Interior coal regions.

Direct liquid conversions naturally produce a high fraction of heavy oils. Since the traditional market for heavy oils (utility and industrial boilers) will probably convert to direct combustion of coal and medium Btu gas, upgrading of product slates into other market fuels is probable. Gulf's "Phase Zero Study" to DOE (also see Market Applications for SRC-11 products, Proceedings of the Sixth Annual International Conference on Coal Gasification, Liquefaction and Conversion of Electricity, Univ. of Pittsburgh, July 31-August 2, 1979) identified a substantial market where coal-derived liquid boiler fuels would have a distinct economic advantage over coal combustion with flue gas desulfurization primarily in congested areas of the Northeast where retrofitting to include flue gas desulfurization is expensive. As an example, projected EDS product slate usage could consist of stationary turbine fuels, special marine diesel fuels, and potentially home heating oils.

In general, direct coal liquefaction yields a high fraction of heavy fuel oil products. Current R&D work (at the laboratory stage) aims at upgrading this yield to the middle distillate, and naptha portion, thus minimizing the residual portion. However, this requires considerable upgrading by hydrogeneration or hydrotreating, as discussed in Chapter 4. In general, the products will be much more aromatic than equivalent petroleun-based products (private communication, Exxon Company, USA, 10/80).

Indirect liquids such as Mobil-M gasoline and methanol have projected use in transportation, and transportation/utility peak usage respectively. These and other product slates (Fischer-Tropsch) have been identified and discussed in Chapters 3 and 4.

Tables 5-10 and 5-11 depict the scenarios constructed from this data. Table 5-12 compares the scenario with other data.

	Bentz	& Associate	s; note feasib		
refers only to PROJECT	SITE	PROCESS	PROJECT SIZE (1000BOE/D)	EST START UP	APPRQX. COST (B\$)
W.R. GRACE <u>STATUS</u> : DOE cost shared demo; conceptual design near com- pletion; construction schedul- ed for 1984.	TN	Texaco Methanol M-Gas	6		0.5
TEXAS EASTERN SYNFUELS <u>STATUS</u> : Feasibility study completed; entered into cooperative agreement with DOE.	КY	Fischer Tropsch	56		
HAMPSHIRE ENERGY <u>STATUS</u> : Beginning DOE funded feasibility study.	WΥ	Methanol M-Gas	18	1985	
NAKOTA CO. <u>STATUS</u> : Beginning DOE funded feasibility study.	ND	Methanol	40	1987	
W.R. GRACE <u>STATUS</u> : Beginning DOE funded feasibility study.	CO	Methanol	14	1986	
AMAX <u>STATUS:</u> Beginning DOE funded feasibility study.	MN	Methanol		1985	
HOUSTON NATURAL GAS/TEXACO <u>STATUS</u> : Beginning DOE funded feasibility study.	LA	Methanol	11	1987	
COOK INLET REGION <u>STATUS:</u> Beginning DOE funded feasibility study.	АК	Methanol	23	1987	
CELANESE <u>STATUS</u> : Beginning DOE funded feasibility study.	TX	Methanol	10	1986	
CLARK OIL & REFINING <u>STATUS</u> : Beginning DOE funded feasibility study.	IL	Methanol M-Gas	12	1987	

		of	Crude (	Dil Equiv	valent		
SC	ENARIO	1980	1985	1987	1990	1995	2000
A	Capacity added in period			3	3	3	5
	Total Capacity			3	б	9	14
В	Capacity added in period			3	5	10	12
	Total Capacity			3	8	18	30
A	Capacity added in period				2	2	2
	Total Capacity				2	4	6
В	Capacity added in period				2	8	10
	Total Capacity				2	10	20

 TABLE 5-lo:
 COAL LIQUIDS BUILD-UP RATE SCENARIOS :
 INDIRECT AND DIRECT\*

(12/80 ) (In Plant Units of 50,000 BPD)

\*Coal liquids build-up scenarios were constructed using interviews and referenced information as cited in Table 5-9, text, and footnotes p and r.

SOURCE: E. J. Bentz & Associates

I N D I R E C T

D I R E C T

	, , , , , , , , , , , , , , , , , , ,	-	e Oil Ed	quivalent				
	Scenario	1980	1985	1987	1990	1995	2000	
A	Capacity added in period			3	5	5	7	
	Total Capacity			3	8	13	20	
В	Capacity added in period			3	7	18	22	
	Total Capacity			3	10	28	50	

#### TABLE 5-11: COAL LIQUIDS BUILD-UP RATE SCENARIOS\* (12/80)

(In plant Units of 50,000BPD)

\*Values derived from Table 5-10.

Source	1980	1985	1987	1990	1992	1995	2000
National Energy Plan <sup>1</sup>							.7-1.8
Frost & Sullivan*				1.0-1.5			9.5
U.S. DOE <sup>3</sup>			.14 .12	.5 .37	.8 .57		
Shell <sup>4</sup>		.03		.25			
Scenario A			•15	.4		.65	1.0
Scenario B			.15	.5		1.4	2.5

## TABLE 5-12: COAL LIQUIDS COMPARISONS

(MMBD) of Crude Oil Equivalent

<sup>1</sup>National Energy Plan II, 5/79.

<sup>2</sup>Synfuel Week reported 2/8/80.

<sup>3</sup>Private communication, DOE, 11/80.

 $^4$ Shell National Energy Outlook, Preliminary Version, Feb. 1980.

#### (D) Summary Tables and Comparisons

Table 5-13 depicts the summed synthetic fuel deployment schedules. Table 5-14 compares our "grass root" scenario build-up with other estimates developed by different approaches. As seen in Figure 5-1, the scenario brackets most estimates.<sup>\*</sup>

Next we will look at the labor requirements associated with the scenarios, as well as identify other impacts and concerns associated with their synfuel deployment.

	(in plant uni of Crud	ts of 50 e Oil Equi			
	1980	1985	1990	1995	2000
Shale Oil		. 5	8	9	9
Coal Liquids			8	13	20
A Coal Gases		1.7	7.11	13.6	18
Total		2.2	23.11	35.6	47
(MMBD)		( .11)	(1.16)	(1.78)	(2.35)
Shale Oil		. 5	10	18.5	19.0
Coal Liquids			10	28	50
B Coal Gases		1.7	13.0	23	30
Total		2.2	33.0	69.5	99
(MMBD)		(.11)	(1.65)	(3.48)	(4.95)

TABLE 5-13 : SUMMED SYNTHETIC FUEL DEPLOYMENT SCHEDULES\* (in plant units of 50,000 BPD)

\*Derived from adding Tables 5-2, S-7, and 5-11.

	of c	rude Oil Eq	uivalent			
Source	1985	1987	1990	1992	1995	2000
Energy Security Act <sup>1</sup>		. 5		2.0		
Exxon Outlook <sup>2</sup>			1.2-1.5			4.0-6.1
Bankers Trust <sup>3</sup>			.5			
Mellon Institute <sup>4</sup>						2.1
Natl. Energy <sup>5</sup> Plan (II)						2.4-4.1
NTPSC <sup>6</sup>						
(Low-Meal)	002		0318		. 2 8 - 1 . 2 7	1 . 3 4 - 5 . 3 4
Shell <sup>7</sup> 2/80	.22		. 8 9			
Scenario A	.11					
Scenario B	.11					

# TABLE 5-14 COMPARISON OF TOTAL SYNTHETIC FUEL PRODUCTION ESTIMATES (TARGET GOALS)

(MMBD)

<sup>1</sup>Energy Security Act, PL 96-294 6/30/80, Sec. 100(a) (2) .

<sup>2</sup>Exxon Energy Outlook, Dec. **1979**.

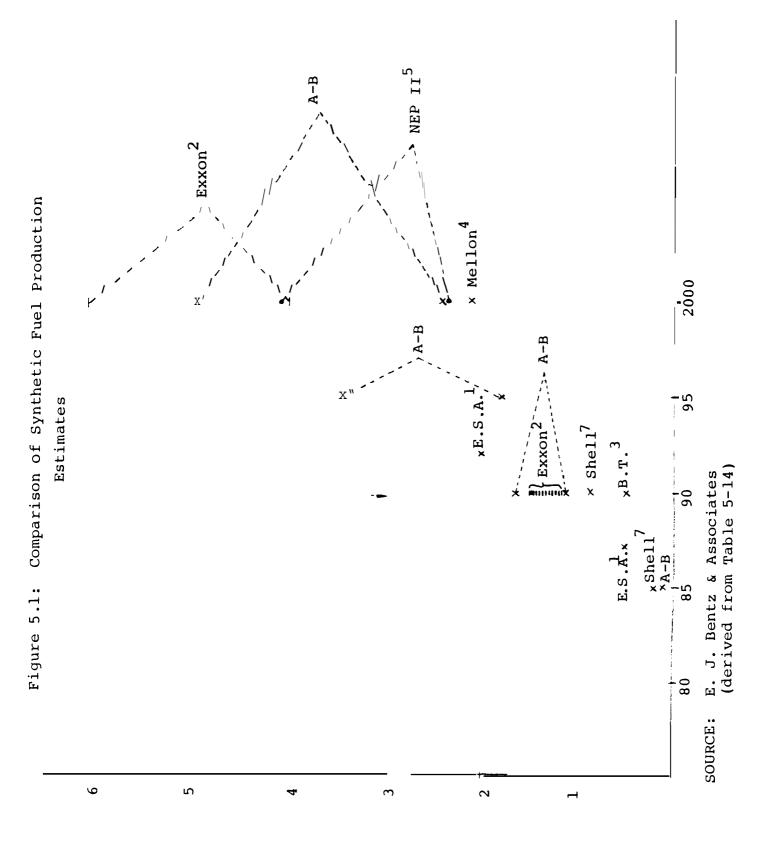
<sup>3</sup>Bankers Trust Forecast--as reported in <u>Synfuels</u>, 8/15/80.

<sup>4</sup>Mellon Institute Forecast -- as reported in <u>Synfuels</u>, 8/22/80.

<sup>5</sup>National Energy Plan II, May 1979.

<sup>6</sup>National Transportation **Policy** Study Commission Report, July 1979.

<sup>7</sup>Shell National Energy Outlook, preliminary version, Feb. **19**, 1980.



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#### 5.4 Labor Requirements Associated With The Scenarios

There are two categories of labor needs: construction labor and operations labor. As discussed in Chapters 2, 3 and 4, construction labor represents a peak employment situation whereas operations represents a steady-state labor requirement associated with the useful life of the facility. In addition, as discussed earlier, there are additional labor requirements in the geographical (and sectoral) area associated with provision of goods and services for the facility or for its labor force. The peak labor force is confined to a limited number of years (4-6) and often is several times the size of the resident population. This is especially so in the West. The impacts of this surge in peak labor can cause numerous community and environmental concerns in addition to severe strain on local infrastructure and even erosion of this infrastructure. Reference No. 52 discusses in detail some of these site impacts and their consequences. In addition, several studies, such as the Sec. 153a Studies of the 1976 Highway Bill, have looked at "Coal Roads" Issues, and the recently passed Energy Security Act mandates further studies to assess and hopefully suggest mitigation to energy impacted communities. The National Transportation Policy Study Commission in its final report (July 1979) specifically addressed the large and growing impacts of coal movement either in unbeneficiated or product form (pp. 141-149: The Commission forecast a large growth in the movement of coal. Associated with these movements will be: physical capacity concerns of a carrier nature; adequacy of service issues associated with carrier capabilities; and potential disruptions associated with these large scale movements) .

#### 5.4.1 Operations Labor Needs

Based on Chapter 4 results, a typical labor composition for operation of a 50,000 barrel/daw synthetic fuel facility is as follows:

Operations	120	people
Operator supervisors	25	people
Maintenance labor		people
Maintenance supervisors	30	people
Administrative	30	people
Total	355	people

Hence, upon applying this typical labor force participation to the scenario deployment estimates we arrive at the following aggregate estimate of needs: (See Table 5-15).

Workers		1985		1990		1995	4	2000
	A	В	A	В	A	В	A	В
Operators	264	264	2773	3960	4272	8340	5640	11,880
Operator Supervisors	55	55	578	825	890	1738	1175	2,475
Maintenance Labor	330	330	3465	4950	5340	10,425	7050	14,850
Maintenance Superviso:	rs 66	66	693	990	1068	2085	1410	2,970
Administrative	66	66	693	990	1068	2085	1410	2,970
Totals	781	781	8202	11,715	12,638	24,673	16,685	35,145

### TABLE 5-15 AGGREGATE OPERATIONS LABOR NEEDS (WORKERS)

Table 5-15 entries derived upon applying Chapter 4 typical labor force estimate to values developed in Table 5-13. Operations labor needs skill mix utilized, Chapter 4, based on ESCOE process estimates.

#### 5.4.2 Construction Labor Needs

For a typical 50,000 BOED synthetic fuel plant, the following construction labor skill mix needs are representative for the generic process as described in Chapters 3 and 4 (manpower requirements provided by Chapter 4 reference information and Reference No. 53 ). These estimates, as described in the reference citation, are associated with the conversion process above. In addition to these estimates will have to be added labor needs associated with mining, transportation, potential upgrading, distribution, and retailing. These requirements, however, will depend upon the specific product produced, the particular resource (fuel or coal) selected, the nature of the site, and other specific features. The Appendix to Chapter 2 gives a representative sample for different specific conditions. Manpower rates used are those based upon the previously referenced ESCOE work, which was part of the original study design.

#### A. Direct Coal Liquids and Shale:

Engineers	958	man	years
Draftsmen/Designers			years
Manual, blue collar	9160	man	years
(including pipefitters,			
welders, skilled labor)			

B. <u>Indirect Coal Liquids</u>:

Engineers	1985 man years
Draftsmen/Designers	<b>1330</b> man years
Manual, blue collar	16,185 man years
(including pipefitters,	
welders, skilled labor)	

#### C. <u>Coal Gases</u>

Engineers			years
Draftsmen/Designers			years
Manual, blue collar	9000	man	years
(including pipefitters,			
welders, skilled labor)			

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The typical construction period is spread such that the spread used for construction personnel labor demand is as follows: (This does not include permitting requirements or delays)

% Deployment	1	ч 2	lear 3	4	5
Engineers	30		15		5
Draftsmen/Designers Manual/Blue Collar	30 0%	40 10%	15 30%	10 40%	5 20%

Using the above estimates, and the previously derived supply deployment scenarios, we estimate the following incremental labor construction requirements (for each indicated time period) for each generic process and scenario (Tables 5-16 to 5-19 ).

