Glycerol to Propylene Glycol

April 12, 2011

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April 12, 2011

Dear Professor Seider and Professor Fabiano,

As proposed by Bruce Vrana, we have designed a plant that will produce one hundred million pounds per year of propylene glycol from crude glycerol, which is a byproduct derivative of biodiesel production. This process integrates electrodeionization, an emerging glycerine refinement technology, with a multiphase reaction in a trickle-bed reactor.

This report includes the design details for the process, an economic analysis, and recommendations and predictions for the extent of its success. Based on the aforementioned analysis, this process is found to be a viable and profitable alternative to current methods of propylene glycol production. Furthermore, the process is environmentally friendly while also complementing the growth of the biodiesel industry that is predicted to surge over the project's lifetime.

Thank you for your consideration and assistance with this endeavor.

Sincerely,

Kelsey Hall

Kumardipti Chatterjee

Samuel Tell

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ABSTRACT

Abstract

A Cu-ZnO-Al₂O₃ catalyst has been observed in laboratory scale tests to effectively produce propylene glycol from glycerol using a liquid phase hydrogenolysis reaction, which occurs at 410 F and 580 psia. A trickle-bed reactor will be used to ensure the full contact of liquid and vapor phases with the solid catalyst. This project aims to successfully scale up this reactor model, which has thus far only been tested in bench scale. The design specification stipulates that this process will produce 100 MM lb/year of propylene glycol. Using crude glycerol harvested from biodiesel production, a final product purity of 99.6% was achieved from a feedstock of 80% glycerol, 15% water, 1% methanol, and 4% sodium chloride by weight, plus trace amounts of organic salts.

The economic analysis that follows assumes a grassroots plant on the US Gulf Coast. The total capital investment was calculated to be \$34.0 million, which includes a working capital of \$9.78 million. Under the assumptions that the prices of crude glycerol, hydrogen, and propylene glycol are \$0.22, \$0.50, and \$1.00 per pound respectively, the net present value (NPV) at the end of the 15 year allotted course of the project is \$88.4 million and the investors' rate of return (IRR) is 58.45%. The price of glycerol is projected to remain stable or decrease in the future and the price of propylene glycol is projected to remain stable or increase suggesting that this project could become even more profitable in the future.

The apparent profitability of this project is largely caused by the efficient and cost effective method of desalting glycerol through electrodeionization. Unfortunately, the proprietary nature of this new process precludes public access to true costing and specifications for the equipment since firms as Dow largely control the technology. Thus, conservative estimations were made in our economic analysis to account for this uncertainty.

INTRODUCTION

Introduction

Propylene glycol (also called 1,2-propanediol) is an incredibly versatile compound that is used in a number of industrial applications that range from transportation and construction to food and pharmaceutical production. Pharmaceutical (USP) grade propylene glycol is at least 99.5% pure by weight and is used in healthsensitive products such as food, personal consumer goods, cosmetics, and pharmaceuticals. Due to its highly sensitive applications, USP grade propylene glycol is regulated carefully by the FDA and producers must comply with strict regulations to ensure the quality and purity of their product. Industrial grade propylene glycol is at least 95% pure and is an important player in the transportation industry as it is used in aircraft de-icer, antifreeze, and brake fluid. It is also used in the construction industry as the primary component in unsaturated polyester resins (UPRs) that are used to make fiberglass reinforced plastics.

Propylene glycol's chemical neutrality and nonreactivity make it very useful as a solvent. It can be used as an emulsifier to stabilize mixtures of two or more immiscible liquids. This often occurs in the preparation of cosmetics, where oil and water must be mixed to produce creams or lotions, and in the preparation and processing of some foods. It is a useful excipient, a pharmacologically inactive substance that acts as a carrier for the active ingredients in medication. It can be used for boiling point elevation or freezing point reduction, which makes it an effective de-icer and antifreeze solution.

Historically, propylene glycol has been produced by the hydration of propylene oxide, which occurs at 392 F and 174 psia, or catalytically at 302-356 F, and produces di-

and tripropylene glycols and small quantities of higher glycols in side reactions (ICIS, 2011). This process has a large negative environmental impact due to pollution and use of valuable resources, so alternative methods, such as production from glycerol, are under investigation worldwide. This option is especially promising in light of the recent surge in biodiesel production. Interest in "greener" technologies has in part been responsible for this boom, creating incentive for companies to invest in research to develop more environmentally friendly methods to produce and distribute their products. In addition to being extremely profitable, the glycerol to propylene glycol manufacturing process described in this report can be marketed as "green" technology, increasing the value of the end product.

The method for propylene glycol production described here consists of three main sections: a pretreatment section, in which crude glycerol purchased from biodiesel manufacturers is desalted by electrodeionization for feed to the reactor; a reactor section, which contains a trickle-bed reactor packed with alumina-supported catalyst for the reaction of glycerol; and a separation section, in which the product is purified to 99.5% (USP grade). Overall, this process describes a profitable, novel way to produce propylene glycol while reducing the negative environmental effects of traditional production methods. It is predicted to eventually become a mainstream process in industry.

Section 3 PROJECT CHARTER

Initial Project Charter

| Project Name | Glycerol to Renewable Propylene Glycol | | |
|-------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--|--|
| Project Champions | Bruce M. Vrana (DuPont) Dr. Warren D. Seider (UPenn) Professor Leonard Fabiano (UPenn) | | |
| Project Leaders | Kelsey Hall Kumar Chatterjee Samuel Tell | | |
| Specific Goals | Develop a process to make 100MM lb/year of propylene glycol from crude glycerol that provides a high Net Present Value and Return On Investment | | |
| Project Scope | In Scope: | | |
| | Selecting an optimal catalyst for the process Designing a trickle-bed reactor model based on reaction kinetics Investigating and choosing an efficient and cost-effective pretreatment equipment for crude glycerol Creating a process flow sheet for the overall process Conducting a financial analysis of the process to determine the ROI of the project Writing a report that summarizes all relevant information | | |
| | Out of Scope: | | |
| | Testing and verifying reaction kinetics, conversions and yields proposed in literature | | |
| Deliverables | Business Opportunity Assessment Technical Feasibility Assessment Manufacturing Capability Assessment Safety and Health Assessment Profitability Analysis | | |
| Timeline | The deliverables included above will be completed in 10 weeks. Future implementation of the project proposal will be determined by the Project Champions. | | |

TECHNOLOGY READINESS ASSESSMENT

Technology Readiness Assessment

One major limiting factor for the synthesis of propylene glycol from glycerol is the extremely high cost and difficulty of glycerol desalting. In the past, vacuum distillation was the only option and was very expensive, precluding the entry of many proprietors into the market. There are a number of companies that specialize in glycerine desalting, but the purified feed glycerol is expensive and drives up the cost of the process. As membrane and resin technologies improve, the electrodeionization process becomes a low cost alternative to vacuum distillation, which allows the use of crude glycerol in the process feed, making the process much more economically feasible. The popularity of biodiesel has increased in recent years, increasing the availability and decreasing the price of the crude glycerol process input.

Although the kinetic model for the glycerol to propylene glycol reaction used in this process is fairly new, the necessary Cu-ZnO-Al₂O₃ catalyst is available commercially. Alumina-supported catalysts such as this are standard for use in many industrial applications. A catalyst very similar to the one in this process is widely used for methanol synthesis and other applications. It is manufactured and sold by a number of companies worldwide, such as Haldor Topsoe (MK-121 catalyst).

Slight manipulation must be done to prepare the catalyst for use in the propylene glycol manufacturing process, as the reaction kinetics call for the catalyst particles to have a diameter less than 0.22 mm. Since the commercial methanol synthesis catalysts are sold in larger (6x5 mm) pellets, simple manual crushing of the catalyst to achieve the desired size will suffice.