# TABLE 5-16 : DIRECT COAL LIQUIDS\*

	1	987	1	990	1	.995	2	000
Scenario:	A	В	A	В	А	В	A	В
Engineers	-	-	1,916	1,916	1,916	7,664	1,916	9,580
Draftsmen/ Designers	-	-	1,250	1,250	1,250	5,000	1,250	6,250
Manual, Blue Collar	-	-	18,320	18,320	18,320	73,280	18,320	91,600

Incremental Construction Labor Requirements for Plants Coming On-line in Period Ending

(Man-years

\* Table 5-16 values based on process construction labor needs identified in Section 5.4.2 applied to values in Table 5-10.

	Г	1987	1	0661	7	1995	2	2000
Scenario:	A	В	A	B	A	В	A	m
Engineers	5,955	5,9=5	5,955	9,925	5,955	19,850	9,925	23,820
Draftsmen/ Designers	3,990	3,99°	3,99°	6,650	3,990	3,990 13,300	6,650	15,960
Manual, Blue Collar	48,555	48,555	48,555	80,925	48,555	48,555 161,850	8° 925 194,22¤	<b>194,</b> 22¤

TABLE 5-17 : **±NDT**RECT COAL LIQUIDS

\*

Table 5-17 values based on process construction labor needs identified in Section 5.4.2 applied to values in Tab $^{1}$ e 5-10.

SOURCE: E. J. Bentz & Associates

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# TABLE 5-18: SHALE OIL \*

Man-Years

Incremental Construction Labor Requirements for Plants Coming On-line in Period Ending

	1	985	1	.990	1	.995	2	000
Scenario:	A	В	A	В	A	В	A	В
Engineers	480	480	7,185	9,100	958	8,143	-	480
Draftsmen/ Designers	313	313	1 4,688	5,938	625	5,313	-	313
Manual, Blue Collar	4,580	4,580	68 <b>,</b> 700	87,020	9,160	77,860	-	4,580

\* Table 5-18 values based on process construction labor needs identified in

Section 5.4.2 applied to values in Table 5-2.

SOURCE: E. J. Bentz & Associates

#### TABLE 5-19 : COAL GASES \*

Incremental Construction Labor Requirements for Plants CominOn-line in Period Ending (Man-Years)

	1	985	1	990	1	995	20	000
Scenario:	А	В	A	В	A	В	A	В
Engineer	1,700	1,700	5,410	11,330	6,480	10,000	4,400	7,000
Draftsmen/ Designers	1,190	1,190	3,78+7777	7,910	4,543	7,000	3,080	4,900
Manual, BlueCollar	15,300	15,300	48,690	101,700	58,410	90,000	39,600	63,000

\*Table 5-19 values based on process construction labor needs identified in Section 5.4.2 applied to values in Table 5-7.

#### 5.4.3 Regional Deployment of Synthetic Fuel Plants Work Force

Not all construction labor needs will be uniformly distributed. As discussed earlier, different generic processes will favor siting in different regions:

- (i) <u>Oil Shale</u>: Almost all shale plants will be sited in the West until the close of the century. Hence all labor needs--both construction labor and operation labor--will be centered at the sites specified earlier (Piceance, Uinta Basin).
- (ii) <u>Coal Liquids</u>: Coal liquids, like coal gases, will be more widely dispersed due to the abundant and regionally varied U.S. coal supplies. As discussed earlier, most of the earlier plants will be of the indirect variety. Later direct plants will be deployed in the Interior and Appalachian regions. Using our previous build-up estimates, and those of other references, (34), we estimate the following regional work force for coal liquids:

Table5-20:Regional Share of Incremental Construction<br/>Work Force for Plants Coming On-Line in<br/>Period Endina:

	1	990	19	95	20	00
Scenario:	A	В	A	В	A	В
South Atlantic	0	11%	<b>29</b> %	15%	0	6%
East North Central	37%	30%	29%	35%	50%	41%
East South Central	13%	13%	14%	13%	17%	12%
West North Central	25%	21%	14%	15%	0	18%
West South Central	0	0	0	5%	17%	12%
North Mountain	25%	25%	14%	5%	17%	11%

(% Share of Totals in Man Years)<sup>t</sup>

(iii) <u>Coal Gases</u>: As discussed earlier, coal gases' characteristic size units are smaller, more numerous and more regionally dispersed. It is expected that they will share the same regional share deployment as do coal liquids reflecting sitings at coal resources, and reflected in the table above.

#### 5.5 Additional Concerns and Impacts: Product Acceptability Concerns

We have already identified key impact concerns and constraints: associated with synfuel development along the entire fuel cycle (Chapter 2); associated with individual technological processes (Chapter 3); with upgrading (Chapter 4); and with actual proposed synfuel plants (Chapter 50. We also have identified and discussed the supply-oriented needs and constraints associated with synfuel development. Many of these concerns are characteristic of the site-process selection (see Footnote to Chapter 5), and others are characteristic of the entire industry build-up to meet synfuel objectives.

In addition to these concerns, there are other concerns associated with synfuel product acceptability in the user marketplace. Traditional end use technology-such as internal combustion engines--have been optimized to meet performance specifications based on power fuel specifications, i.e., fuel product specificity must match engine tolerances on a physical and material basis. In addition to these performance specifications, additional institutional requirements have been placed on the utilization of end use technologies. Choosing the automobile again, automotive emission standards for criteria pollutants have been established with scheduled decreases in emissions over time. In addition, automotive fleets are subject to meeting the CAFE standards for fuel economy. Hence the optimization process of matching automotive performance with fuel specifications is a constrained one.

The potential changes in automotive standards (emission standards for diesel exhaust), as well as the potential introduction of new regulations and procedures which impact on fuel production (such as regulations pursuant to TSCA, RCRA, and Hazardous Waste Act), will further constrain the choices available and the time available to find them. Also in the achievement of these choices, tradeoffs between preservation of performance goals and removal of potential contaminants may have to be made. Several examples of the types and nature of these product acceptability concerns follow:

• Severe hydrotreating of syncrudes may alter or destroy certain fuel characteristics such as lubricity. In recent tests (Reference No. 54 of hydrotreated Alaskan crude, the Navy found that the hydrotreating affected the lubricity of the resulting fuel, which in turn affected the operation of their fuel pumps in aircraft engines.

- •The handling and burning of heavy fuel oils, especially from coal, may raise <u>potential</u> concerns due to their high aromaticity and potential toxicity. Potential carcinogenic concerns have also been raised (Reference No. 55). These concerns require further testing.
- •Nitrogen removal: Several concerns have been raised about the relatively higher concentration of nitrogen in synfuels. Among them:
  - Higher nitrogen content in synfuels has been found by Navy to be a factor in "gumming" (reference above).
  - . Meeting present  $NO_x$  automotive emission standards (1.2 grams/mile) has been difficult for the industry. With the higher fuel-bound nitrogen content of oil shale liquids, this difficulty is expected to increase. Although severe hydrotreating of the oil shale would certainly improve this situation, it would involve, as discussed in Chapter 4, additional upgrading costs. (In general, shale oil would be hydrotreated to reduce nitrogen content prior to pipelining to refinery. Also arsenic contaminants would be removed as-they would poison refinery catalysts, a key question in the degree of upgrading to meet anticipated specs, and at what cost?)
  - . Most SRC liquids have been found to be too high in sulfur and nitrogen content. Recent tests sponsored by EPRI at Con-Ed in New York with SRC-II liquids have required combustion modifications.
- The storage of incomplete refined or upgrated products may pose disposal problems (and costs) especially in more fragile ecosystems (see reference 56 for discussion of aggregate waste requirements).

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#### Next Step(s)

Next Step(s) will require that additional research and testing be performed both at the fuel supplier and end-user levels so that optimum changes can be made between fuel upgrading requirements and end-use combustion changes. As shown in Chapter 4, synfuels can, in principle, be processed to resemble current fuel production "specs" (e.g., gasoline produced from the Sun Oil refinery at Toledo from tar sands feedstocks) . Similarly, redesign of end-use technologies to meet less expensively produced synfuel yields are potential research options.

The potential use of the higher aromatic content of coal liquids for efficiency improvements in highercompression engines is one example. The use of neat methanol is another. The essential series of suboptimization "match-ups" --constrained by health, environmental, safety and other concerns such as liability for technology warantees--will also reflect the utilization of current infrastructure (e.g., refinery capacity), and the projected composition of natural crude supplies (Alaskan and Saudi sour crudes, Venezuelan and Bakersfield heavy crudes; Overthrust production), to which synthetic fuels contribute. This, however, is beyond the scope of this study."

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#### 5.6 FOOTNOTES TO CHAPTER 5

- a. (i) The general methodology used in developing the "bottoms-up" assessments has used the following sources of information:
  - (1) referenced literature and data cited in text and footnotes
  - (2) numerous interviews with industrial and governmental sources, including members of the OTA Synfuel Advisory Group
  - (3) proprietary information heretofore developed by EJB&A, as cited

\*Much of the interview information built upon existing and on-going studies being performed by EJB&A. As such, the data base used was much larger than the study scope allowed in itself. Among the key sources of interview information were:

- (1) <u>Governmental</u> interviews were conducted with numerous federal- and state offices including: the U.S. Department of Energy [Policy Office, Fossil Fuel" Office, Resource Applications, Conservation Office, National Laboratories (Oak Ridge)], the U.S. Environmental Protection Agency (Toxics Substance Office, R&D Office), Kentucky Department of Energy; California Energy Commission; and the Massachusetts Energy Office.
- (2) Industrial interviews were conducted with numerous staff of the major oil companies; chemical companies; automotive companies; and utility companies.

The OTA Synfuels Advisory Group, as well as the OTA staff, were particularly helpful in their sound advice, judgment, and insights in developing information.

(ii) The overall guiding general assumptions used in the methodological approach were:

 There will be no major international conflict which would preclude supply of foreign raw materials and manufactured equipment. (2) There will be no dramatic increase in the consumption of energy related materials or equipment by other segments of industry which will impact on the synfuels fuels program.

(iii) The overall approach methodology is given in footnote p.

(iv) Specific assumptions associated with the development of each of the scenario assessments have been given in the text, and in footnotes: p (general and for shale oil); q (for coal gases); and r (for coal liquids. Furthermore, regionalization techniques are cited in footnote t.

(v) Scenario scope was chosen in consultation with OTA staff at initial and interim briefings, and as reflected in contract study scope.

- b. As discussed later in the individual scenario sections, "high" refers to a maximum deployment schedule, which pushes the limits of material and skill mix availability. However, it does not represent an emergency, supply interruption contingency scenario. Development in the high scenario is conducted by the private sector with fiscal and R&D incentives being provided by the government so as to minimize commercial risk, and to accelerate the pace of development. The "business-as-usual" deployment schedule represents a more historical growth characteristic of capital-intensive new growth industries, as discussed in Chapter 4. High capital demands, technical uncertainties, and other factors discussed in Sections 5.2 and 5.3 dictate a more cautious approach that minimizes financial exposure. The governmental role is mainly an R&D role, especially in high-risk, yet potentially high payoff beneficial technologies. Government fiscal incentives are very minimal as compared to the high scenario. High and low scenario choices were chosen in conjunction with guidance from the OTA staff in initial, and subsequent interim briefings.
- c. I.e., in the mid-term (1980-2000), we have attempted using existing information on scheduled supply projects to match supply concerns with demand needs. An aggregate approach reveals little as to the "make-up" of the fuel composition, although macro aggregate techniques can be valuable in long-term analysis, and in investigating macro-economic effects such as capital formation and monetary effects.
- d. Post 2000 fuel demand slate requirements are dictated more by an assessment of long-term economic market forces, and post 1980 mid-stream supply corrections that by 1980 "current" supply deployment constraints. This is especially

so since there is ample time (for the 2000+ period) to remedy longer-term constraints and because of the inherent uncertainties associated with projecting long-term supply projections. This will be more fully discussed later in footnote p, subsection (v).

- e. "Transition period" here simply refers to the time period 1980-2000 in which we are introducing new fuel supply sources to complement our existing sources. Post 2000 fuel supplies may consist of considerable numerous, nonrenewable, and renewable fuel sources contributions. As such, the 1980-2000 period reflects a period of decisionmaking and change to achieve alternate fuel goals.
- f. Examples of these are: fuel cell use in automobiles; electric vehicles; and extensive use of active device solar heating and cooling. For a more detailed description of potential automotive end use technology changes see Report of the National Transportation Policy Study Commission, June 1979, p. 93.
- g. As an example of an alternate integrated approach see <u>Forecasts of Freight System Demand and Related Research</u> <u>Needs</u>, National Academy of Sciences, June 1978; "Transportation Modeling and Freight Demand Trends, " p" 33, E. J. Bentz & Associates
- h. Already defined in (e) above.
- i. These alternative assessments, as referenced, reflect the use of a variety of different techniques. The specific techniques used differ greatly. Whereas some forecasts rely heavily on the use of macroeconomic models (e.g., DRI, Wharton, Chase), others use more industry-specific survey approaches. In the cited references for each alternative forecast, the specific methodology employed is identified. It should be clearly recognized that there are no "best and only" approaches, since different technique highlight different effects, e.g., an industry survey may give good insight on industry-specific technology changes, but give little insight on the impacts of how potential external changes in national interest rates may affect the industry.
- j. Capital formation concerns including availability and rate concerns are a key ingredient to synfuel project development. However, scope, budget, and time precluded a discussion of an analysis of these concerns. A general discussion of these concerns can be found in "Synthetic Fuels," Report by the Subcommittee on Synthetic Fuels of the Committee on the Budget, U.S. Senate, September 27, 1979, Chapter IV, p. 23, and Appendix I, p. 55.
- k. Exhibits 5.1 and 5.2 identify respectively the <u>potentially</u> critical material and equipment requirements for coal

liquid plants (and associated mines), and overall selected material and equipment items required.

They both represent a series of computer runs using the ESPM model described in reference 44. The key implications of these tables and reference 44 are:

- . for <u>most</u> equipment items, projected requirements represent a relatively small percentage of overall manufacturing capacity
- . in general, domestic manufacturers can expand production as demand develops
- . in addition to domestic capacity, there is foreign manufacturing capacity that can supplement U.S. domestic capacity
- . there are key items, as discussed above in the text (such as draglines), where there may be a <u>potential</u> constraint of a capacity or leadtime nature

Furthermore, as illustrated in Table 5.2, reference 44 assessed for two different deployment schedules, peak needs for equipment as a function of current production capacity. In this regard" "peak" was used to represent the maximum annual equipment requirements associated with the deployment schedules.- Once again, we see that "draglines" and "heat exchangers" are items of concern in that peak requirements are a significant fraction of existing domestic capacity. These peak concerns are further constrained in that some items such as draglines, air separation plants, and large pumps and reactor vessels require substantial supply leadtimes. Although foreign purchases may alleviate potential shortfalls, early programmatic planning can facilitate domestic manufacturing expansions. These plans would include not only equipment planning but planning concerning: transportation needs, capital formation, siting concerns, water needs, and technical personnel needs. These will be discussed later in text.