The Cu-ZnO-Al₂O₃ catalyst with a molar metal ratio of 1:1:0.5 was chosen out of a group of similar options with slight variations (Zhou 2010). In activity tests, this showed the highest conversion and selectivity to propylene glycol. It is assumed in the process analysis that a 100% conversion may be reached by extrapolating the mass of catalyst to a large enough value based on the expressions for r_1 and r_2 . In reality, this is an optimistic estimate, and the actual conversion will be less than 100%. It is expected that the actual conversion will fall between the 82% achieved in Zhou's activity test and the 100% predicted with the extrapolation model (Appendix B). To account for this, further separation may be necessary following the reactor section in the process.

Prior to beginning plant construction, a midsize pilot plant should be used to determine the reliability of scale-up models used for the reactor. This will also provide a more precise value for the conversion of glycerol in the reactor. The final separation train should not be designed until the reactor product composition can be determined.

CONCEPT STAGE

Market Analysis

In the following sections, the various inputs and products of this process are analyzed in the context of their individual markets to determine consumer, technical and financial needs. Propylene glycol historically has been synthesized from a petroleum derivate known as propylene oxide. In recent years, novel reaction routes have been discovered to synthesize propylene glycol from glycerol. In a patent issued to the BASF chemical company, a Cu-ZnO-Al₂O₃ catalyst was identified that converts glycerol to propylene glycol with very high conversion. Crude glycerol, a naturally occurring by-product of biodiesel production, will be used as a feed to this process.

Crude Glycerol Market

For every nine kilograms of biodiesel produced, about one kilogram of glycerol is formed; thus, the market for crude glycerol is intimately linked with the biodiesel industry (Chiu *et al.* 2006). Biodiesel represents a very promising green alternative as it can be used in conventional diesel engines. Furthermore, the synthesis mechanism for biodiesel is well studied and can be carried out using a variety of feed stocks. As of 2009, 148 plants were producing 2 billion gallons of biodiesel a year in the US. As a result, 200 million gallons of glycerol was produced as well (Johnson *et al.* 2007). Although purified glycerol can be sold for \$0.60/lb-\$1.20/lb depending on the grade, crude glycerol is currently sold for \$0.22/lb, and the price is projected to fall as biodiesel production continues to increase.

Propylene glycol represents an inherently profitable value-added chemical produced from crude glycerol due to the high margin between the feed and final product. In addition to propylene glycol, however, a number of other alternatives also exist as potential markets for

crude glycerol as outlined in the article "Glycerin – Emerging Opportunity". Solvay Chemicals commercialized a process to reverse the conventional route of making synthetic glycerol from epichlorohydrin. After the success of their pilot plant in 2007, a full scale commercial plant is scheduled to be built in Map Tha Phut, Thailand. Epichlorohydrin can be used to make epoxy resins, paper-reinforcing agents and other similar products. Alkylene carbonate (reactive intermediate or solvent), biomenthanol, acrylic acid are additional examples of potential products produced from glycerol feed material. (Chemical Business 2010).

Companies have also invested in developing cost effective methods to purify crude glycerol. Pure glycerol has a number of uses ranging from consumer goods to food and beverages to industrial chemicals. Pharmaceutical grade glycerol can be sold for up to \$1.20/lb, making it a very profitable venture. Purada Processing and Cargill are two examples of companies that have chosen to invest in glycerol refineries. However, traditional methods of crude glycerol refinement require vacuum distillation and thus are capital and energy intensive. New purification systems, such as electrodionization (EDI), offer good alternatives to costly and labor intensive vacuum distillation, which could further equalize glycerol supply and demand.

Propylene Glycol

Propylene glycol is a value-added derivative of crude glycerol that has the potential to be a profitable venture. PG is used in a number of markets including unsaturated polyester resins, plastics, antifreeze products, plane de-iceres, industrial solvents, and consumer goods such as detergents, cosmetics and other personal care products. The global demand for propylene glycol was 3.5 billion lb/yr as of 2007 (Shelley 2007). According to a recent report by ICIS, a subsidiary of Reed Business Information and trusted information provider for the chemical and oil industry, the demand in the unsaturated polyester resin (UPR) industry has grown

approximately 2.5% per year while demand in the cosmetics and consumer goods industries has grown 3.0% and 3.5% a year, respectively (2010). Because propylene glycol is not toxic, it is gradually replacing ethylene glycol as the primary ingredient in the 2.4 billion lb/yr market of antifreeze de-icing liquids (Suppes 2008).

Although the emergence of the renewable propylene glycol market is relatively new, a number of large industry leaders have already begun to embrace the change. Dow Chemical, currently the world's leading manufacturer of petroleum based propylene glycol, introduced product called renewable propylene glycol (RPG). A successful pilot plant was built at their Dow Halterman Custom Processing Facility in Houston and a full scale plant will soon be implemented at that site. Huntsman Corp. is another major company that has begun producing RPG using a proprietary method. Located in Conroe Texas, the company started to scale up the process to produce 100 million lb/yr (Shelley 2007).

Senergy Chemical famously licensed a novel low temperature, low pressure reaction route developed by two of the most famous researchers in the field, Dr. Suppes and Dr. Sutterlin from the University of Missouri. And finally, Ashland and Cargill have signed a joint venture to build a facility to produce 65,000 tonnes per year of RPG from glycerol. Furthermore, the plant will include a large glycerol refinery that will enable it to simultaneously produce high purity glycerol to be sold in the chemical market. (Chemical Business 2010)

Consumer Requirements

| Consumer Requirement Category | Propylene Glycol Requirement | Type (NUD,FTS) | Weighting Factor |
|----------------------------------|--------------------------------------|-------------------|---------------------|
| Non-Toxic | Non-toxic to both people and animals | FTS | 35 |
| High Purity | Non-toxic for food/cosmetics | NUD | 35 |
| Low Freezing Point | Appropriate for anti-freeze | FTS | 12 |
| Miscibility | Use in detergents | FTS | 6 |
| Polar Solvent | Use in paints and detergents | FTS | 6 |
| Green Manufacturing | Biodiesel Byproduct | NUD | 3 |
| Degradability | Use in detergents and solvents | FTS | 2 |
| Ease of Handling | Use as a working fluid | FTS | 1 |

Table 1 Consumer requirements.

The consumer requirements of non-toxicity and high purity were weighted as the most important factors because the food, consumer goods and pharmaceutical industries have been identified as the most profitable target markets. As a result, our process is designed to produce UPS-grade propylene glycol, which is defined as greater than 99.8% purity by mass. The high purity requirement is labeled as New-Unique-Difficult because it relies on new catalysts that have been developed for very high selectivity. Furthermore, these criteria are important when positioning propylene glycol in the antifreeze and de-icing markets as an alternative to the toxic ethylene glycol.

Our process is also designed to accommodate consumers in the industrial chemical market. Low freezing point, miscibility and polarity were all weighted to account for their needs. Finally, a weighting factor of 3 was assigned to "Green Manufacturing" because there is

significant marketing potential if the product is characterized as an environmentally friendly chemical. Furthermore, designing a plant with a limited carbon footprint may be advantageous when seeking government subsidies or tax breaks in the future.

PRELIMINARY PROCESS SYNTHESIS

Pretreatment Section

The growing biodiesel industry supplies an abundance of crude glycerol products that can be used as the feed for the propylene glycol synthesis process. However, this crude glycerol cannot be fed directly to the process because it contains chloride and sulfate salts and organic alcohols, which must be removed prior to reaction. If they were to enter the reactor, these salts would cause corrosion of the reactor and preclude the success of the process.