1. Overall employment statistics are of limited value in assessing potential labor constraints. The shortages which may occur will be for a particular technical or craft skill. For this reason, exhibits 5.3-5.4 are broken down by skill mix. Similarly, since project construction-- as described in later section--is location specific, an overall regional assessment is illustrated in exhibit 5.5. As reference 44 discusses, the key labor constraint concerns are:

- . the availability of chemical engineers may be a key limiting factor in the availability of engineering manpower
- . the most serious challenge in meeting engineering requirements will probably be in the early peak years, as Exhibit 5.3 shows for design and construction. This simply reflects the early intensive use of these skills in normal project deployment
- that the supply of civil, electrical, industrial, and mechanical engineers will probably not present as severe a concern as meeting chemical engineering requirements (Exhibit 5.3)
- of skilled construction labor needs, the critical needs are those of pipefitters, welders, boilermakers, and electricians (Exhibit 5.4). For some sparsely settled regions of the nation where there is a limited skilled labor force, this will mean bringing in considerable new labor (such as in the Alaskan pipeline). Exhibit 5.5 illustrates this regional pattern of potential skilled labor needs.
- Water supply and availability is of key concern to the m. siting of synfuel plants. As mentioned in the text (p. 5-13) and in Chapter 2, this is particularly true for arid regions of the West- Under the prevailing system of purchased water rights, most of the available surface water supply in these Western regions has already been allocated. As such, these rights will have to be acquired for prospective projects. It has been estimated in The Nation's Water Resources, the Second National Water Assessment, U.S. Water Resources Council, 'Washington, D.C., vol. A-2, April 1978, that the characteristic maximum water consumption in the most water-scarce areas likely to contain synfuel plants would be about 5% of current consumption. State Water Law in the West: Implications for Energy Development, Los Alamos Scientific Laboratory, January 1979, gives a comprehensive discussion of current water rights, and transfer in the West, especially as they affect potential energy site development.
- n. Ranges of shale oil capacity vary greatly depending on key assumptions. As an example, the OTA's "An Assessment of Oil Shale Technology, " June 1980, lists a 1990 production target of 400,000 barrels/day as being "consistent with achieving an efficient and cost-effective energy supply system" (p. 10) and an alternate 1990 production target of 200,000 barrels/day as a target "to maximize ultimate environmental information and production" (p. 11). Similarly, Exxon, in its 1980 Report to the Business Roundtable, lists a target of 8 million barrels/day by the year 2010 in the

Piceance and Uinta Basin. These ranges which depict the uncertainty of many key technical and socioeconomic variables are illustrated in Tables 5-3 and 5-4.

- O. As discussed earlier, the determination of site choice for different processes is affected by many factors. There are several critical factors that are common to the siting of any synthetic fuel facility. They have been discussed at length in the literature of both coal and oil shale facilities (Reference Nos. 31, 32 and 33 ). One such review (Reference No. 32 includes a detailed evaluation of seven representative facilities for various critical factors, which include both physical and institutional aspects. The situations for coal and oil shale conversion facilities. The critical factors considered are:
  - Capital availability
  - Industrial marketing decisions such as transportation availability
  - Resource depletion
  - Air pollution control
  - Water availability
  - Surface mine reclamation
  - Socioeconomic disruption
  - ownership of land and the management of federally owned lands

The main objective is to determine on a regional basis the potential for development of a synthetic fuels industry with minimal conflicts. Assessment of the ability to mitigate some of the environmental constraining impacts have been studied (above references).

Among the characteristics that have been identified and assessed are:

(1) <u>Air Quality Characteristics</u>: Special attention has been paid to constraints due to Prevention of Significant Deterioration and non-attainment areas.

(2) <u>Water Availability</u>: Institutional factors (e.g., competing uses, allocation policies, water rights) as well as physical factors (e.g., stream flows, quality of the water) have been identified. (3) <u>Socioeconomic Capacity</u>. The capability of communities to adjust successfully to the potential social disturbances associated with the construction and operation of large synthetic fuel facilities have been identified as the key factor in affecting public acceptance. This factor is particularly important for synthetic fuel facilities to be located in western states where the communities are small relative to the size of the facilities. Socioeconomic capacity is evaluated with respect to population size of the affected communities, their infrastructure level of services, and growth history.

(4) Ecological Sensitivity. This factor is evaluated with respect to susceptibility of natural ecosystems to disturbances associated with large scale industrial activity. Waste disposal operations and reclamation of mined lands and disposal sites of spent shales are considered important considerations.

(5) <u>Human Health</u>. There is an undetermined potential risk to both the health of occupational workers employed in the synfuel plants, and to the population surrounding the plants. As discussed in reference <u>31</u>, the risk factors are still largely undefined because knowledge is lacking about the kinds and quantities of toxic materials to be released from actual synfuel plants. (See

(6) Land ownership. This factor, and particularly the management of federally owned lands, is particularly important in the West. There, the federal government is a major land holder, and some critical lands are owned by Indians. Policies established under the Federal Land Policy and Management Act of 1976, as well as existing management practices are in conflict with extensive exploration and development of coal and oil shale resources and with the siting of synfuel facilities.

- P\* Table 5-2 shale oil build-up scenarios were constructed using the following iterative process. This same approach was used in the build-up scenarios of coal liquids and gases:
  - (1) Utilize General Methodological Assumptions stated in footnote (a) (i.e., not supply interruption concerns).
  - (2) Specific Approach:
    - (i) From Table 5-1 develop initial project schedules baseline reflecting "business-as-usual conditions. In developing baseline schedules utilize specific

project information; interviews with industry and government officials and comparisons with other individual and aggregate companions (referenced in Table 5-3)

- (ii) After developing initial baseline, iterate by reviewing against above referenced comparisons and additional interviews. Using a modified Delphi-type approval, develop a final baseline schedule.
- (iii) using final baseline schedule, repeat steps (i) and (ii) above, under new "upper limit" conditions. These conditions reflect a maximum conditions. possible rate-of-growth schedule consistent with pushing material, manpower, and siting concerns discussed in Sections 5.1 and 5.2. They mostly closely reflect an environment of significant governmental fiscal incentives to minimize market commercial risk and accelerate development, as reflected in the economic climate of the fall of They do, however, reflect utilization of 1980. private market forces, and not large-scale direct governmental intervention. For a more detailed discussion of governmental assistance see "Synthetic Fuels, Report of the Senate Budget Committee, September 1979, Chapters IV and V. As such, this "high" scenario does <u>not</u> reflect an emergency planning, oil supply disruption scenario. Such-a scenario, although very useful in its own right, was not in the directed scope of work, and would require significantly different methodological assumptions and techniques.
  - (iv) After developing a final "high" and "low" scenario, specific scenario characteristics, such as differences in rate of growth, peaking of scheduled outputs, and leveling "off" phenomena were compared to above referenced interviews and literature. A comparison of several of these alternate "scenarios," albeit using different, and mostly proprietary techniques, is given in Table 5-3.
    - (v) post 2000 deployment schedules are mostly "second-round" decisions which would be based on both results of first round (1980-2000) successes and failures, as well as an assessment of the market needs for synthetic fuels in light of the supply, availability, and price of conventional fuels, as well as end-uses. For these reasons, extreme values (at 2000) reflect first round decisions on deployment, and not second round decisions. As such, they are subject to more uncertainty. A

long-term overall energy supply, demand, and price forecast was outside of the scope of this effort. Also, for the numerous uncertainties in Sections 5.1 and 5.2, as well as the technical and methodological uncertainties inherent in longrange forecasting. A discussion of the methodological and data needs associated with long-range energy forecasting is given in Forecasts of Freight System Demand and Related Research Needs, National Academy of Sciences, 1978, p. 33; "Transportation Modeling and Freight Demand Trends," E. J. Bentz & Associates. A discussion of the supply and availability of energy for future transportation needs is given in Alternate Energy Sources, Part B, Academic Press, 1981, p. 733, Transportation and Energy, Outlook to 2000, E. J. Bentz & Associates.

- (vi) There are additional product quality and acceptability concerns associated with the use of the synfuel products. These concerns, already introduced in Chapter 4, are discussed in Section 5.4 and accompanying footnote. They add an additional element of uncertainty into the deployment schedule, but at this early research stage are at best difficult to bracket.
- From Tables 5-5 and 5-6 and referenced literature and q\* interviews, the low/reed Btu and high Btu coal gas build-up scenarios were constructed from Table 5-7, using the iterative methodology described in foot note p, and the general assumptions outlined in footnote a. As discussed in the text (Section B), particular reference 50 was made to the National Coal Association Coal Synfuel Survey reference as well as detailed proprietary information developed by E. J. Bentz & Associates, and numerous private communications with industry and governmental officials (federal and state). As stated on p. 5-24, the eventual regulatory treatment of high Btu gas (pricing, advances to 'pipelines) will greatly affect the scenario schedules. Although the scenarios assured that high Btu gas will be treated as natural gas, this realization will be affected not only by the treatment of high Btu gas, but also on the pricing schedule of natural gas itself (i.e., natural gas deregulation). Table 5-8 summarizes comparisons with current alternative forecasts. Note, as discussed in footnote p, these alternative forecasts employed a variety of different proprietary methodological techniques. As such "bottoms-up" comparisons are not appropriate.
- r. From Table 5-9, and identified literature and interviews, using the iterative methodology described in footnote p, and the general assumptions outlined in footnote a, Table 5-10 was constructed. Of specific assistance were references 50 and 51, as well as proprietary information developed by

E. J. Bentz & Associates, in the deployment schedules. In brief, indirect liquefaction technology is a known, commercially proven technology. Although on-going R&D will improve this technology (such as alternate gasifier designs), it is building upon a known baseline. Also much of the equipment needed is commercially available. As such, early development in the coal liquids area will utilize indirect liquefaction techniques (including the Mobil-M gasoline process) . Direct liquefaction offers great promise, but requires more R&D to achieve a similar commercial-type status. Also, as discussed in Chapter 4, many of the direct products will have to be upgraded, at additional costs, for use in existing enduse technology. Hence, "direct liquids" will be introduced later in our deployment schedules. Because of the variety and complexity of coal liquid sources, as well as the shale oil liquid contribution to our liquid supply (discussed earlier), additional iterations had to be undertaken sequencing individual supply sources (e.g., shale and indirect liquids earlier) and then reiterating the sums against independent numbered comparisons and previous interview results. As such, the "coal liquids" scenarios -- high and low -- represented the greatest number of iterations. The comparisons of the developed build-up rates with alternate estimates (derived using different proprietary methodologies) is given in Table 5-12.

- s. As discussed in footes a, 'p, q, and 5, Table 5-14 depicts alternative macro-estimates developed by the referenced sources using alternate (and often proprietary) techniques.
- t. Table 5-20 developed by distributing on regional basis each of the incremental construction work forces for each of the processes, described in Tables 5-16, 5-17, 5-18, 5-19 and then adding regional sums. In tiers, these regional factors were first obtained using following independent sources:
  - . reference 34 regional factors developed for coal liquids
  - Tables 5-1, 5-5, 5-6, and 5-9
  - . reference 20 for coal liquids (indirect) and reference 6 for all synthetics
  - . proprietary information developed by E. J. Bentz & Associates

NOTE: It should be noted that Figure 2-3 on p. 2-4 represents the geological coal resource region. Because such a breakdown does not include all supply resources (e.g., shale) as well as the fact that site location is dependent upon a variety of factors (see footnote o), the

regions chosen for regionalization were the well-known and used (in all the above references) census regions.

u. An example of the diversity of aromatic chemical properties associated with coal-derived gasoline is given in the following table.

Aroma- tics (Wt %)	Gasoline from Petroleum	Gasoline from SRC-II Naphtha Hydrotreated	Gasoline from EDS Naphtha Hydrocracked	Gasoline from H-Coal Gas-Oil Heavy Hydrocracking
Benzene	.12	18.0	.08	5.1
Toluene	21.8	19.0	12.6	6.5
Alkyl- benzene °8 <sup>-</sup> °1 3	7.0	27.9	43.6	14.6

SOURCE: U.S. Environmental Protection Agency, Research Triangle Park, 1980 APPENDICES

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# APPENDIX TO CHAPTER 2

Typical Mining Characteristics: Tables 3-8 From Reference: "Technology Characterizations" U.S. DOE, June **1980** 

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1     100004     mutatial     1.1.1 ± 100     1.0.00004       0.11     0.11 ± 100     0.100000000000     0.1000000000000       0.11     0.11 ± 100     0.1000000000000     0.11 ± 100       0.11     0.11 ± 100     0.11 ± 100     0.10000000000000       0.11     0.11 ± 100     0.11 ± 100     0.11 ± 100       0.11     0.11 ± 100     0.11 ± 100     0.110       0.11     0.11 ± 100     0.110     0.110       0.11     0.11 ± 100     0.110     0.110       0.11     0.110     0.110     0.110       0.11     0.110     0.110     0.110       11     0.00000000     0.110     0.110       11     0.000000000     0.110     0.110       11     0.000000000     0.110     0.110       11     0.000000000     0.110     0.110       11     0.0000000000     0.110     0.110       11     0.0000000000     0.110     0.110       11     0.00000000000     0.110     0.110       11     0.00000000000000     0.110       11     0.0000000000000000000000     0.110       11     0.00000000000000000000000000000000000	
America     0.1     1.0     1.0       America     0.1     0.1     0.1       America     0.1 <td></td>	
Quantity     etutement     1.4 ± 105       Quantity     ether costs     1.6 ± 105       1     ether costs     1.6 ± 105       1     restruction     1.1 ± 105	
PERSONNEL         Mortacia           1         Construction         Mortacia           1         construction         1.3           1         non-manual, technical         1.3           1         non-manual, technical         0.1	
I     constrated       II     constrated	
11 Inc. Inc. [active] and [active] 0.1 12 approximation (.) 13 approximation (.) 14 approximation (.) 15 approximation (.) 16 approximation (.)	
11 operation 12 ann-mail, technical 3.9 3 ann-mail, technical 3.9	
sometric and the second s	
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b selec Acalcium cerbunate and maintained are unacter than calcium o boas and ba and ba and ba and ba and base and and part of the treatmant process.	7 7 7 3
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reatory, and a start a transmission and Operating Coste for Goal Stafp Mines, 1976. An base man, Padaral Cost Minagenesi and Operating Coste for Goal Stafferent, 1979.	

BRACT STSTDI	RESONACES USED: (Per 19 0 IM Produced)		RESIDUALS AND PRODUCTS: (Per 10 <sup>12</sup> Bru Produced)	
5138 a 73,300 tong of raw shalo par day a 0.413 x 10 <sup>12</sup> Bcu/day	<u>PUEL</u> Ter unbined shale	Tone 178,450	AIR POLLUTANTS 9 articdoiso	1000 21.79
e 2,800 Btu/pound of raw shale			50,2	0.21
· 30 gallons/ton shale ofl content	DIENCY		#G	2.95
o mine operatos 320.5 deye/year	electricity for operating	MA	hyðrocarbona	3.44
• 24.2 x 10 <sup>6</sup> tons of shale minod/year • total annual output to 136.67 x 10 <sup>12</sup> Btu	drilling equipment and		CO	1.80
	trucke			
e mine life is 30 years	6 (5 / 6 / 7 / m)	• ()	NATER POLLUTANTS	
DESCR 177100	COMPOSITION	<u>X (by weight)</u> 17.1	probability of • 110 C	
o is sufface mining, the everburden is	organic Motorial Vatar	17.1	contamination of under-	
removed appealing the underlying shale.	inorganic materia)	ei.5	ground unter by also water	
Shale is mined using the bench technique.	•	•1.7	SOLID WASTE	
Shale is fractured through drilling	LAND <sup>(1)</sup>	• • •	negligible (.00 Precessing)	
and blasting and transported by trucks	nine development	<u>Acces</u> 0.8	maftifiate (100 stocestift)	
to primary crushing site.	disposal of permanent everbuiden	7.4	ENERGY PRODUCT	178.430
CONFORMENTS	storage of apant shale	1.1		,
e drilling equipment	disposal of spent shale	1.1		
· escevetion equipment (cranes)		1.1		
• cluchete	WATER	Acres-Fest		
e trucks	mining and crushing	Acres-Feet 2.8 (2.2 - 3.3)		
TTAL BOUNDITAL CONCERNS	<u>00575</u> (2)	Pollara (1978)		
• sir quality deterioration	construction (*)			
e moise	Babychet	226,033		
é vater regulrement	moteriolo	28,809		
e contamination of underground water	Qutpmoat	339,649		
supplies with saline mine water	• cher Coet	14,792		
	total	609,283		
	operation & maintenance	MA		
	PERSONNEL	Nurberg		
	construction	MA		

A-4

(1) This represents land committed to use over the lifetime of the plant, divided by the <u>annual</u> output of the plant, expressed in trillion Btu. (2) This represents total cost of constructing the plant, divided by the <u>annual</u> output of the plant, expressed in trillion Btu.

MA

operation & maintenance

SCURCES: Environmental Protection Agency, <u>Homitoring Environmental Impacts of the Coal and Oil Shale Industrian</u>, 600/7-77-015, February 1977, Camaron Engineers Incorporated, <u>Symiheric Fuela Mandbook</u>, 1975. Department of Energy, <u>Dreft Environmental Impact Statement for the (updated) Prototype Oil Shale Leaging Prograp</u>, 1979. University of Oklahama, <u>Taergy Alternatives: A Comparative Analysia</u>, 1975. Bechtel Corporation<u>, Energy Supply Flanning Model</u>, 1978.

UndergroundOilShale	Mining
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BIENCT STATEN:	RESOURCES USED: (Per 10 <sup>12</sup> Beu Produced)	)	RESIDUALS AND PRODUCTS: (Per 10 <sup>12</sup> Bin Produced)	
\$125 o 73,700 tong of raw shale per day	<b>FUEL</b>	Tone	ATR POLLUTANTS	Tone
• 0.413 x 1012 Btu/day	Tay unplaced thalo	170,450	particulates	4.46
a 2,800 Btu/pound of raw shale			50 NO 2	0.012
a 30 gallons/ton shale oil content	ENERGY			0.17
a aine opergias 328.5 days/year	electricity for operating	MA.	hyðrocarbona	0.019
a 24.2 x 10 <sup>®</sup> tons of shale minod/year,	drilling equipment and		C0	0.10
e total sanual output is 135.67 x 10 <sup>22</sup> Btu	trucko			
e mine life is 30 years		• •	WATER POLLUTANTS	
	CONPUSITION	<u>1 (by wright)</u>	probability of o dIBO	
DESCAIPTION	organic material	17.1	contamination • r under-	
e Underground mining uses roum and piller	water	1.4	ground valot with 0 im4 water	
cechnique. The oil shale deposit is	inorganic material	81.5	501 IB 111575	
entered through a tunnel dug into the	LAND <sup>(1)</sup>		SOLID WASTE	
elde of a valley where an outcrop appears. Pillere are left in place to provide roof	mine development	Acree 0.15	negligible (nee Proceesing)	
support at appropriate intervals. Entrac-	crushing	2.93	ENERGY PRODUCT	
tion is also accompliated by drilling and	( ) uentas	2.93	e trod e IM10 reck	178.450
blasting. The broken shale is transported	WATER	Acre-Feet		
to portable crusher for primary crushing.	mining and crushing	2.8 (1.3 - 3.3)		
CONPOSE IT S	COSTS	Dollere (1978)		
e drilling equipment	construction <sup>(2)</sup>			
e escavation equipment (creese)	manpower	164,224		
e cruehere	materiala	43,249		
e trucke	quipanc	254,842		
• • • • • • • • • • • • • • • • • • • •	other cost	02,341		
BIVIEUNERTAL CONCERNS	tot <b>al</b>	344,655		
• sir quality deterioration	Nrttl= & mintense	Le MA		
e noise				
e water requirement	PERSONNEL	Vorkere		
e contamination of underground water supplies	construction	484		
with saling mine water	operation & maintenance	MA		
		1		

(1) This represents land committed to use over the lifetime of the plant, divided by the <u>annual</u> output of the plant, aspressed in trillion Btu. (2) This represents total cost of constructing the plant, divided by the <u>annual</u> output of the plant, aspressed in trillion Btu.

SOURCES: Environmental Protection Agency, Monitoring Environmental Impacts of the Coal and Oll Shale Industries, 600/7-77-015, February, 1977. Cameron Engineers Incorporated, Synthetic Fuels Manibook, 1975. Department of Energy, Draft Environmental Tapact Statement for the (updated) Prototype Oll Shale Leasing Program, 1970. University of Oklahoma, Energy Alternatives: A Comparative Analysis, 1975. Bechtel Corporation, Energy Supply Fishning Model, 1976.

# TABLE 8 -

Coal	Beeneficiation

•

ENGT STSTEM:	NESOUNCES USED: (Per 70° Bew Produced)		RESIDUALS AND PRODUCTS: (Per 10 <sup>12</sup> BLu Produced)		
28 9 Process 2,037,000 tone of run-of-nine (BON) coal each year to produce 2 million tone of	PUEL FUE-of-oine (BOH) of raw coal	<u>Tone</u> 31,943	AIR POLLUTANTS Berticulates	Tene (Grees)	<u>Tene<sup>(3)</sup>(Het)</u>
cool ooch yoof to produce 2 million tone of clean cool	(assuming one ton of him coal	<i><i>M</i>.<i>MJ</i></i>	80,	2.1	0.005
• Nourly capacity 950 tase of NGM coel	hes an energy content 0 1 11		80	1.5	0.6
<ul> <li>Operates 3,000 hours per year, representing tan shifts per wesh, 230 days per year</li> </ul>	• Iltloa Dtwo per 800)		hy <b>f</b> recerbene CQ	1.1	0.1 0.2
e 20 year plant life	EMERCY <sup>(1)</sup>	_			
e \$7.5% efficiency (in terms of Btue)	alacticity	2.0 x 10 <sup>2</sup> him	VATER POLLUTANTS	Tone (Grees)	<u>Tone ( )) (Het)</u>
e yield by weight in 702	•11	2.8 ± 10 <sup>5</sup> kih 5.9 ± 10 <sup>8</sup> Biu	total dissolved solids	-143	- 11 -
	1.488	Acres(2)	iron December	0.2	0,007 0.03
© Coal beneficiation to a process for upgrading		0.2	aluminum	i.i	0.04
coal prior to its use for metallurgical of	loading facility	1.6	siac	0.04	0.005
utility purposes. The purpose of beneficia-	estiling pond	2.3	nichol	0.01	0.00)
tion is to remove imputities (i.e. ash and/or	•••		sulfatos	94	10
oulfur) from row cool. The degree of bonefi-	WATER	Acto-PL.	total suspended colide	5,670	0.6
cation depends on the type of coal and its	consumption	3.7	iron	4.4	0.06
ultimate use. The system described on this		B-11 (1834)	annos 1 a	0.2	0.05
summery sheet (level & per Phillips et al.)	COSTS	Pollers (1976) 4.3 # 107	SOLID WASTE	Ime <sup>(4)</sup>	
to a relatively intense procedure. It removes	construction execution and maintenance	3.2 x 10 <sup>5</sup>	primery breaking	<b>ATTE</b> .	
more pultur and ach than most other types of beneficiation, and it is also more costly.	operation and maintenance	3.4 H 10-	coores cleaning[5)	2	
beneficiation, and it is also more costly. The regultant classed cash would be used for	PER SOUMEL	Workere	tew-coal sizing	ě	
netallurgical purpose.	construction (1 year)	<b>1</b> 1	orimory cleaning	10,157	
mentionficat berbesser	operation and maintenance	1.5	froth flotation	5,341	
2001D/T1			thornal drying	0	
• •calping ecrees			breaking 004 O ItiS@	1	
0 crusher			tetal	13,302	
• tetery be-bar					
• vibrater ecreens			HEAT		
e jim			little Or mene		
40uatorlagmtpt			80158		
• thickeners			Noise ney e ffoct werkerei	and in	
• filtere			cleaning ceal, but there sh		
O concontrating tables O r hydroclosod O flotation circuits		-	little or no adverse impact		
o thermal drying		1	mear beneficiation PIO010.	•	
WEBOINERTAL CONCERNS			ENENCY PRODUCT	Tone	Heat Content
e erticul.t. entestene			cleaned coal	36, 360	13,750 Bcu/16
ealid wests disposal					
• • wrtaco water centamination from settling					
pond everties and/or refuse pile runoff					
· possible ground water contamination from					
• octilg pendleaching					
• octile pendiesching • poise					
♥ octilg <b>pendjesching</b> e seise					
• 10100					
• bolte	tent e f 12,000 btu/16 e t coal (Mit to alabarata (1 Jeval II) havefiet	tmon, 1974). They a	) . O otiooal everages (assuming		
<ul> <li>below</li> <li>These figures war, celculated accounting an emergy con an energy           filot=or e t \$1.33 and 40 oat           PQ1?     </li> </ul>	to elaberate (1 level 0) beneficia	at ion in particular.	- •		
<ul> <li>below</li> <li>These figures war, calculated accursing in emergy con a seargy filter or a fill. Bland 40 cat P PQ1?</li> <li>These coefficients my b. O ojoct to ersor 0 laco plant's ne" 1 were to pict to ene".</li> </ul>	to elaborate (1 level B) beneficial the data source presented only the find	ation in particular, a second of land upp	d without O pecifying the		
<ul> <li>below</li> <li>These figures war. calculated accusing an emergy con an energy fittet=or e t \$1.33 and 40 oat \$ PQ1?</li> <li>These coefficients may b. O ojoct to ersor 0 laco plant to merit surprise f cool. Is calculating the in the "size" of the other.</li> </ul>	P to elaborate (1 level B) beneficial the data source presented only the first se coefficients, it woo e aoad here	ation in particular, of amount of land upon that plant output we	d vithout O pecifying the ne the ex. B that O pacific4		
<ul> <li>beise</li> <li>These figures war. celculated accounting an emergy con an energy of filt=cr o t 91.33 and 40 oat 0 PQ1?</li> <li>These celficients may b. 0 ojoct to orsor o laco plant's o ne" every of cool. In celculating the in the "site" soction of this obset.</li> <li>These constants of the order of every of a strenges b</li> </ul>	? to elaberate (1 level 8) beneficial the data source presented on ly the first se coefficients, it woo • aoad here based upon regional coefficients project	ation in particular, of amount of land upon that plant output we cted by SEAS for 197	d vithout O petifying the sether x . B that O patified 9. The regional coefficients		
<ul> <li>beise</li> <li>These figures war, calculated accusing in energy con an energy filtct=cr e t \$1.13 and 40 oat P P017</li> <li>These cellications b. O ojoct to ersor 0 iaco plast 'o ene'l output of cool. In calculating the in the "size" sociate of this obtact.</li> <li>These figures o ro wighted o celciaal everages berge berge to the use of the bus used. Each of the series of the subscript set of the s</li></ul>	? to alaberate (1 level B) benefic: the data source presented only the fine as coefficients, it woo • acad here paced upon regional coefficients project coefficients above on this pheet is •	ation in particular, ad amount of land upp b that plant output up cted by SEAS for 197 gtrol to total nati	d without O pecifying the no the e x . 0 that O pacified 9. The regional coefficients implices of regional divided		
<ul> <li>below</li> <li>Theose figures war. calculated assuming an emergy con an energy of filt=cr o t \$1.33 and 40 oat O PQ1?</li> <li>Theose coefficients may b. O ojoct to o rsor 0 laco plant o ne"   output o f cool. Is calculating their in the "list" of the oheat.</li> <li>Theose figures o ro weighted o celciaal everages b ware weighted is terms o t bus used. Each o f the by tetal mational but output. Theose figures included</li> </ul>	? to alaborate (1 lovel B) beneficial the data source presented only the first as coefficients, it woo • aoad here pased upon regional coefficients project coefficients shown on this phase is 0 treativele from refuge pilse • 0 the	ation in particular, a amount of land upp b that plant output up cted by SEAS for 197 qtrol to total nation beneficiation prece	d without O pecifying the he theo x. B that O pacified 9. The regional coefficients longi tese of residual divided we itself. They 0 0 W4 that		
<ul> <li>below</li> <li>These figures war. calculated assuming memorgy con an essergy of filt=cr st 91.33 and 40 oat of PQ17.</li> <li>These coefficients and b. O ojoct to ersor elaco plant's one*i sutput of cool. In calculating the in the "sise" sociar of this sheet.</li> <li>These figures or on wighted o calculat a everages b were weighted in terms of thus used. Each of the by total national fits output. These figures include 400 et tool or noortion planto pr. closed erclose.</li> </ul>	? to alaborate (1 lovel B) beneficial the data source presented only the first as coefficients, it woo • aoad here pased upon regional coefficients project coefficients shown on this phase is 0 treativele from refuge pilse • 0 the	ation in particular, a amount of land upp b that plant output up cted by SEAS for 197 qtrol to total nation beneficiation prece	d without O pecifying the he theo x. B that O pacified 9. The regional coefficients longi tese of residual divided we itself. They 0 0 W4 that		
<ul> <li>below</li> <li>These figures war. calculated assuming memorgy con an essergy of filt=cr st 91.33 and 40 oat of PQ17.</li> <li>These coefficients and b. O ojoct to ersor elaco plant's one*i sutput of cool. In calculating the in the "sise" sociar of this sheet.</li> <li>These figures or on wighted o calculat a everages b were weighted in terms of thus used. Each of the by total national fits output. These figures include 400 et tool or noortion planto pr. closed erclose.</li> </ul>	? to alaborate (1 lovel B) beneficial the data source presented only the first as coefficients, it woo • aoad here pased upon regional coefficients project coefficients shown on this phase is 0 treativele from refuge pilse • 0 the	ation in particular, a amount of land upp b that plant output up cted by SEAS for 197 qtrol to total nation beneficiation prece	d without O pecifying the he theo x. B that O pacified 9. The regional coefficients longi tese of residual divided we itself. They 0 0 W4 that		
<ul> <li>moise</li> <li>These figures war, calculated accousing the energy cos an energy filter or a til. 181 and 40 oat PQ17</li> <li>These coefficients may b. O ojoct to 0 roor 0 laco plant's one"! eutput of cool. In calculating the in the "site" section 0 f this sheet.</li> <li>These figures 0 ro weighted 0 ociciaal everages b warg weighted in terms of the used. Such of the by total national bu output. These figures include 400 et coel or porotion plants or r. figures include 400 et coel or porotion plants or r. for figures include based ac ottocal 0 verages a 0 ittmm.</li> </ul>	to elaborate (1 level8) benefici. the data source presented only the find so coefficients, it woo e acad here beed upon regional coefficients project coefficients shown on this sheat is e e residuals from refuse pilos e d the 4 that e it refuse is treated. In e the Economics of Steam Coal Property.	ation in particular, and amount of land use b that plant entput we cted by SEAS for 197 qtrol to tetel metl beseficiation proce tticloacy of 908 (ind tton", Coal Hining a	d without • pecifying the to the • x . • that • pacified 9. The regional coefficients lemai tone of residual divided we itself. They • 0 0 W 4 that Btue) was • cau004.		
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<ul> <li>beise</li> <li>These figures war. calculated assuming momergy con memorgy filtet=cr e t \$1.33 and 40 oat PQ1?</li> <li>These coefficients my b. O ojoct to ersor elaco plant's e ne"! sutput of cool. is calculating the is the "size" section of this sheet.</li> <li>These figures ero weighted e celcia al everages b were weighted is terms o t bus used. Each of the by cetal matienal bu output. These figures include 901 e t cool e r.porotica plants e r. closed cycle .</li> <li>Based es e ottocal e verages is elitmm.</li> <li>UNCES: Phillips, Pater and Paul Delienze, "Assessing BOT and EFA, <u>Engineering/Econopic Analysis of</u> Witness Associates, Environmental Impatte, Eff</li> </ul>	to elaborate (1 level8) benefici. the data source presented only the find so coefficients, it woo e acad here pased upon regional coefficients proje- coefficients shown on this sheet is e e residuals from refuse pilos e d the 4 that e it refuse is treated. In Con- the Economics of Steam Coal Propers Coal Proparation with S0, Cleanup P ficience, and Coat of Enefect Supply	ation in particular, and amount of land use b that plant output use cued by SEAS for 197 qtrol to tests and beneficiation proce thicloacy of 908 (had clon", <u>Coal Mining</u> / processo, 1978,	d without • pecifying the to the • x . • that • pacified 9. The regional coefficients lemai tone of residual divided we itself. They • 0 0 W 4 that Btue) was • cau004.		
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#### APPENDIX TO CHAPTER 5, A: STATUS OF PROCESS DEVELOPMENT