Conventional methods for accomplishing this difficult purification require a great deal of energy and resources. A number of companies, such as SRS Engineering Corporation of Murrieta, California (SRS 2010), turn a considerable profit by refining and selling purified glycerol. The conventional method of desalting glycerol involves its distillation under vacuum conditions. Glycerol is extremely stable and nonvolatile, with a normal boiling point of 554 F. Because the polymerization temperature of glycerol (approximately 500 F) is lower than its boiling point, distillation that requires glycerol evaporation is impossible to achieve at atmospheric pressure. In order to maintain the condenser's temperature below the polymerization temperature of glycerol flows out of the column in an overhead stream through the condenser, leaving a highly concentrated salt solution with other heavy components in the bottoms product from the reboiler.

The high temperatures required to achieve this separation result in excessive utility usage in the form of high pressure steam and electricity. The vacuum inside the column necessitates thick walls and effective seals to prevent leakage of gases into the column and a powerful pump

to maintain the vacuum. These things are expensive and contribute to the high price of glycerol purification.

Recently, new technologies involving cation/anion selective membranes have been developed. These allow for electricity to be used instead of heat as the separation agent for removal of salts from glycerol. One such technique is electrodeionization, which involves flowing the mixture of glycerol, water, and salt through a mixture of resins bound together with fluoroelastomers. The resins adsorb the free ions, which are then attracted to opposite sides of the EDI device when an electrical current is applied. The ions are transported through an ionically selective semi-permeable membrane and accumulate in a concentrate stream of process water. The concentrate is then disposed of as wastewater.

EDI is a proprietary process, and there are only a few companies that specialize in desalting glycerol through electrodeionization, such as EET and Dow. Due to the competitive nature of this new market, process specifications for EDI equipment are not publicly available. However, from some basic figures provided by EET, it was estimated that the desalting of the crude glycerol stream in this process to 10 ppm salt content should require approximately 200 kW of electricity. This estimation was made using the current efficiency equation (Shaffer 1980).

$$\xi = \frac{zFQ_f(C_{inlet}^d - C_{outlet}^d)}{NI}$$

Details of this calculation are located in the Appendix. Additionally, an estimated \$2MM/year was assumed for equipment leasing. Both the capital and annual costs of using the electrodeionization process to desalt the crude glycerol is substantially lower than the cost of vacuum distillation.

Reactor Section

A trickle-bed reactor was selected for use in this process to accommodate the multiphase reaction. Non-negligible volumes of liquid and vapor phase reactants must be able to simultaneously come into contact with the solid surface of the catalyst in order for the reaction to proceed. In a trickle bed reactor, a liquid feed and a vapor feed enter the reactor co-currently at the top of the packed catalyst bed. The liquid "trickles" down over the catalyst and reaction occurs as described by the kinetic model. A more traditional reactor such as a packed bed plug flow reactor is unacceptable for this application due to the impracticality of sufficient contact between the hydrogen gas and the catalyst in the presence of liquid. This effect is amplified by the presence of a large excess of hydrogen gas in the reactor, which is intended to ensure complete conversion of the glycerol input.

Trickle bed reactors are traditionally used in industry for hydrodesulfurization, the kinetics and conditions of which are similar to propylene glycol synthesis from glycerol. Both reactions occur at severe conditions (temperatures above 572 F and pressures above 514 psia) and require a vapor and a liquid to react on a solid catalyst. Using the catalytic hydrodesulfurization of thiophene as described in Chapter Five of *Chemistry of Catalytic Processes* by Gates, Katzer, and Schuit as a basis for comparison, it is expected that the use of an industrial trickle-bed reactor will be successful for propylene glycol manufacture in this process.
CHEMICAL DATABASE

Chemical Database

The notable thermophysical and chemical properties of each chemical species present in the process are shown below. Room temperature values are displayed for all temperature-dependent properties (viscosity, thermal conductivity, heat capacity).

| Propylene Glycol (| (1,2 propanediol) |
|-------------------------------|-------------------|
| Molecular Weight (g/mol) | 76.09 |
| Melting Point Temp (°C) | -60 |
| Boiling Point Temp (°C) | 189 |
| Density (lb/ft ³) | 64.207 |
| Standard enthalpy of | -209880 |
| formation (Btu/lbmol) | |
| Standard molar entropy | -130.306 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 40.4 |
| Thermal Conductivity | 0.116 |
| $(Btu-ft/hr-ft^2-R)$ | |
| Heat Capacity, CP | 38.735 |
| (Btu/lbmol-R) | |

| Glyce | erol |
|-------------------------------|---------|
| Molecular Weight (g/mol) | 92.1 |
| Melting Point Temp (°F) | 64.2 |
| Boiling Point Temp (°F) | 554 |
| Density (lb/ft ³) | 78.7 |
| Standard enthalpy of | -287069 |
| formation (Btu/lbmol) | |
| Standard molar entropy | -146.3 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 934 |
| Thermal Conductivity | 0.169 |
| $(Btu-ft/hr-ft^2-R)$ | |
| Heat Capacity, CP | 46.327 |
| (Btu/lbmol-R) | |

| Wat | ær |
|-------------------------------|--------|
| Molecular Weight (g/mol) | 18.0 |
| Melting Point Temp (°F) | 32 |
| Boiling Point Temp (°C) | 212 |
| Density (lb/ft ³) | 62.4 |
| Standard enthalpy of | -229.2 |
| formation (Btu/lbmol) | |
| Standard molar entropy | 38.9 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 1 |
| Thermal Conductivity | 0.335 |
| $(Btu-ft/hr-ft^2-R)$ | |
| Heat Capacity, CP | 1.80 |
| (Btu/lbmol-R) | |

| Hydrogen | (gaseous) |
|-------------------------------|-----------|
| Molecular Weight (g/mol) | 2.01 |
| Melting Point Temp (°C) | -259 |
| Boiling Point Temp (°C) | -253 |
| Density (lb/ft ³) | 0.005 |
| Standard enthalpy of | 0 |
| formation (Btu/lbmol) | |
| Standard molar entropy | 22.430 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 0.009 |
| Thermal Conductivity | 0.103 |
| $(Btu-ft/hr-ft^2-R)$ | |
| Heat Capacity, CP | 6.873 |
| (Btu/lbmol-R) | |

| n-Propanol (1 | l-propanol) |
|--------------------------------------------------------|-------------|
| Molecular Weight (g/mol) | 60.1 |
| Melting Point Temp (°C) | -127 |
| Boiling Point Temp (°C) | 97 |
| Density (lb/ft ³) | 50.255 |
| Standard enthalpy of | -130079 |
| formation (Btu/lbmol) | |
| Standard molar entropy (Btu/lbmol-R) | -107.168 |
| (Dtu/I0III0I-R) | 1.052 |
| viscosity (cr) | 1.932 |
| Thermal Conductivity (Btu-ft/hr-ft ² -R) | 0.090 |
| Heat Capacity, C _P (Btu/lbmol-R) | 37.301 |

| Isopropanol (| 2-propanol) |
|--------------------------------|-------------|
| Molecular Weight (g/mol) | 60.1 |
| Melting Point Temp (°C) | -89 |
| Boiling Point Temp (°C) | 82.5 |
| Density (lb/ft ³) | 49.458 |
| Standard enthalpy of | -136421 |
| formation (Btu/lbmol) | |
| Standard molar entropy | -109.637 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 2.055 |
| Thermal Conductivity | 0.078 |
| (Btu-ft/hr-ft ² -R) | |
| Heat Capacity, CP | 41.309 |
| (Btu/lbmol-R) | |