The status of development of coal conversion and oil shale retorting both in this country and abroad are reviewed at length in the literature (Rogers and Hill, 1979; National Coal Association, 1980; Fluor Engineers and Constructors, Inc., 1979a,b,c).

The status of synfuel commercialization is summarized in the following tables:

- "Table A: Coal Gasification (Fluor Engineers and Constructors, Inc., 1979a).
- Table B: Coal Liquefaction (Fluor Engineers and Constructors, Inc., 1979b).
- Table c: Shale Oil Retorting (Fluor Engineers and Constructors, Inc., 1979c) .

These tables also review the major characteristics of these technologies. More details about the processes are given in Bentz, E.J., 1980.

# COVT EVRIFICATION - COMMERCIALLY PROVEN PROCESSES

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IABLE A-1 (continued) Coal Gasheration - Processes Ready for Demonstration

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cual Gasification - 1	PRODUC15	Hogin RTU gas, char, and tuttur	Fuel ges writelie fue SNG: clast, within and by product out	Low BTU ges and utilu	fuel ger suitable for SNC, cher, and auflur	Low BTU ges and suffur	Synthesis ges and auflur	adiu: adiu:
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# TABLE A-1 (continued) COAL GASHFICATION - PROCESSES READY FOR COMMERCIAL DEVELOPMENT

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8	. INC Corporation		ł	1	secologia	1.400 L (1) Ien dat 1.80.00 L (1) Ien dat 8000 SC/ Ien ga 1276 d (10,600)		Buccostulty piloted an 38 TD pilot, dange computed permutation	herunia an el 200 FD pera a combra 200 FD pera a combra an esta COGAE P. e.c.a Perty Council, liturata		1 Limited Io here beduc const 2 Limited press require press command
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OCCIDENTAL	Accelerate Deserves and Deserves and	Lightia Lightia Bitantiana		Continued too	Armoughur to	1100 Lives Am 1100 Lives Am 100 S67 (an there 2000 S67 (an there 2000 B10/BCF)	Part I control into Part I control into Part Part of Part of Part of Part of Part Part of Part of Part	36 TPD oriest parts	Terringues lo Bendinus de Dunge el 260 F/D Polet pens under any	rands from the results	1 High willin cher
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# TABLE A-2 Coal Liquefaction - Pyrolysis process

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	TABLE $A-2'$	(continued)
COAL LMMJEFACTION	- HYDROGENA	FION PROCESSES

MAME	DEVELOPER	PEEDSTOCKS	REACTOR BYET SM	CONVERSION CONDITIONS	TYPICAL PRODUCTS VIELDS	HYDROGEN CONSUMPTION	REMARKE	PILOT PLANT	PUTURE PLANS FOR PROCESS	PROCESS ADVANTAGES	PROCESS DIBADVANTAGES
CSF	Canaca Cual Development Campany	Lignitë Sub-bitumineus Bitumineus	Stirred surry Satistics plus shullated bod catolytic Nydrogenesies	EM TRACTOR 180 pro, 148 <sup>4</sup> 7 HY DROTREATIN 2000 pro, 800 <sup>4</sup> 7	2 0 88L/Ion ol 3400 SCF /Ion geo 1633 8TU/SCP	7600 SCF /Te , limw 16 000 e Sc F/T hydrotrening	H <sub>2</sub> supplied by donse solvent in actiector which is continuously recovered and rehydrogeneted ber recycle to an wector	20 TPD pilot plant In start up at Grange, West Virginia			Pressurred storry Ned system     Michani di ostactea     under frassira     Nydrotrasing of     autract required
	Érron Rossersh and Engineering Chargony	Lignita Sub-bitumingus Bituminous	Proprietary Ba- tractor Flace bed catalytig solvent hydrogenation	1800 - 2600 prig 7000 - 8004p	~3088L/ion love sulfur fuel eli	6000 6000 Basification of institute Isoal and reforming of product gas to supply H	Politowa C&F scheme, where dance solvent is Continuously racovered and ra hydrogenated fei macycle to she awyector	I TPD © Wi plant in sparation	Design and constructing at 250 TPD priori glant	<ol> <li>Different product date per be obtained by usying operating conditions</li> </ol>	I High pressure shurry Bood system
SRC .	04.11 Gu Corporation	Lignite Bub bitumingus Bitumingus	M. Is pie vertical tubo dissolver	1000 pris 81847	1100 L S/ian Ikw sulfur, ian shi char (18 000 STU/LS) 9 1% osh	3000-6000 SCF/mn	Gaseo ut hydrogen uaed m driasius far hydrotresiung	50 TDP plant operating or F1 Laws, Wash 6 TPO pilot plant at Wildowillig, Alabama	Design af two 6000 M fermonalization please Des suring SPC1 ge other SPC11	High conversion room     Phip physical column     properation required	1. High pressure durry lead synam
						7					
H-COAL	Hydracarbon Raeserch, Inc.	Lignite Bubibituminoue Bituminous	Ebuilated bod Coloryst	2200 2100 pag 800° 878 <sup>9</sup> P	~ 3.0 BBL/Ion eyninotic crude 18º API 30% fuel gas	13,000-18,000 SCF/T	Lover H3 consumption yields now within fuel or higher H3 Lagors yields the 18 <sup>®</sup> API syncrude	Buccessfully tested in 3 TPD plant	Construction of BOD 7/D domensisation plant underway	Based on commercialized H Orl process for heavy residues Continuous repronative cations produces constant attivity Leve Rydrogen consumption	<ol> <li>High prosure sturry load system</li> </ol>
SYNTHOIL	U S. Burseu of blings	Lignita Bibbatuminaus Bituminaus	Frank bad Coblyst	2000 4000 paig 800° f	~ 3 9 88L/ion 200 ( 8/ion chy (17 000 8TU/L8) 3000 8CP/T fuel ges	4000 + SCF/T Suggest using a gesification schema, but none specified ar zested	The apparent very and relidence time required mee attractive H ackey mine prociscal Later teers phowed H wee not	1/2 170 pilotpiont Ad longer in operation	Construction completed in 18 TPD plot plant, lever utilized.	High Injurd yields     Con handle moderataly     heavy sturies     Low hydrogen requirement	E High pressure sture " feed system 2 Solid resulue contains high aniumta 0 functionwerted coal 3 High hydrugen recycle retrie 1. Lare work manad teccien retrie
CCL	Guil Ós Corporation	Lignite Gub bituminaus Bituminaus	Franci band Coloriyat	2000+ proj 750* 1000*r	~ 3 0 BBL/ion tow suitur lust ad	16.000 22.000 SCF/mi Indicate reforming of product per for Hg	Short residence teme Bipecial reactor design Cohe resistant, high Bitiwity catalyst	1 TPD priot unit in aperation	The 600 TPD p-log plant is currently in decap	<ol> <li>Cotoryst exhibits high resistance to carbon deposition</li> <li>High liquid yields</li> </ol>	Brigh prossure stars " feed system

.

# table A-3

#### SUMMARY OF RETORTING PROCESSES

	GAS COMBLISTION		ARAHO KILN	ARAHO KILN IALT.J	'ETROSIX	SUPERIOR'S GRATE RETORT	0600	MION RETORT "A"	union retort ""	LANCH SQR	OCCIDENTAL IN SITE
VELOPER	reau of Mines	.urgi Ruhigas	'araha Dev Carp	irsha Dav. Corp	<sup>1</sup> etrobres, Cameron & Iones	iuperier Oli Compony	ha Cui Shala Carp.	Jinen Oil Co. of Calif	Union Oll Co. of Calif.	Union Ott Co. of Colif.	Occidental Potroloum C
ILE FOR FING IANSFERF	ternai cumbusian ad ant shala carbon jidua	- Sut circulating polid Asht Nu internet ombustion	isternal combustion of perit shale carbon asidue.	ut recycle gas Nu Imbusian in retart	tus recycle ges No lombustion in reters.	isher direct (with period simplication of recycle just or indirect.	lot circulating caraftic alls. No internal ambietten.	nierzei combustien ef part shale earben pridue.	Hat recycle gas Ale sombusudit in retort.	Het recycle gas, with escen and het gas from each gas-lar.	internel someration of grant shale carbon Radue phis recycle ga
Dele ()µsreted	160 1. 1952 6, 1964 6	1 <b>860</b> s, 1976	. Invisione (Comm.) 1969 se dele 1974 se dele	D76	1972-To date.	1876 to delo.	965-7, 1885-8, 971-2	) <b>848, 1866-8</b>		1073-74	1872 to data.
Tare Per Day	26,160	•			2000	NO TPH Manimum cape- iily — operated © about 16. TPH	16,1000	2,30,380 to 1200	Laborasory anly.	3	Nat readily gravitable for articl
AI OPERATION	ior yas — suistă intacting resulting slicitor formation, autil not operate web grade shale laver 30 gel/tan),		Yi ugi am fur oli ghate julut prent operation jurigi ated Erecuting junitect to produce 100 000 blut of oli for jelining taste	itot plant operatos sping i Patrosia operation sing U.S. shale at 20-38 gi/lon.	Uporating with Brazil Ini shale (23 gol/tun) Have had sume rotiun- ing of shale oil which furms terry substation and pings bod.	Circular traveling grath, plapted from von ore untering Operated by Superior at M. Daviel Nellingin facilities – Claveland Ohio.	<sup>1</sup> ilos plent operation uccessful Full scale Hent (50 000 870) In protiminary angen- tering stage.	Utilizes rock pump Process appears werk- pho II commercial prod oquipment apn be developed.	Uniters rach pump. Process approves wark- able if commercial- wied equipment aph in developed.	Unilians rach pump and apparase aska gentlar	First large scale unit (1976) produced 20.00 bbil, but had low yraide due te poor rubbilap tion Second gwe wyr poor yrbith due te ungegleined channelin Thind is under way.
/I NEQUARED	14'' ω 3''	Loss than 1/2"	1/4 " 103"+	/4" 10 3'"	1/4" to 3"	-3"	Loss than 1/2"	I/8" ⊷ 3"	1/8" to 3"	1/6" es l'	Nat reparted
*AP1	¥6		21.4	11.7	19.6		21	21	31	21.6	26.0
L Surlur, Will S	<b>4</b> 0		0.61		1.06		38	6.7		£7	• 44
Cl Notrugen We	16		1 <b>#</b> 5		094		1.8	1.0		1.0	911
Pour Point, D	80		-	•	*	90	80	89	m	20	*
MALE UNE VIEL D PREMER ASSAY	2 - 87		94 96	)7 )7	00 - 00 (Est )	96 - 409		<b>9</b> 1		100	Much depends on depres of rubbilitation
~	lesorting is pithter to estru apparach. De etypnen: work needed in suind distribution nd er ges distribution yttem	Currenterical plant of 1700-1710 in spectrum using pradriad in uses coal Hair 101-1 of Latingue brane in late 1976, Lin 2 week, pilot test. Date not available but repurchally can very smoothip	Haturt has wurked well in deniuntration prent built at 8 of M Anvel Pronts facilities Have nade end stored 80,000 bail of the 100,000 bbt to be refined for U.S. Newy	An alterñais echama tarrionsraisd al Anus Painte plant,	Process should work using two grade shale - * 30 yearson. Autosteene devolupment work neeted to utilize high grade shale.	Claims so have eliminate must environmental ubjections. Recovers nationale, polos ean, and elimina es setette by- produces. Reduced volume of esclus permits returning of solid waste to exces	Machanical and technical problems asem to be tasolosid. Process appears to Le one of most efficient to uses. Tooca clama it is tuity devid- oped on dema scale and is ready for communical application. Final E15 has been tasee by BLM.	Union believe technol- ogy established for design of 60001/D rotuit. They are pro- conting with plane for privately financed full-acate module.	Retort "8" could be expressed in combin- ation w/Retort "A" to yoold 20 Vol 3, to Fecher Assay.	High 182%) sharmal efficiency: Proceeding with 1800 170 dome reset und 6000 170 grossitype colide pum	Model an priv, wrth 16 or 2016 of deposet removed, followed by probume fracturing by probum retoriates rubble Courds be connermical even for lean shakes 110 pa/TI-

# APPENDIX TO CHAPTER 5, B

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#### Appendix B-1: EXISTING U.S. AND FOREIGN LOW- AND MEDIUM-BTU GASIFICATION SYSTEMS

(Courtesy: "Low and Medium Btu Gasification Systems: Technology Overview U.S. EPA, 1978; EPA-600/7/78-061)

Gasifier	Licensor/developer	Number of gasif Low-Btu gas	iers currently o Medium-Btú gas	perating (No. Synthesis gas	of gasifiers Location	built) Scale
Lurgi	Lurgi Mineralöltechnik GmbH	5	(39)	(22)	Foreign	Commercial
Wellman-Galusha	McDowell Wellman Engineering Co.	8(150)			US/Poreign	Commercial
Woodall-Duckman/ Gas Integrale	Woodall-Duckham (USA) Ltd.	(72)@@		(8)**	Foreign	Commercial
Koppers-Totzek	Koppers Company, Inc.			(39)**	Foreign	Commercial
Winkler	Davy Powergas		(23)**	6(14)	Poreign	commercial
Chapman (Wilputte)	Wilputte Corp.	2(12)	-	-	us	Commercial
Riley Morgan	Riley Stoker Corp.	1	-	-	us	commercial
Wellman Incadenscent	Applied Technology Corp.	(2+) ++	-	-	US/Foreign	Commercial/ Demonstration
BGC/Lurgi Slagging	British Gae Corp. Ond Lurgi Mineralöltechnik GebH		1		Poreign	Demonstration
Bi-Gas	Bituminous Coal Research, Inc.	· ·	1		us	Demonstration
Foster Wheeler/Stoic	Foster "Wheeler/Stoic Corp	1*(2)**	-	-	us	Demonstration
Pressurized W <b>ellman-</b> Galu <b>sha (NERC)</b>	ERDA	1*			us	Demonstration
GFERC Slagging	ERDA	-	1+		us	Demonstration
Texaco	Texaco Development Corp.	-		1*	us	Demonstration
BCR LOW-Btu	Bituminous Coal Research, Inc.	1•			us	Demonstration
combustion <b>Engineering</b>	Combustion Engineering Corp.	1+			us	Demonstration
Hygas	Institute of Gae Technology	-	1		us	Demonstration (High-Btu)
Synthane	ERDA	-	1		us	Demonstration (High-Btu)
C0 <sub>2</sub> Acceptor	ERDA	-	1		us	Demonstration (High-Btu)
Foster Wheeler	Foster Wheeler Energy Corp.	1			us	Pilot
Babcock & Wilcox	The Babcock & Wilcox Co,	1			us	Pilot
U-Gas	Institute <b>of</b> Gas Technology, Phillips Petroleum Corp.	1			us	Pilot (400 lb/hr coal)
Westinghouse	Westinghouse Electric Corp.	1			us	Pilot
Coalex	Inex Resources, Inc.	(1+)			us	Pilot

Under construction. Demonstration scale indicates 2000 to 10,000 lb/hr coal feed. Pilot scale indicates 400 to 1500 lb/hr coal feed.

Undetermined number overseas currently in operation.

# Appendix B-2: Continued

# POPULATION OF LOW/MEDIUM-BTU GASIFIERS

Gasifier name	Licensor/Developer	Stacus
d-Sod, Dry Ash		
Jurgi	American Lurgi Corp.(USA)	Present commercial operation
Wellman-Galusha	McDowell Wellman Engr. Co. (USA)	Present commercial operation
Chapman (Willputte)	Wilputte Corp. (USA)	Present counsercial o p e r a t
Voodall-Duckhan/Gas Integrale	Woodall-Duckhan, Ltd. (USA)	Present commercial operation
Liley Morgan	<b>Eiley</b> Stoker Corp. (USA)	Present demonscration unit testing; commercially available
Pressurized Wellman-Gelushs (MERC)	Norgantown Energy Res Center/ERDA (USA)	estebent development unit testing
oster Wheeler/Stoic	Foster Wheeler Energy Corp. (USA)	Demonstration unit planned
iinges	Allis Chaimers Corp. (USA)	Present development unit testing ; commercially • vallabla
Lellogg Fixed Bad	M. u. Kellogg Go. (USA)	Present 118v'dome unit testing
IBGAS	General Electric Research and Development (USA)	Present development unit testing
consol Fixed Bed	<b>Consolidation Coal</b> co. (USA)	Present development unit testing
FE Two Stage	International Furnace Equipment Co., Ltd.	Past commercial operation
erpely Producer	Bureau of Mines/ERDA (USA)	Past commercial operation
larischka	Daizova	Past commercial operation; anthracite or coke only
Matsch Hillebrand	Unknown (Germany)	Past commercial operation
I.G.I. Blue Water Gas	U.G. I .Corp. /DuPost (USA)	Fast commercial operation; coke only
over Gas	Forer Gas Co. (USA)	Past commercial operation
lellman Incandescent	Applied Technology (uDA)	Present commercial operation
CR/Kaiser	Unknown	Past development unit testing
ed-Bed. Slagging Ash		
BGC/Lurgi Slagging Gasifier	British <b>Ges</b> Council (G3) Lurgi Mineralöltechnik (W. Germany)	Present development unit testing
GYERC Slagging Gasifier	Grand Forks Energy Research Career/ERM (USA)	Present development unit testing; lignite only
Luena	Uaknown	Put Commercial operation; coke only

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# POPULATION OF LOW/MEDIUM-BTU GASIFIERS

Gasifier name	Licenser/Developer	Status	
dized-Bed, Dry Ash			
Winkler	Davy Powergas Co. (USA)	Present commercial operation	
Hygas	Institute of Gas Technology (USA)	Presentdevelopment unit testing	
Synthane	Pittsburgh Energy Research Center/ERDA (USA)	Present development unit testing	
Hydrane	Pittsburgh Energy Research Center/ERDA (USA)	Present development unit testing	
Cogas	Cogas Development Co. (USA)	Present development unit testing	
Exxon	Exxon Corp. (USA)	Present development unit testing	
BCR Low-Btu	Bituminous Coal Research (USA)	Present development unittesting	
CO <sub>2</sub> Acceptor	Consolidation Coal co. (USA)	Present development unit testing	
Electrofluidic Gasification	Iowa State Univ./ERDA (USA)	Presentdevelopment unittesting	
LR Fluid Bed	Unknown (Germany)	Past commercial operation	
<b>BRI</b> Fluidized Bed	Hydrocarbon Research Inc. (USA)	Past development unit cesting	
BASF-Flesch-Demag	Badische Anilin und Soda Fabrik (West Germany)	Past development unit testing	
GECB Marchwood	Unicova	Past development unit testing	
Heller	Unknown (Gerffiny)	Put development unit testing	
uidized-Bed, Agglomerating Ash			
U-Gas	Institute of Gas Technology (USA)	Present development unit resting	
Bactmlle/Carbide	Battelle Memorial Institute (USA)	Present development unit testing	
Westinghouse	Vestinghouse Electric Corp. (USA)	Present development unit testing	
City College of NY Mark 1	Hydrocarbon Research Inc./ A.M. Squires (USA)	Present development unit cesting	
Two-stage Fluidized	British Gas Council (England)	Present development unit testing	
ICI Hoving Burden	Imperial Chemical Industries, Ltd. (England)	Past development unit testing	
trained-Bed, Dry Ash			
<b>Garrett</b> Flash Pyrolysis	Garrect Research and Development co. (USA)	Present development unit testing	
Bianchi	Unknown (France)	Pase development unit testing; lignite only	

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Gasifier name	Licenser/Developer	Status
Pamindco	<b>Unknown</b> (France)	Past development unit testing; lignite only
USBN Annular Recort	Bureau of Mines/ERDA (USA)	Past development unit testing; lignite only
USBM Electrically Heated	Bureau of Mines/ERDA (USA)	Past development unit testing
atrained-Bed, Slagging Ash		
Koppers-Totzek	Roppers Co. (USA)	Present comercial operation
Bi-Gas	<b>Bituminous Coal Research, Inc.</b> (USA)	Present development unit testing
Texaco	Taxaco Development Corp. (USA)	Present development unit eating
Coalex	Inex Resources, Inc. (USA)	Present development unit testing; commercially ● milabla
PAMCO/Foster Wheeler	Pittsburgh O od Midvey Coel co. / Foster Wheeler (USA)	Present development unit testing
Combustion Engineering	Combustion Engineering (USA)	Present development unit cesting
Brighen Young University	Brighen Young University/ Bituminous Ceal Research (USA)	Present development unit testing
Babcock and Wilcox	The Babcock and Vilcox CO. (USA)	Past comercial operation
Ruhrgas Vortex	Ruhrgas A. G. (West Germany)	Past commercial operation
IGT Cyclonizer	Institute of Gas Technology $( USA)$	Pastdevelopment unit testing
Inland Steel	Inland Steel CO. (USA)	Past dcvdopment whit testing
USBN, Morgantown	Morgantown Energy Research Canter/IIDA (USA)	Past development unit testing
Great Northern Railway	Great Morthern Railway Co. (USA)	Past development unit testing
TRS Cyclone	Unknown (England)	Past development unit testing
olten Media, Slagging Ash		
Kellogg Holten Salt	N. W. Kellogg CO. (USA)	Present development unit testing
Atgas/Patgas	Applied Technology Corp. (USA)	Present development unit testing
lockgas	Atomics International (USA)	Present development unit testing
Rummel Single Shaft	Union Rheinische Braun Kohlen Kraftstoff A. G. (Vest Germany)	Past comercial operation
Sum Gasification	Suo Research and Development Co. (USA)	Past development unit testing
Otto-Rumei Double Shaft	Dr. C. Otto and Co.	Past development unittesting

# POPULATION OF LOW/MEDIUM-BTU GASIFIERS

# APPENDIX TO CHAPTER 5, C

# COOPERATIVE AGREEMENTS AND FEASIBILITY STUDY GRANTS FOR SYNFUELS

(Source: U.S. DOE 7/80)

AGREEMENTS	DESCRIPTION/SITE		Texas Eastern Synfuels proposes to construct a coal liquefaction facility which will produce the equi-	valent of 56,000 barrels of oil per day. Texas Eastern Synfuels is a joint venture of Texas Eastern Corporation, and Texas Gas Transmission Corporation. Proposed project is a Fischer Tropsch plantlike the SASOL facility in South Africathat would convert appruximately 28,000 tons per day of coal into a mixture of transportation fuels, Synthetic Matural Gas (SNG), and chemicals. Approximately 44 percent of the output is SNG (145 nmSCF/D); about 30 percent transportation fuel, and the test chemicals Site is near Hendersen, Kentucky.	The project will employ a Lurgi pressurized, fixed bed gasificat on process with Lurgi methanization re- quiring 14,000 tons/day of lignite coal to produce 137.5 mmCF/day of synthetic gas, 9∃ tons of ammonia/ day and 85 tons/day of sulfur. The facility will be sited in the Beulah Hazen area of Mercer County, North Dakota and has a total capital requirement of \$1.5 billion
COOPERATIVE AGREEMENTS	REQUESTED FROM DOE		<b>\$</b> 24 <b>,</b> 300.∞	-	<b>\$</b> 22 <b>,</b> 000 <b>,000</b>
	TECHNOLOGY	Coal Liguids	Texas Eastern Synfuels	<u>High Btu Gas</u>	Great Plains Gasification Associates

Wycoal Gas \$13, 155, 000	Wycoal plans to construct a facility using Lurgi and Texaco gasification units to process 16,000 tons of sub-bituminous coal daily to produce high Btu gas. All liquid by-products will also be gasified. The facility is to be located in Douglas, Wyoming. The Statement of work <b>proposed</b> will involve developing a definitive basis for plant design estimating costs, securing permits and approvals, obtaining financing and identifying long-lead delivery items. There is a market for the SNG via a pipeline system to the midwest owned by the participants. The project would produce the equivalent of 51,000 barrels of oil per day.
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	FEASI BI LI TY STUDY GRANTS						
TECHNOLOGY	REQUESTED_FROM_DOE "	<u>DESCRIPTION/SITE</u>					
<u>Coal Liquids</u>							
Cook Inlet Region Anchorage, Alaska 99509	\$3,900,000	Feasibility study of producing 54,000 barrels per day of methanol from low sulfur coal using Winkler gasi- fier and ICI methanol synthesis. Site: West side of Cook Inlet, Alaska					
₩. R. Grace Denver, Colorado 80223	\$786, 477	'Stage 111 of a feasibility study of a coal sourced methanol plant using a Koppers/Totzek Gasifier. Site: Moffat County, NW Colorado					
Clark Oil & Refining Milwaukee, Wisconsin 53227	\$4, 000, 000	Feasibility study of producing synthesis gas from coal, steam, oxygen & methanol from synthesis gas using a KT Gasifier, ICI & the Mobil M Process. Site: S. Illinois					