| | 1 |
|--------------------------------|------------|
| Acet | tol |
| Molecular Weight (g/mol) | 74.1 |
| Melting Point Temp (°C) | -17 |
| Boiling Point Temp (°C) | 145 |
| Density (lb/ft ³) | 66.939 |
| Standard enthalpy of | -178054 |
| formation (Btu/lbmol) | |
| Standard molar entropy | -92.820 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 6.701 |
| Thermal Conductivity | 0.088 |
| (Btu-ft/hr-ft ² -R) | |
| Heat Capacity, CP | 29.739 |
| (Btu/lbmol-R) | |

| Metha | anol |
|--------------------------------|---------|
| Molecular Weight (g/mol) | 32.0 |
| Melting Point Temp (°C) | -98 |
| Boiling Point Temp (°C) | 65 |
| Density (lb/ft ³) | 49.5 |
| Standard enthalpy of | -102573 |
| formation (Btu/lbmol) | |
| Standard molar entropy | -57.5 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 0.538 |
| Thermal Conductivity | 0.116 |
| (Btu-ft/hr-ft ² -R) | |
| Heat Capacity, CP | 24.6 |
| (Btu/lbmol-R) | |

| Ethylene | Glycol |
|-------------------------------|----------|
| Molecular Weight (g/mol) | 62.1 |
| Melting Point Temp (°C) | -12.9 |
| Boiling Point Temp (°C) | 197.3 |
| Density (lb/ft ³) | 69.865 |
| Standard enthalpy of | -197205 |
| formation (Btu/lbmol) | |
| Standard molar entropy | -107.699 |
| (Btu/lbmol-R) | |
| Viscosity (cP) | 16.746 |
| Thermal Conductivity | 0.147 |
| $(Btu-ft/hr-ft^2-R)$ | |
| Heat Capacity, CP | 34.608 |
| (Btu/lbmol-R) | |

The T-xy diagrams for the major separations in the process are shown below. As in column D101, the separation of glycerol and water can be described by the curves shown in Figure 1.



Figure 1 T-xy diagram for the separation of glycerol and water.

The major separation in column D102 is the extraction of propylene glycol from water. This separation, at a pressure of 20 psia as in the column, is described by the curves in Figure 2.



Figure 2 T-xy diagram for the separation of propylene glycol and water.

N-propanol and water exist simultaneously in the system. An azeotrope exists as shown in Figure 3, but it is of very little concern as the mole fraction of water in such mixtures is significantly higher than the point of azeotropy, which in this case is approximately 60% water. The azeotrope only becomes a concern at compositions near this value, which are not present in the process.



Figure 3 T-xy diagram for the separation of n-propanol and water. An azeotrope exists at a composition of approximately 60% water.

REACTION KINETICS

Reaction Kinetics

The kinetics of the reaction of glycerol to propylene glycol used in this process were developed largely in "Kinetics of Hydrogenolysis of Glycerol to Propylene Glycol over Cu-ZnO-Al₂O₃ Catalysts" (Zhou et al. 2010). Their model was developed to enable the simulation and optimization of different-scale reactors for glycerol hydrogenolysis, as used in the propylene glycol production process.

Cu-ZnO- Al₂O₃ catalysts of varying metal compositions were prepared using coprecipitation. Activity tests showed that a molar ratio of 1:1:0.5 Cu:Zn:Al in the catalyst resulted in the highest glycerol conversion as well as the highest selectivity for propylene glycol. Henceforth, this preparation was used in catalytic studies. The experiments were performed over a hydrogen pressure of 431-725 psia and a temperature range of 428-464 K.

A two-step mechanism for the hydrogenolysis of glycerol to propylene glycol, originally proposed by Suppes *et al.*, has been investigated by many researchers and is widely accepted for copper-based catalysts. This mechanism, shown below, consists of the dehydration of glycerol to intermediate acetol, followed by the hydrogenation of acetol to propylene glycol.



Figure 4 Reaction mechanism of glycerol hydrogenolysis to propylene glycol (Zhou et al. 2010)

From this reaction, a two-site Langmuir-Hinshelwood kinetic model was proposed as follows, and validated by the experimental work of Li, Zhou et al.:

$$H_{2} + 2\theta_{1} \underbrace{\xrightarrow{b_{1}}}{2H\theta_{1}} 2H\theta_{1}$$

$$G + \theta_{2} \underbrace{\xrightarrow{b_{0}}}{G\theta_{2}} G\theta_{2}$$

$$G\theta_{2} \underbrace{\xrightarrow{k_{1}}}{A\theta_{2}} A\theta_{2} + H_{2}O$$

$$2H\theta_{1} + A\theta_{2} \underbrace{\xrightarrow{k_{2}}}{P\theta_{2}} P\theta_{2} + 2\theta_{1}$$

$$A\theta_{2} \underbrace{\xrightarrow{b_{0}^{-1}}}{P} A + \theta_{2}$$

$$P\theta_{2} \underbrace{\xrightarrow{b_{0}^{-1}}}{P} P + \theta_{2}$$

Figure 5 Langmuir-Hinshelwood kinetics for glycerol dehydrogenation reaction.

Where θ_1 is the active site for hydrogen adsorption, θ_2 is the active site for organic molecule adsorption, k_1 and k_2 are the rate constants for the first and second reaction steps respectively, and b_H , b_G , b_A , b_p are the adsorption constants of each species. The temperature dependence of these parameters is expressed in the following equations.

$$k_i = k_i^0 \exp[\frac{-E_i}{R_g T}], \qquad i=1,2$$
$$b_j = b_j^0 \exp[\frac{Q_j}{R_g T}], \qquad j=G, A, P, H$$

Assuming that a pseudo-steady-state analysis for adsorbed intermediates is valid, the rate expressions for the system are derived and presented below, where r_1 is the consumption of glycerol per mass of catalyst and r_2 is the production of propylene glycol per mass of catalyst, c_i are the molar concentrations of each species and P_H is the hydrogen pressure.

$$r_{1} = \frac{k_{1}b_{G}c_{G}}{1 + b_{G}c_{G} + b_{A}c_{A} + b_{P}c_{P}}$$
$$r_{2} = \frac{k_{2}b_{A}c_{A}b_{H}P_{H}}{(1 + b_{G}c_{G} + b_{A}c_{A} + b_{P}c_{P})(1 + \sqrt{b_{H}P_{H}})^{2}}$$

Figure 6 Rate expressions for propylene glycol formation.

The kinetic model described above was used as a basis for the design of the trickle-bed reactor that appears in the glycerol to propylene glycol process. To determine the mass of catalyst needed, an Excel model was developed to compute the kinetic parameters and species flow rates based on the temperature, hydrogen to glycerol feed ratio, and the feed composition using a Riemann sum over the total catalyst mass. This calculation is shown in the Appendix. The conversion of glycerol is plotted against the mass of catalyst in kilograms in Figure 7. Based on this analysis, it was determined that 4190 kg of catalyst is necessary to achieve a 99.8% conversion of glycerol, for a hydrogen to glycerol feed ratio of 5:1, temperature of 423 F and the specified feed composition.



Figure 7 Conversion of glycerol versus mass of catalyst. 4190 kg of catalyst are necessary to achieve 99.8% conversion.

Higher conversions have a very low rate of return. To achieve the remaining 0.1% conversion would require almost 2000 kg of additional catalyst, or an increase of almost 150%, which is not economically justifiable.