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ejb&a

Fourteen month feasibility study of producing fuel grade methanol from coal using Ziegler ≓oal deposits. Site: Covent, Louisiana	Feasibility study of a coal to methanol plant pro- ducing 14,910 barrels per day using Koppers or Lurgi Gasifiers. Site: Diluth, Minnesota	Feasibility study for constructing an 85,000 barrel/ day coal to methanol plant using Lurgi gasifier and Lurgi methan⊂l synthesis. Site: Dunn, North Dakota	Feasibility study of gasification, in-situ deep Texas lignite and conversion of remaining medium BTU syn- thesis gas to methanol and high octane gasoline. Site: Somewhere in Texas Gulf Coast	Ten-month feas bil ty study of converting 15,∞ tons of coal/day to 20,∞ barrels/day of gasoline * S te: G llette, Wyoming		Nine month feasibility study, High Btu Gas (Lurgi Process - SNG) at Crow Reservation, MT. Site: East of Billings, MT.	Nine mon\$h feasibility study, High Btu Gas (Lurg Process ∺ SNG, Methanol) at San Juan County, New Mexico. Site: Easc of Navajo Indian Reservation
				-			
\$3,260,000	\$2,190,000	\$4,000,000	\$808,781	\$1, <del></del> , 000		\$2,729,393	<b>\$3,</b> 018,00 <b>~</b>
Houston Natura Gas/Texaco Houston, Texas 77001	AMAX, Inc. Grenwich, Connecticut	Dakota Company B smark, North Dakota 58501	Republic of Texas Coal Co. and Mitchell Energy Corp. Houston, Texas 77002	Hampshire Energy Hilwaukee, Wisconsin	<u>High Btu Gas</u>	Crow Tribe of Ind ans Washington, D.C. 20036	Texas Eastern Synfuels, Inc. Nouston, Texas 77001

	Twelve month feas bil ty study of Med um Btu Gas Combined cycle. Site: Pinellas Courty, Florida	Nine montıfeasibil ty study of low Btu Indus ral Fuel Gas. Site: Florence, Kentucky	Fifteen month feasibility study of combined cycle, medium Etu gas at Sears Island, ME. (Process: Texaco Gasifier) Site: Waldo County, Maine	Feasibility study for a medium Btu gasificati∽n fa- cility producing combined cycle power and methanol. Choice or process technol⊖gies between Koppers-Totzek or Slagging Lurgi. Site: Fall River, Massachusetts	Twelve month feasib lity study of medium Btu gas (Process: TRD). Site: Philadelphia, Pernsylvan≎a	Feasibility study to determine the egin sal and economic viability of developing a carbon moroxide and hydrogen syngas from either a high Btu coal or a Texas lignite. Site: Hear Bishop, Texas	Eighteen-month feas bility study of low/Med um Btu Gas. Site: Texas City, Houston, Texas
	<b>\$1,</b> 380,796	\$922,555	<b>\$3,624,558</b>	\$4,000,000	<b>\$1,</b> 168,108	No cost	<b>\$</b> 3,945,676
Low/Hed um Btu Gas	Florida Power St.Petersburg, Florida 33/33	General Refractories Bala Cynwyd, PA. 19004	Central Ma ne Power Augusta, Ma e 04336	EG&G Wesley, Massachusetts 02181	Philadelphia Gas Works Philadelphia, PA 19102	Celanese Corp Dallas, Texas 75247	Union Carbide/Linde Division Tonawanda, New York 14150

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	Feasibility study for upgrading crude ~ ] shale to gasoline jet fuels, DFO and res dual us ng UOP hydro- processing & hydro-cracking. Site: Fru ta, Colorado	Eighteen month feasibility of 2000 BPD (or larger) module of a 50,000 BPD plant. Site: Lewis County, Kentucky		Eight-month feasibility study of extracting 20, <sup>000</sup> barrels/day of oil from domestic tar sands - Bitumen. Site: Site may be in Utah or California	. Feasibility study of a 5º 000 barrel/day Tar Sands Bitumen facility. Site: Sunnyside, Utah		Feasibility study of anaerobic digestion of sewer water to obtain methane. Site: Possibly Oakland, California	Feasibility study of the recovery of natural gas from Devonian Shales - vertical wells. Methane from Devonian Shale. Site: Salamanca, New York
					-			
	<b>\$3,009,399</b>	\$3,778,267		\$357,511	0		\$440,261	<b>\$</b> 896 <b>,</b> 638
Oil Shale	Gary Energy Corp. Fruita, Colorado 81521	Transco Energy Co. Houston, Texas	Tar Sands	Natomas Energy Co. San Francisco, Cal forn a 94108	Standard Oil of Indiana Ch'sago, Illinois 60601	Unconventional Gas	Acruex Corporation Mt. View, California 94-42	Seneca Indian Nat on Salamarca, New York 14779

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Republic of Texas Coal Co. and Mitchell Energy Corp. Houston, Texas 77002		Feasibility study of gasification, in-situ deep Texas lignite and conversion <b>of</b> remaining medium Btu synthesis gas to methanol and high octane gasoline. Site: Calvert, Robertson County, Texas
Mountain Fuel Supply Co. Salt Lake City, Utah 84139	\$1, 810, 762	Two-year feasibility study of unconventional natural gas in the Pinedale field. Product is natural gas and condensate. Site: Sublette County, Wyoming
Peat		
Minnesota Gas Co. Minneapolis, Minnesota 55402	\$3, 996, 554	Nineteen month feasibility study for the production <b>of</b> high Btu substitute natural gas from peat. Site: Minnesota
<u>Shal</u> e Liquid Upgrading		
Union Oil Energy Mining Los Angeles, CA 90017	, \$4, 000, 000 ,	Feasibility study for operation of a 10,000 BPD up- grading plant producing premium quality syncrude. Site: Grand Valley, Colorado

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It Cliffol. OGY     REQUEST O FROM DOE       Unconvention     \$600,000       U.S. Steel Corporation     \$600,000       Bankl ck Corporation     \$999,500	COOLEWALTKE MANEEMEINIS	DESCR PT ON/SITE	U.S. Steel Corporation proposes to build a collection and compression system to capture methane from a mine pre-drainage program. The gas, currently being vented, will be injected into an interstate pipel ne system for sale. The project will produce the equ valent of 200 barrels of oil per day. Site s Oak Grove, Alabama.			Banklick Corporation proposes to design and construct a Coal Mining Mixture (COM) preparation plant on a site on Blount Island, Florida owned by the Jacksonville Port Authority and to market the products. In this proposal, the approach is to first grind the coal, then mix it with oil and pulverize the result, and, finally, to mix the product more thoroughly using ultrasonic agitators. A COM prep plant is relativley simple and, in addition to the above equipment, cons sts o <sup>c</sup> coal storage and handling equipment (including a coal pile), oil and COM piping and storage hardware, and associated hardware. Coal would be delivered by rail. The project will produce 6,000 barrels per day.			
The Chinol. OGY Unconventional Gas U.S. Steel Corporation Bankl ck Corporation		REQUEST O FROM DOE		<b>\$600,0</b> 00		<b>\$989 ,</b> 500			
C-8		TECH101.0GY	<u>Unconven</u> eional Gas	U.S. Steel Corporation					

COOPERATIVE AGREEMENTS

- GLOSSARY (Courtesy: Coal Liquefaction Quarterly Report, U.S. DoE, May 1979)
- absorptio an imprecise term suggesting the taking up of one substance by another by either a physical process or a chemical combination.
- acceptor calcined carbonate that absorbs carbon dioxide evolved during gasification. liberating heat.
- acid gas removal the process of selectively removing hydrogen sulfide and carbon dioxide from a gas stream.
- activated carbon carbon obtained by carbonization in the absence of air, preferably'in a vacuum; has the property of absorbing large quantities of gases. solvent vapors; used also for clarifying liquids.
- diabatic any process where heat is neither given off nor absorbed.
- dsorption the process by which the surface of a solid or liquid attracts and holds any atom. molecule. or ion from a solution or gas with which it is in contact.
- gglomerate assemblage of ash particles rigidly joined together. as by partial fusion (sintering).
- nttmcits coal hard coal containing 86 to 98 percent fixed car- " bon and small percentages of volatile material and ash.
- API American Petroleum institute.
- API gravity— a scale adopted by the API for measuring the density of oils; °API = \_\_\_\_\_\_ 141.5 - \_\_\_\_\_ - 131.5

#### Specific gravity. 60° F 60° F

- romatic hydrocarbon a cyclic hydrocarbon containing one or more six-carbon (benzene) rings.
- sh solid residue remaining after the combustion of coal.

ASTM — American Society for Testing Materials.

- autoclave a vessel, constructed of thick-walled steel for carrying out chemical reactions under high pressures and temperatures.
- bench-scale. unit a small-scale laboratory unit for testing process concepts and operating parameters as a first step in the evaluation of a process.
- . binder carbon products. tars. etc.. used to impart cohesion to the body to be formed: a coai-extract binder may be used to prepare formed-coke pellets from non-coking coals.
- bituminous coal a broad class of coals containing 46 to 86 percent fixed carbon and 20 to 40 percent volatile matter.
- blow down— periodic or continuous removal of water from a boiler to prevent accumulation of solids.
- bottoming cycle~ the lower temperature thermodynamic power cycle of a combined-cycle system.
- Btu British thermal unit. the quantity of energy required to raise the temperature of one pound of water one degree Fahrenheit.
- BTX— benzene. toluene. xylene; aromaticc hydrocarbons.
- caking the softening and agglomerartion of coal as a result of the application of heat.
- calcination the process of heating a solid to a high temperature to cause the decomposition of hydrates and carbonates.
- calorific value— the quantity of heat obtained by the complete combustion of a unit mass of a fuel under prescribed conditions.

- carbon  $_{\rm fiber\,-}$  fine filaments of carbon about eight microns in diameter which  $are\,$  used in composite materials, being bound with resins.
- carbonization destructive heating of carbonaceous substances with the production of a solid. porous residue or coke. and the evolution of a number of volatile products. For coal. There are two principal classes of carbonization. high-temperature coking (about 900" C) and low-temperature carbonation (about 700° C).
- $_{\tt catalyst}$  a substance that accelerates the rate of a chemical  $_{\tt reaction}$  without itself undergoing a permanent chemical change.
- centrifuge an apparatus rotating at high speed which utilizes the centrifugal force generated to separate materials of different densities. e.g.. undissolved residue from coal solution in the SRC process.
- char the solid residue remaining alter the removal of moisture and volatile matter from coal.
- Claus process— industrial method of obtaining elemental sulfur through the partial oxidation of gaseous hydrogen sulfide in air followed by catalytic conversion to molten sulfur.
- coal a readily combustible rock containing more than 50 weight percent and more than 70 volume percent of carbonaceous material including inherent moisture. formed from compaction and induration of variously altered plant remains similar to those in peat.
- coalification metamorphosis of vegetable debris into coal.
- coke strong porous residue cons] sting of carbon and mineral ash formed when bituminous coal is heated in a limited air supply or in the absence of air. Coke may, also be formed by thermal decomposition of petroleum residues.
- coke breeze the fine screenings from crushed co kc usually passing a 1 2 inch or 3 4 inch screen opening.
- combined cycle two sequentai thermodynamic power conversion systems operating at different temperatures.
- combustion gas gas formed by the combustion of coal. e.g.. burning.
- combustor a vessel in which combustion taken place
- coupon a polished metal strip used to measure the rate 01 corrosion of the metal in a specific gaseous or liquid environment.
- cracking the partial decomposition of h l~h-m~)lecuiar-weight organic compounds into lower-molecular-weight compounds. generally as a result of high temperatures
- crude <sub>gas</sub> impure gas produced in a gasifier
- culm the waste or slack from anthracite mines or preparation  $_{\rm plans}$  consisting of fine coal, coal dust. and dirt.
- cyclone separator essentially a settling chamber to separate solid panicles from a gas. in which gravitational acceleration is replaced by centrifugal acceleration.
- degasification a process for removing  ${\tt nat}~{\tt urally}$  occurring methane from coal seams.
- delayed cokinq a process wherein coal is subjected to a long period of carbonization at moderate temperatures to form coke.
- demineralization removal of mineral matter (ash) from coal by solvent extraction. usually under hydrogen atmosphere.

- depolymerization the change of a large molecule into simpler molecules usually accompanied by the substitution of hydrogen for oxygen in the molecular structure.
- destructive distillation the distillation of coal accompanied by Its thermal decomposition.
- desulfurization the removal of sulfur from hydrocarbonaceous substances by chemical reactions.
- devolatization the removal of a portortion of the volatile matter from medium- and high-voiatile coals.
- diatomaceous earth a yellow. white. or light-gray, siliceous porous deposit made up of opaline shells of diatoms: used as a filter aid. paint filler, adsorbent. abrasive. and thermal insulator. Also known as kieselguhr.
- diatomite See Diatomaceous Earth.
- dissolution the taking up of a substance by a liquid with the formation of a homogeneous solution.
- distillation a process of vaporizing a liquid and condensing the vapor by cooling: used for separating liquids into various fractions according to their boiling points or boiling ranges.
- dotomite- a carbonate of calcium and magnesium having the chemical formula CaMg(CO3)2
- Dowtherm trademark for a series of eutectic mixtures of diphenyl oxide and dlphenyl used as high-temperature heat-transfer fluids.
- bullated bed gas containing a relatively small proportion of suspended solids. bubbles through a higher density fluidized phase with the result that the system takes on the appearance of "boiling liquid.
- economizer beat exchanging mechanism for recovering heat from flue gases.
- effluent gas gas given off from a process vessel.
- etutriation the preferential removal of the small constituents of a mixture of solid particles by a stream of high-velocity gas.
- ndoth\_mic reactton a process in which heat is absorbed.
- rtthdpy change the increase or decrease in heat content of a substance or system which accompanies its change from one state to another under constant pressure.
- ntrmncd bed (flow) a bed in which solid particles are suspended in a moving fluid and are continuously carried over in the effluent stream.
- eutectic that combination of two or more components which produces the lowest melting temperature.
- xotttermte reaction a process in which heat is liberated.
- extraction a method of separation in which a solid or solution is contacted with a liquid solvent (the two being essentially mutually Insoluble) to transfer components into the solvent.
- xtractive coking similar to delayed coking process. with the emphasis on high tar yields to produce liquids.
- filter and finely divided solids used to increase efficiency of filtering.
- filter cake -- the moist residue remaining from the filtration of a slurry to produce a clean filtrate.
- filtrate a liquid free of solid matter after having passed through a filter.
- filtration the separation of solids from liquids by passing the mixture through a suitable medium. e.g.. cloth, paper. diatonaccous earth.
- Fischer assay method for determining the tar and light oil yields from coal or oil shale: conducted in a retort under an inert atmosphere with a presribed increase in temperature to 500°C.
- Fischer-Tropsch catatyst catalysts developed for the catalytic synthesis of liquid fuels from coal-derived synthesis gas; catalysts contain principay iron. cobalt. nickel. or ruthenium.