PROCESS FLOW DIAGRAMS AND MATERIAL BALANCES



| | S-101 | S-102 | S-103 | S-104 | S-105 | S-106 | S-107 | S-108 | S-109 | S-110 | S-111 | S-112 | S-113 | S-114 |
|------------------|--------|--------|--------|-------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| | LIQUID | LIQUID | LIQUID | SOLID | LIQUID |
| Temperature F | 70.0 | 70.0 | 54.8 | 54.8 | 69.8 | 6.69 | 151.8 | 242.0 | 221.4 | 80.0 | 80.0 | 80.1 | 80.0 | 247.9 |
| Pressure psia | 30.0 | 30.0 | 25.0 | 20.0 | 20.0 | 42.0 | 41.8 | 36.8 | 20.0 | 19.7 | 19.7 | 30.0 | 19.7 | 20.0 |
| Total Flow lb/hr | 19392 | 5384 | 34007 | 5876 | 33231 | 33231 | 33230 | 33230 | 13985 | 13985 | 9230 | 9230 | 4755 | 19245 |
| Mass Flow | | | | | | | | | | | | | | |
| GLYCEROL | 15514 | 0 | 15514 | 0 | 15514 | 15514 | 15514 | 15514 | 0 | 0 | 0 | 0 | 0 | 15514 |
| WATER | 2909 | 5384 | 17151 | 5100 | 17151 | 17151 | 17149 | 17149 | 13421 | 13421 | 8858 | 8858 | 4563 | 3729 |
| HYDROGEN | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| ЪС | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| N-PROP | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| ISO-PROP | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| SODIU-01 | 776 | 0 | 776 | 776 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| METHA-01 | 194 | 0 | 566 | 0 | 566 | 566 | 566 | 566 | 564 | 564 | 372 | 372 | 192 | 2 |
| Mass Frac | | | | | | | | | | | | | | |
| GLYCEROL | 0.800 | 0.000 | 0.456 | 0.000 | 0.467 | 0.467 | 0.467 | 0.467 | Trace | Trace | Trace | Trace | Trace | 0.806 |
| WATER | 0.150 | 1.000 | 0.504 | 0.000 | 0.516 | 0.516 | 0.516 | 0.516 | 0.960 | 0.960 | 096.0 | 0.960 | 0.960 | 0.194 |
| HYDROGEN | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| Ъд | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| N-PROP | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| ISO-PROP | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| SODIU-01 | 0.040 | 0.000 | 0.023 | 1.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| METHA-01 | 0.010 | 0.000 | 0.017 | 0.000 | 0.017 | 0.017 | 0.017 | 0.017 | 0.040 | 0.040 | 0.040 | 0.040 | 0.040 | Trace |
| | | | | | | | | | | | | | | |



| | S-114 | S-115 | S-116 | S-117 | S-118 | S-119 | S-120 | S-121 | S-122 | S-123 | S-124 | S-125 | S-126 | S-127 | S-128 | S-129 | S-130 | S-131 |
|------------------|--------|--------|--------|--------|-------|-------|-------|-------|--------|-------|-------|--------|-------|-------|-------|--------|-------|-------|
| | LIQUID | LIQUID | LIQUID | VAPOR | VAPOR | MIXED | VAPOR | MIXED | LIQUID | VAPOR | MIXED | LIQUID | VAPOR | MIXED | MIXED | LIQUID | VAPOR | VAPOR |
| Temperature F | 247.9 | 252.1 | 410.0 | 70.0 | 11.7 | 202.4 | 416.5 | 455.8 | 455.2 | 455.2 | 429.0 | 429.0 | 429.0 | 409.0 | 225.0 | 225.0 | 225.0 | 244.4 |
| Pressure psia | 20.0 | 587.0 | 581.0 | 1000.0 | 600.0 | 600.0 | 598.9 | 566.0 | 560.0 | 560.0 | 559.1 | 559.1 | 559.1 | 558.3 | 553.3 | 553.3 | 553.3 | 600.0 |
| Total Flow lb/hr | 19245 | 19245 | 19245 | 346 | 346 | 1952 | 1952 | 21197 | 4317 | 16880 | 16880 | 4876 | 12004 | 12004 | 12004 | 10398 | 1606 | 1606 |
| Mass Flow | | | | | | | | | | | | | | | | | | |
| GLYCEROL | 15514 | 15514 | 15514 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| WATER | 3729 | 3729 | 3729 | 0 | 0 | 308 | 308 | 7117 | 491 | 6626 | 6626 | 759 | 5867 | 5867 | 5867 | 5559 | 308 | 308 |
| HYDROGEN | 0 | 0 | 0 | 346 | 346 | 1342 | 1342 | 667 | 1 | 966 | 966 | 1 | 966 | 966 | 966 | 0 | 995 | 995 |
| ЪG | 0 | 0 | 0 | 0 | 0 | 18 | 18 | 12645 | 3805 | 8840 | 8840 | 4088 | 4752 | 4752 | 4752 | 4734 | 18 | 18 |
| N-PROP | 0 | 0 | 0 | 0 | 0 | 60 | 60 | 136 | 6 | 127 | 127 | 12 | 116 | 116 | 116 | 55 | 60 | 60 |
| ISO-PROP | 0 | 0 | 0 | 0 | 0 | 224 | 224 | 300 | 12 | 288 | 288 | 16 | 272 | 272 | 272 | 48 | 224 | 224 |
| SODIU-01 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| METHA-01 | 2 | 2 | 2 | 0 | 0 | 0 | 0 | m | 0 | 2 | 2 | 0 | 2 | 2 | 2 | 2 | 0 | 0 |
| Mass Frac | | | | | | | | | | | | | | | | | | |
| GLYCEROL | 0.806 | 0.806 | 0.806 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| WATER | 0.194 | 0.194 | 0.194 | 0.000 | 0.000 | 0.158 | 0.158 | 0.336 | 0.114 | 0.393 | 0.393 | 0.156 | 0.489 | 0.489 | 0.489 | 0.535 | 0.192 | 0.192 |
| HYDROGEN | 0.000 | 0.000 | 0.000 | 1.000 | 1.000 | 0.687 | 0.687 | 0.047 | Trace | 0.059 | 0.059 | Trace | 0.083 | 0.083 | 0.083 | 0.000 | 0.620 | 0.620 |
| Ъд | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.009 | 0.009 | 0.597 | 0.881 | 0.524 | 0.524 | 0.838 | 0.396 | 0.396 | 0.396 | 0.455 | 0.011 | 0.011 |
| N-PROP | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.031 | 0.031 | 0.006 | 0.002 | 0.008 | 0.008 | 0.002 | 0.010 | 0.010 | 0.010 | 0.005 | 0.037 | 0.037 |
| ISO-PROP | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.115 | 0.115 | 0.014 | 0.003 | 0.017 | 0.017 | 0.003 | 0.023 | 0.023 | 0.023 | 0.005 | 0.139 | 0.139 |
| SODIU-01 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| METHA-01 | Trace | Trace | Trace | 0.000 | 0.000 | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace |
| | | | | | | | | | | | | | | | | | | |



| | S-129 | S-130 | S-131 | S-132 | S-133 | S-134 | S-135 | S-136 | S-137 | S-138 |
|------------------|--------|-------|-------|-------|-------|--------|--------|--------|-------|--------|
| | LIQUID | VAPOR | VAPOR | MIXED | MIXED | LIQUID | LIQUID | LIQUID | VAPOR | LIQUID |
| Temperature F | 225.0 | 225.0 | 244.4 | 317.3 | 254.8 | 364.0 | 90.06 | 130.0 | 130.0 | 364.0 |
| Pressure psia | 553.3 | 553.3 | 600.0 | 548.3 | 30.0 | 20.0 | 15.0 | 20.0 | 20.0 | 20.0 |
| Total Flow lb/hr | 10398 | 1606 | 1606 | 19591 | 19591 | 12380 | 12380 | 7075 | 136 | Trace |
| Mass Flow | | | | | | | | | | |
| GLYCEROL | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| WATER | 5559 | 308 | 308 | 6809 | 6809 | 38 | 38 | 6765 | 9 | Trace |
| HYDROGEN | 0 | 995 | 995 | 2 | 2 | 0 | 0 | 0 | 2 | 0 |
| PG | 4734 | 18 | 18 | 12626 | 12626 | 12342 | 12342 | 285 | 0 | Trace |
| N-PROP | 55 | 60 | 60 | 76 | 76 | 0 | 0 | 21 | 55 | 0 |
| ISO-PROP | 48 | 224 | 224 | 76 | 76 | 0 | 0 | 2 | 74 | 0 |
| SODIU-01 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| METHA-01 | 2 | 0 | 0 | 2 | 2 | 0 | 0 | 2 | 0 | 0 |
| Mass Frac | | | | | | | | | | |
| GLYCEROL | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| WATER | 0.535 | 0.192 | 0.192 | 0.348 | 0.348 | 0.003 | 0.003 | 0.956 | 0.043 | Trace |
| HYDROGEN | 0.000 | 0.620 | 0.620 | Trace | Trace | Trace | Trace | Trace | 0.012 | 0.000 |
| PG | 0.455 | 0.011 | 0.011 | 0.644 | 0.644 | 0.997 | 0.997 | 0.040 | Trace | Trace |
| N-PROP | 0.005 | 0.037 | 0.037 | 0.004 | 0.004 | Trace | Trace | 0.003 | 0.404 | 0.000 |
| ISO-PROP | 0.005 | 0.139 | 0.139 | 0.004 | 0.004 | Trace | Trace | Trace | 0.540 | 0.000 |
| SODIU-01 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 | 0.000 |
| METHA-01 | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace | Trace | 0.000 |
| | | | | | | | | | | |

BENCH SCALE LABORATORY WORK

Bench-Scale Laboratory Work

The kinetic model described in the work of Zhou, Li, et al. was tested in a very small bench scale reactor with inner diameter 10 mm and length 450 mm. One major concern in the development of the propylene glycol production process is the plausibility of scaling this process up to the large industrial reactors that are necessary to accommodate the desired 100 million lb/year throughput. The process-scale reactor has a projected diameter of 5.3 feet $(1.6 \times 10^3 \text{ mm})$, which is over 100 times larger than the bench-scale reactor.

Most models used to describe the scale-up of trickle bed reactors make the following assumptions (Dudukovic 1984):

- i. The gaseous reactant is limiting;
- ii. Internal particle diffusion resistance is present;
- iii. Catalyst particles are completely externally and internally wetted;
- iv. Gas solubility can be treated by Henry's law;
- v. Isothermal operation;
- vi. Axial dispersion model can be used to describe deviations from plug flow;
- vii. Intrinsic reaction kinetics exhibit first order behavior.

The glycerol to propylene glycol reaction is novel among reactions of its kind because it is not vapor phase reactant limiting; the hydrogen gas is flowing at a 5:1 molar excess in the reactor, so the limiting reactant is liquid glycerol. Because of this, the existing models must be modified considerably for use with this reaction.

One method to approach this challenge might be to, with a considerable amount of additional bench-scale lab work, develop a new correlation and model to fit the reaction's parameters, and use it to determine the size of an industrial-scale reactor. However, the lack of the necessary time and materials to perform this research precluded this as an option.

Instead, a model was built to calculate the conversion achieved in the reactor as a function of catalyst mass, using the reaction kinetics to couple the reaction rate to the input flow rates and compositions (see Appendix). An assumption was made that it is possible to reach 100% conversion of glycerol by extrapolating the catalyst's mass to a large enough value. However, this assumption is highly optimistic. In the work of Zhou et. al., the actual conversion of 81.5% was achieved using the Cu-ZnO-Al₂O₃ catalyst with a 1:1:0.5 molar metal ratio. A more precise value of the conversion should be determined using an intermediate scale reactor prior to plant construction. It can be expected that the actual value of conversion in the reactor should fall between the 81.5% in the activity test and the 99.8% that was extrapolated using the Excel model.

The financial risk involved with undertaking an endeavor such as this process calls for a very high degree of confidence in the plant's scalability, which can be provided by a pilot plant test of a mid-size reactor. This is recommended prior to plant construction to ensure the reliability of the scale-up model described above.

PROCESS DESCRIPTION

Pretreatment

The first step in the process involves removing the salt and volatile organic compounds from the crude glycerol. Crude glycerol arrives at our plant directly from biodiesel refineries. It contains by mass, 1% methanol, 4% sodium chloride, 15% water, and trace amounts of organic salts. Almost 20000 lbs/hr of crude glycerol are required, which equates to about one tanker truck each hour. Depending on specific plant geography and transportation networks, rail and ship options are viable; however, 24 tanker trucks per day for raw material and finished product shipment is a reasonable estimate.

The crude glycerol is then mixed with process water in order to reduce the viscosity of the mixture to put through the EDI machine. The resultant mix is 50% wt. glycerol. Process water requirements are 5390 lb/hr from a source, and the rest is the recycle from the distillate from column D-101 in S-112.

The glycerol and water are mixed in M-102. This mixer contains a paddle and electrical input because while glycerol and water are readily miscible, the difference in viscosity requires agitation to achieve homogeneity.

The resultant stream, S-103, is then fed into the electrodeionization (EDI) machine, which proceeds to remove the NaCl and trace sulfates and phosphates. The salt is concentrated in a concentrate H₂O stream, S-104 which is usually 15% of the total mass flow flow through the EDI device in stream S-103. This is about 5100 lb/hr of process water. The assumption for the project is that the amount of organic salt is negligible, and so the salt water concentrate can be disposed of at normal sewerage rates for commercial enterprises in Louisiana. However, if organic salts are higher than expected, or the plant is located in a marshland or other salinity

sensitive zone, than sewerage rates will be higher as the salt cannot simply be sent into the environment.

The desalted glycerol in S-105 is then pumped to replace the head loss across the mixer and EDI device. This is then heated in a feed-product heat exchanger with the distillate from D-101, S-109. The glycerol stream is then heated further with medium pressure steam. This preheating reduces the duty requirement on the reboiler in the column.

The glycerol stream, S-108, enters column D-101 in order to both remove the methanol and reduce the water content to 20% wt, which is the optimal concentration to use in the trickle bed reactor, due to viscosity concerns. The column contains 4 theoretical trays, a total condenser, and a reboiler. The tray efficiency is approximately 9.3% due to the viscosity of glycerol, so the actual column requires 43 trays to operate. The condenser uses cooling water from 80 F to 130F.

This is a relatively easy separation. The water is the light key and it, with methanol, concentrate in the distillate. The distillate, S-109, exits as a saturated liquid at 20 psia and 221 F. The flow rate is 14000 lb/hr and 4% MeOH by weight. This liquid is cooled to 80 F in HX-101 and then 34% of the total flow is purged in S-113 in order to prevent methanol buildup in the pretreatment section. The rest of the flow is then pumped in P-102 to restore pressure to 15 psig in order to flow freely through the mixer and EDI. The purge stream is combined with the propanol/water distillate, S-136, at the end of the process and sent to wastewater treatment. Since this stream contains organic material, we used the Vrana organic wastewater correlation to determine the cost of treatment.

$$Cost \left[\frac{\$}{lb}\right] = 0.044 + 0.048 * (\% \text{ organic impurities by weig } ht)$$

The clean glycerol and 15% water exits the reboiler at 5 psig and 248 F in S-114. The reboiler is heated with medium pressure steam.