Fischer-Tropsch process - method of hydrogenating mixtures of

carbon monoxide and hydrogen produced from coal. lignite. or natural gas by means of steam. at 1-10 atmospheres and 360-410°F to yield liquid and gaseous fuels. and a wide spectrum of industrial chemicals.

- fixed-bed stationary solid particles in intimate contact with fluid passing through them.
- fixed carbon the solid residue. other than ash. obtained by destructive distillation; determined by definite prescribed methods.
- flash carbonization a carbonization process characterized by short residence times of coal in the reactor to optimize tar yields.
- ftue gas- gaseous combustion products.
- fluidization (dense phase) the turbulent motion of solid particles in a fluid stream; the particles arc close enough as to interact and give the appearance of a boiling liquid.
- fluidization (entrained) gas-solid contacting process in which a bed of finely divided solid particles is lifted and agitated by a rising stream of gas.
- fluidized-bed assemblage of small solid panicles maintained in balanced suspension against gravity by the upward motion of a gas.
- fly ash— a fine ash from the pulverized urned in power station boilers. or entrained ash carried over from a gasifier.
- fractionation distillation process for the separation of the various components of liquid mixtures.
- freeboard the space in a fluidized-bed reactor between the top of the bed and the top of the reactor.
- free swelling index a standard test that indicates the caking characteristics of coal when burned as a fuel.
- Friedel-Crafts reaction a substitution reaction. catalyzed by aluminum chloride in which an alkyl R-) or acyl (RCO-) group replaces a hydrogen atom of an aromatic nucleus to produce a hydrocarbon or a ketone.
- fuel cell a galvanic cell in which the chemical energy of a conventional fuel is utilized to produce electricity.
- fuel gas low heating value (150-350 BTU, scf) product generally utilized on sitc for power generation or industrial use.
- gasification of coal the conversion of solid coal into a gaseous form by various chemical reactions with steam.
- gasifier a vessel in which gasification occurs. usually utilizing fluidized-bed. fixed-bed. or entrained-bed units.
- he@ capacity— quantity of heat required to raise the temperature of one pound of a substance one degree Fahrenheit.
- high-Btu gas— a gas having a heating value of 900 to 1,000 Btu per standard cubic foot. which approaches the value for natural gas.
- higher-heating value  $(HHV) th_{-}$  heat liberated during a combustion process in which the product water vapor is condensed to a liquid and the heat of condensation is recovered.
- hydroclone a small cyclone extractor for removal of suspended solids from a flowing liquid by means of the centrifugal form set up when the liquid is made to flow through a tight conical vortex.
- hydrocoking coking of tars. SRC. etc.. under hydrogenating conditions to form liquid products.
- hydrocracking the combination of cracking and hydrogenation of organic compounds.
- hydrogasification gasification that involves the direct reaction of fuels with hydrogen 10 optimize formation of methane.
- hydrogenation chemical reactions involving the addition of gaseous hydrogen to a substance in the presence of a catalyst under high temperatures and pressures.
- hydrogen donor solvent solvent. such as anthracene oil tetralin (tetrahydronaphthalene), decalin. etc.. which transfers hydro-

gen to coal constituents causing depolymerization and consequent conversion to liquid products of lower boiling range which are then dissolved by the solvent.

- hydrotreating a process to catalytically stabilize petroleum *or* other liquid hydrocarbon products and or remove objectionable elements from products or feedstocks by reacting them with hydrogen.
- Ideal gas— any gas whose equation of state is expressed by the ideal gas law, namely PV = nRT where P is the pressure. V is the volume. R is the gas constant. T is the absolute temperature, and n = number of moles.
- Ignition temperature— the minimum temperature necessary to initiate self-sustained combustion of a substance.

Industrial gas - see fuel gas.

 $\mbox{Inerts----}$  constituents of a coal which decrease its efficiency in  $\mbox{usc.}$ 

- e.g.. mineral matter (ash) and moisture in fuel for combustion. In situ— in its original place. wengerground gasification coal scam.
- intermediate-Btu synthesis gas product with a higher heating value between 350 and 500 Btu per standard cubic foot.
- Ilgnite— brownish-black coal containing 65-72 percent carbon on a mineral-matter-free basis. with a rank between peat and subbituminous coal.
- limestone— sedimentary rock containing 50 percent carbonate (CO3) of lime or magnesia. Chemical formula (for calcite limestone) is CaCO3
- liquefaction— conversion of a solid to a liquid: with coal. this appears to involve the thermal fracture of carbon-carbon and carbon-oxygen bonds, forming free radicals. These radicals abstract hydrogen atoms yielding low molecular weight gaseous and condensed aromatic liquids.
- liquefied patroleum gas (LPG)— those hydrocarbons that have a vapor pressure (at 70°F) slightly above atmospheric (such as propane and butane): kept in liquid form under a pressure higher than 1 atm.
- lock hopper a mechanical device that permits the introduction of a solid into an environment of different pressure.
- low-Btu gas a gas having a heating value up to 350 Btu per standard cubic foot.
- lower heating value the heat liberated by a combustion process assuming that none of the water vapor resulting from the process is condensed, so that its latent heat is not available.
- MAF- moisure and ash-free; a term that relates to the organic fraction in coal.
- mesh measure of fineness of a screen. e.g.. a MO-mesh sieve has 400 openings per linear inch.
- methanation the production of methane (CH4) from carbon monoxide or dioxide and hydrogen.
- methane a CH, a colorless. odorless. and tasteless gas. lighter than air; the chief component of natural gas.
- methane methanol alcohol. CH<sub>3</sub>OH.
- micron a unit of length equal to one millionth of a meter:  $10^{\circ}$  meter.
- moving bed particlized solids in a process vessel that are circulated (moved) either mechanically or by gravity flow.
- natural gas naturally occurring gas extracted from sedimental structures consisting mainly of methane and having a higher heating value of approximately 1,050 Btu per standard cubic foot.
- noncoking coal a coal that does not form coke under normal coking conditions.

- olefinic hydrocarbon a class of unsaturated hydrocarbons containing one or more double bonds and having the general chemical formula CnH2n
- open cycle a thermodynamic power cycee in which the working fluid passes through the system only once and is then exhausted to the atmosphere.
- peat— an unconsolidated. hydrophilic. yellowish-brown to brownish-black. carbonaceous sediment. formed by accumulation of partially fragmented and decomposed plant remains in swamps and marshes which retains more than 75 percent inherent moisture and less than 12 percent mineral matter in saturated natural deposits.
- petrochemicals— those derived from crude oil or natural gas. or their coal-derived substitutes: they include light hydrocarbons such as butylene. ethylene and propylene. the raw materials for the production of plastics by polymerization.
- of phenols— a group of aromatic compounds having the hydroxyl ( OH ) group directly attached to the benzene ring.
  - pilot plant chemical process plant containing all the processes of a commercial unit, but on a smaller scale. for the purpose of studying the technical and economic feasibility of the process.
  - pipeline gas— a methane-rich gas that conforms to certain standards and has a higher heating value between 950 and 1.050 Btu per standard cubic foot.
  - plenum chamber an enclosed space through which air is forced for slow distribution through ducts.
  - present layer of suitable filtering medium. e.g.. diatomaceous earth. laid down on a rotary filter cloth prior to operation.
  - prilling tower— a tower that produces small solid agglomerates by spraying a liquid solution in the top and blowing air up from the bottom.
  - process development unit a system used to study the effects of process variables on performance: sized between a bench-scale unit and a pilot plant.
  - proximate analysis— analysis of coal based on the percentages 01 moisture, volatile matter. fixed carbon (by difference). and asn. using prescribed methods. Reported on different bases. such as as-received (or as-fired), dry, mlncral-matter-free (mmf), and dry mineral- matter-free (dmmf).
  - purification removal of a wide range of impurities present in gases from coal gasification.
  - pyrolysis thermal decomposition of organic compounds in the absence of oxygen.
  - quenching cooling by immersion in oil. water bath. or water spray.
  - Raney nickel <sub>catalyst</sub> specially prepared nickel catalyst used in the hydrogenation of organic materals and the methanation of synthesis gas to methane.
  - raw gas— see crude gas.
  - reactivity susceptibility to chemical change: for example, in coal liquefaction. the reactivity of the coal for conversion to liquid products is a function of the coal rank. among other things.
  - reactor vessel in which coal-conversion reactions take place.
  - Rectisol process— a process for the purification of coal-gasification gas based on the capability of cold methanol to absorb all gas impurities in a single step: gas naphtha. unsaturated hydrocarbons. sulfur compounds. hydrogen cyanide. and carbon dioxide are removed from the gas stream by the methanol at temperatures below O°C.
  - reducing gas a gas which, at high temperatures, lowers the state of oxidation of other chemicals.

reforming processes — a group of proprietary processes in which low-grade or low molecular weight hydrocarbons are catalytically converted to higher grade or higher molecular weight materials: also applies to the endothemic reforming of methane. for the production of hydrogen, by the reaction of methane and steam in the presence of nickel catalysts.

refractory — a material capable of withstanding extremely high temperatures and having a relatively low thermal conductivity)'.

- residence time time spent by a typical particle in a particular zone.
- saturated hydrocarbon a carbon-hydrogen compound with all carbon bonds filled; that is. there are no double or triple bonds as in olefins and acetylenes.
- scrubber apparatus in which a gas stream is freed of tar. ammonia. and hydrogen sulfide.
- seam coal coal which is intermediate in rank between bituminous coal and anthracite: contains 8 to 22 percent volatile matter and from 9 I to 93 percent carbon.
- semi- water gas a mixture of carbon monoxide. carbon dioxide. hydrogen. and nitrogen. obtained by passing an air-stream mixture through a hot bed of coke. having a higher heating value of about 120 Btu per standard cubic foot.
- sensible heat— that heat which results in only the elevation of the temperature of a substance with no phase changes.
- shift conversin process for the production of gas with a desired carbon monoxide content from crude gases derived from coal gasification; carbon monoxide-rich gas is saturated with steam and passed through a Catalytic reactor where the carbon monoxide reacts with steam to produce hydrogen and carbon dioxide. the latter being subsequently removed in a scrubber employing a suitable sorbent.
- sintering the agglomeration of solids at temperatures below their melting point, usually as a consequence of heat and pressure.
- slag molten coal ash composed primarily of silica, alumina iron oxides. and calcium and magnesium oxides.
- slurry a suspension of pulverized solid in a liquid.
- solvation the assocation or combination of molecules of solvent with solute ions or molecules.
- solvent that component of a solution which is present in excess: liquid used to dissolve a substance.
- solvent extraction selective solution of coal constituents from finely divided coal particles into a suitable solvent after intimate mixing, usually at high temperatures and pressures in the presence of hydrogen, with or without a catalyst. followed by phase separation.
- solvent refined coal (SRC) a coal extract derived by solvent extraction; a brittle. vitreous solid (m.p. 300° F to 400° F) containing about 0.1 percent ash and about 10 percent of the sulfur in the original coal feedstock; calorific value is about 16.000 Btu per pound: may be used as a clean fuel for power generation by combustion: utilized for the production of h:gh-grade metallurgical coke. anode carbon. and activated carbon by coking. or hydrogenated to produce synthetic crude oil.
- space Velocity volume of a gas (measured at standard temperature and pressure) or liquid passing through a given volume of catalyst in a unit time.
- spacific gravity ratio of the weight of any volume of a substance to the weight of an equal volume of water at  $4^{\circ}C$ .
- specific heat heat capacity of a substance as compared with the heat capacity of an equal weight of water.
- standard cubic fool (SCF) the volume of a gas at standard conditions of temperature and pressure. The American Gas Associa-

tion uses moisture-free gas at 60° F and 30 inches of mercury (i.0037 atm) as Its standard conditions. The pressure standard is not universal in the gas industry: 14.7 psia (1.000 atm) and 14.4 psia (0.980 atm) are also used. The scientific community uses  $32^{\circ}F$  and ! atm as standard conditions.

- stoichometry the definite proportions in which molecules react chemically to form new molecules.
- stripping the removal of the more volatile components from a liquid mixture of compounds.
- subbituminous coal the rank of coal between bituminous and lignite. classified by ASTM as having a range of heating values between 8.300 and i 1.000 Btu per pound on a moist mineral-matter-free basis.
- **substitute** natural gas (SNG) a gas produced from coal. oil sands. or oil shale conforming to natural gas standards.
- Superficial velocity the linear velocity of a fluid flowing through a bed of solid particles calculated as though the particles were not present.
- superheater a heat exchanger ,which adds heat to the saturated steam leaving a boiler.
- syncrude synthetic crude oil: oil produced by the hydrogenation of coal.. coal extracts. oil sands. or oil shale. which is similar to petroleum crude.
- synthesia gas a mixture of hydrogen and carbon monoxide which can be reacted to yield a hydrocarbon.
- tail gas a gas issuing from a gas-treatment unit which may be recycled to the process or exhausted.
- tar (coal) a dark brown or black. viscous, combustible liquid formed by the destructive distillation of coai.
- therm a unit of heat used as a basis for the sale of natural gas; equal to 100.000 Btu.
- topping cycle the higher temperature thermodynamic power cycle of a combined-cycle system.
- turndown ratio- the minimum ratio of actual flow rate to design flowrate at which a process unit can be operated.
- ultimate analysis the determination by prescribed method of carbon and hydrogcn in the material as found in the gaseous products of its complete combustion. the determination of sulfur. nitrogen. and ash in the material as a whole and the estimation of oxygen by difference: may be reported on different bases. such as as-received (or as-fired). dry.. mineral-matterfree (mmf). and dry mInerai-matter-free (dmmf).
- Venturi scrubber— a gas cleaning device which involves the injection of water into a stream of dust-laden gas flowing at a high velocity through a contracted portion of a duct. thus transferring the dust particles to the water droplets which are subsequently removed.
- volatile matter those constituents of coal, exclusive of moisture. that are liberated from a sample when heated to 1750° F for seven minutes in the absence of oxygen.
- water gas— gas produced by the reaction of carbon (in coal or coke) and steam to yield mixtures of carbon monoxide and hydrogen: similar to synthesis gas.
- water gas shift the reaction between water vapor and carbon monoxide to produce hydrogen and carbon dioxide or the reverse: CO + H.O ≓ H: + CO<sub>2</sub>.
- working fluid a gas stream which directly does work. e.g.. powering a gas turbine.

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