Reactor Section

The refined glycerol from the separation column D-101in S-114 is pumped to 585 psia in P-103 and heated in HX-102 with the reactor vapor effluent, S-123, to 416 F. Hydrogen gas is drawn from a pipeline at 1000 psig in S-117, and is depressurized in the turbine T-101 to reach the operating pressure, 600 psia. This hydrogen stream is mixed with the stream of recycled hydrogen, S-131, to maintain a 5:1 molar ratio of hydrogen to glycerol in S-119. The combined hydrogen stream is heated to process temperature (416.5 F) by condensing the vapor effluent, S-126, from the second flash vessel, F-102.

In order to achieve the maximum conversion of glycerol to propylene glycol in a multiphase system, a trickle-bed reactor was selected. The trickle bed reactor is the simplest reactor type for performing multiphase catalytic reactions, and is used widely in industry for such applications as liquid-phase hydrogenation and oxidation of harmful chemicals in waste streams. Trickle-bed reactors are used industrially for hydrodesulfurization processes, which lends credibility to the use of scale-up from the bench-scale analysis. The reactor, R-101, consists of a packed bed of solid catalyst particles, over which streams of gas (S-120) and liquid (S-116) flow down the reactor.

The catalyst that was selected for this process is Cu-ZnO-Al₂O₃ in a 1:1:0.5 molar ratio of Cu:Zn:Al, as it was determined to exhibit the best performance for glycerol hydrogenolysis by (Zhou 2010). The catalyst particles that showed optimal performance in studies by Zhou, Li, et al. had a diameter of 0.17 mm.

Process Description

The overall goal of this process is to achieve 100% conversion and 98.5% selectivity of the incoming glycerol to propylene glycol. Based on this specification, it was determined that a mass of 4190 kg of catalyst is necessary. The small diameter of the catalyst pellets allows for the assumption of a void fraction in the bed of 0.35, a typical value for catalyst pellets of small diameter. 21,000 lb/hr of reactants are fed to the reactor. Professor Leonard Fabiano suggested the assumption of a flux of 1000 lb/ft²*hr through the packed bed, and using this assumption, a 21 ft² cross section is required to accommodate the flow. In a cylindrical vessel, this corresponds to a diameter of 5.2 feet. Based on the fluid properties of the liquid and vapor streams, trickle bed reactor of these dimensions, the Ergun equation can be used to calculate a pressure drop of 20 psi along the length of the reactor (See Appendix for sample calculation).

Additionally, a 5:1 molar excess of hydrogen gas must be accounted for in the reactor sizing. At the operating temperature (455.8 F) and pressure (556 psia), the excess volume of hydrogen is 11,112 ft³/hr. Assuming a retention time of 1 minute in the vessel, calculated based on the effluent flow rate, the total volume of the reactor vessel must be 240 ft³, an increase of almost double the volume for stoichiometric reactants of 142 ft³. Accounting for the hydrogen excess, the new reactor vessel requires a diameter of 6 ft and height of 11 ft.

For the economic analysis of the reactor, catalyst prices were assumed equivalent to those of MK-121, a Cu-Zn-Al high activity catalyst for methanol synthesis that is prepared by Haldor Topsoe, a leading manufacturer of commercial catalysts. Consultations with company representative Mr. Henrik Rasmussen of Haldor Topsoe revealed that this prepared catalyst is sold for \$27/kg in 6 x 4 mm tablets. The commercial catalyst has dimensions too large for use with the chosen kinetic model; the pellets must have a diameter of 0.22 mm or smaller to eliminate the internal diffusion limitation effects of the transport of fluid around the catalyst. To

crush the tablets into small-diameter catalyst particles of diameter approximately equal to 0.17 mm for use in the reactor, an additional \$3/kg surcharge is assumed for processing, for a total of \$30/kg. Therefore, the total purchase cost of one load of 4190 kg catalyst is \$125,700. To account for replacing the catalyst for the main reactor and alternate reactor every two years over the project's life, the total purchase cost of \$1.97 million was used in the economic analysis.

Cu/ZnO/Alumina catalysts have a typical life of approximately 2-3 years and cannot be reliably regenerated by burning. To ensure that there is negligible loss in catalyst performance throughout the plant's lifetime, the catalyst mass will be replaced every year. Two reactor vessels of equal size will be purchased for the plant and connected in parallel. The reactor section process flowsheet displays two sets of valves, labeled R-101A and R-101B corresponding to each reactor vessel. It is relatively simple to switch the reactor vessel that is in operation by alternating which set of valves is open and which is closed. At all times, only set of either R-101A or R-101B will be open. When a reactor is not operational, it may be detached from the plant and repacked with new catalyst. Each year, the plant operators will alternate which reactor is active in the process, to allow the second vessel to be cleaned and packed with new catalyst. This setup also allows for fast and relatively easy introduction of the replacement reactor in case of any kind of malfunction during the operational year.

Both the refined glycerol and mixed hydrogen streams, S-116 and S-120, enter the trickle bed reactor, where the glycerol is completely converted into propylene glycol and some propanol. The heat of reaction varies slightly with temperature. The reactor is assumed to operate adiabatically, as the heat released is relatively small, and it can be removed through bulk transport in the vapor stream. The mixed-phase effluent flowing out of the reactor, S-121, is collected in a flash vessel, F-101, where the liquid stream, S-122, is sent to the refinement section. The vapor, S-123, is then partially condensed in heat exchanger HX-102 and then flows to another flash vessel, F-102. The liquid stream from the flash vessel, S-125, becomes one of the input streams to the separation section for refinement and purification. The vapor, S-126, that exits the flash vessel is partially condensed in HX-103 and further cooled in C-102 until the vapor phase contains pure hydrogen. This stream is sent to a third flash vessel, F-103. The vapor is then recompressed in CP-101 to the operating pressure, 600 psia, and recycled, and the liquid in S-129 is fed to the separation section.

Separation Section

The separation section purifies the propylene glycol to 99.8% purity by weight. There is only one 2-product distillation column, D-102, required because the propanol/water mixture, S-136, is sent to wastewater treatment. The incoming feed to the separation column, S-133, contains propylene glycol, propanol, and water. Since the patent suggested that there would be 100% conversion, no glycerol will be present in the reactor effluent, and the only design constraint other than product purity was keeping the reboiler temperature, 364 F, below the degradation temperature of propylene glycol, which is 369 F.

The RADFRAC block in ASPEN suggested that 14 theoretical trays would be required to obtain the desired separation. In order to calculate the overall efficiency of the column, the O'Connell correlation was used. This relates the ratio of the number of theoretical trays to the total number of trays, η , as a function of the feed viscosity (in cP) and relative volatility of the key components.

$$\eta = 0.492(\mu_F \alpha)^{-0.245}$$

The O'Connell correlation suggested that the efficiency was 9.8%, therefore the total number of trays required for this column was 70. This is the upper limit on column height. The condenser is a partial condenser, with a vapor pilot flame of hydrogen (S-137) and a liquid water/propanol distillate (S-136). The bottoms product is 99.6% pure propylene glycol at 364 F, which is then cooled in C-103 to handling temperature.

At the bottom of the column there will be a boot attached in order to collect the heavy compounds produced, such as ethylene glycol. The bottoms product will be drawn off as a vapor which will increase the cooling load on C-103. The boot residue will be drawn off periodically in S-138 to prevent contamination.
Section 12

HEAT INTEGRATION: ENERGY BALANCE AND UTILITY REQUIREMENTS

Heat Integration

Because EDI was used for the glycerol pretreatment, the overall utilities required for the process are less than the requirements for other processes on this scale. Additionally, no heater or reboiler is required to operate above 365 F, which is the saturation temperature of 150 psig steam. Since it is less expensive, low to medium pressure steam can be used. Heat integration was achieved primarily through feed/product heat exchange, and required just 2.41 million Btu/hr of heating and 10.08 Btu/hr of cooling utilities, excluding the condensers and reboilers.

In the pretreatment section, the hot water/methanol distillate, S-109, is exchanged with the stream fed to the distillation column, S-106, in HX-101. S-109 is cooled to room temperature, and S-106 receives additional heat in H-101. This design was chosen in order to reduce the size of the reboiler in the first distillation column, D-101. HX-101 exchanges 2.14 million Btu/hr and has an area of 2052 sq. ft.

In the reactor section, the design objectives are to heat the incoming glycerol and hydrogen streams (S-114 and S-120) to the reactor inlet temperature of 416 F, and to recover all of the propylene glycol from the reactor outlet. Since a substantial amount of PG is in the vapor phase at the outlet of the reactor, we decided to design a system of flash vessels and heat exchangers in order to condense PG from hydrogen. The reactor effluent at 456 F is sent to F-101, which collects the PG and water that is liquid in S-122. The vapor (S-123) is then used to heat the incoming glycerol stream to the reactor inlet temperature in HX-102, exchanging 1.99 million Btu/hr. This heat exchange condenses more PG, which is collected in a second flash vessel, F-102, operating at 426 F. The vapor from this vessel in S-126 is then used to heat the incoming hydrogen in S-119 to the reactor inlet temperature in HX-103, exchanging 1.13 million

Btu/hr. The vapor stream, S-127, is then cooled further to 225 F in C-102, at which all of the PG is found in the liquid phase. C-102 exchanges 7.50 million Btu/hr. The reason that we were able to eliminate external heating requirements for the reactor is because hydrogen has a very high heat capacity, and the latent heat of condensation for propylene glycol is also high, which means that there is a lot of available thermal energy stored in the reactor effluent.

In the refinement section, the feed stream is sufficiently hot not to require preheating, and the only requirement is that pure PG be cooled to handling temperature which is approximately 90 F. This occurs in C-103, which has a duty of -2.08 million Btu/hr. The liquid distillate, S-136, is mixed with the methanol purge from the pretreatment section in order to prevent flash boiling once the liquid is sent to the organic wastewater storage tank.

Energy balances, computed by ASPEN PLUS, are in the Appendix.

Section 13

EQUIPMENT LIST AND UNIT DESCRIPTIONS

| Unit Number | Unit Type | Function | Material | Size | Temperature (F) | Pressure (psia) |
|-------------|--------------------|-------------------------------------------------|-----------------|-----------------|-----------------|-------------------|
| | | Pre-Treatment Section | n I | | | |
| M-102 | Mixer | Mixes crude Glycerol, fresh H20 and recyle H20 | Stainless Steel | 1 HP | | |
| S-101 | EDI Equipment | Extracts all chloride and organic salts | | | | |
| P-101 | Pump | Transports clean glycerol | Stainless Steel | 75 HP | | |
| HX-101 | Heat Exchanger | Raise temp of liquid stream | Carbon Steel | 2052 sqft. | 221 | 35 |
| H-101 | Fired Heater | Raise temp of liquid stream | Carbon Steel | | 242 | 30 |
| D-101 | Tower | H20 LK, Glycerol HK | Carbon Steel | 3.9ft x 100ft | | 20 |
| D-101a | Heat Exchanger | Column Condensor | Carbon Steel | 654 sqft. | 221 | 20 |
| D-101b | Reflux Accumulator | Reflux Accumulator | Carbon Steel | 3.21ft x 1.61ft | | |
| D-101c | Pump | Pumps bottoms into Reboiler | Carbon Steel | 75 HP | | |
| D-101d | Heat Exchanger | Column Reboiler | Carbon Steel | 2438 sqft. | 248 | 20 |
| P-103 | Pump | Transports clean glycerol | Stainless Steel | 250 HP | 252 | 587 |
| P-102 | Pump | Transports H20 recycle | Stainless Steel | 75 HP | 80 | 30 |
| | _ | Reactor Section | | | | |
| R-101a | Reactor | Contains catalyst | Stainless Steel | 5.2ft x 11.5ft | 455 | 565 |
| R-1010 | Reactor | | Stainless Steel | 5.2ft x 11.5ft | 455 | 565 |
| F-101 | Flash | Separates liquid PG from H2 | Stainless Steel | 1.7ft x 6.75ft | 455 | 560 |
| HX-102 | Heat Exchanger | Raise temp of Glycerol feed | Stainless Steel | 73 sqft. | 455 | 587 |
| F-102 | Flash | Separates liquid PG from H2 | Stainless Steel | 1.51ft x 5.36ft | 426 | 540 |
| HX-103 | Heat Exchanger | Raise temp of H2 feed | Stainless Steel | 165 sqft. | 426 | 585 |
| C-102 | Heat Exchanger | Cools PG/H2 mixture | Stainless Steel | 256 sqt. | 225 | 520 |
| F-103 | Flash | Separates liquid PG from H2 | Stainless Steel | 0.74ft x 1.5ft | 225 | 520 |
| CP-101 | Compressor | Repressurize H2 Recycle | Stainless Steel | 123 HP | 225 | 520 |
| TR-100 | Turbine | Reduces pressure of fresh H2 | Stainless Steel | 27.7 HP | 12 | 400 psia drop |
| M-101 | Mixer | Mix fresh H2 and recycle H2 | Stainless Steel | 7 HP | 600 | 210 |
| | | Separation Section | | | | |
| M-103 | Mixer | Mix PG product streams | Stainless Steel | 1 HP | 317 | 515 |
| V-101 | Valve | Decrease pressure of product stream | | | | 485 pressure drop |
| D-102 | Tower | H20 LK, PG HK | Carbon Steel | 2.8ft x 154ft | | 20 |
| D-102a | Heat Exchanger | Column Condensor | Carbon Steel | 538 sqft. | 130 | 20 |
| D-102b | Reflux Accumulator | Reflux Accumulator | Carbon Steel | 1.58ft x 3.15ft | | |
| D-102c | Pump | Pumps bottoms liquid into Reboiler | Carbon Steel | 75 HP | | |
| D-102d | Heat Exchanger | Column Reboiler | Carbon Steel | 419 sqft. | 364 | 20 |
| C-103 | Heat Exchanger | Cools final PG stream | Carbon Steel | 191 sqft. | 90 | 15 |
| | | Storage Vessels | [| Γ | 1 | |
| T-101 | Floating Roof Tank | Store 2-wks of Glycerol Feedstock | Carbon Steel | 724,559 gal | | 30 |
| T-102 | Floating Roof Tank | Store 2-wks of NaCl/H20 EDI Concentrate Streams | Carbon Steel | 424,453 gal | | 20 |
| T-103 | Floating Roof Tank | Store 2-wks of Methanol/H20 Purge | Carbon Steel | 25,907 gal | | 15 |
| T-104 | Floating Roof Tank | Store 2-wks Propanol/H20 Product | Carbon Steel | 8,400 gal | | 20 |
| T-105 | Floating Roof Tank | Store 8-hr of PG Product | Carbon Steel | 11,607 gal | | 15 |
| T-106 | Floating Roof Tank | Store 8-hr of PG Product | Carbon Steel | 11,607 gal | | 15 |
| T-107 | Floating Roof Tank | Store 2-wks of PG Product | Carbon Steel | 482,718 gal | | 15 |

Detailed Equipment Descriptions

The following bare-module costs were calculated using correlations, calculations and heuristics were taken from the text <u>Product & Process Design Principles</u> by Sader, Seider and Lewin. The price index for all calculations were based on CE Index = 560.4 from the most recent reported value from December 2010 (Chemical Engineering 2011). This index was used to account for inflation and provide the most accurate estimation possible for the financial analysis presented later in the report.

M-102 (Mixer)

Bare-Module Cost: \$3,699

The crude glycerol feed, fresh water feed and water recycle are mixed in M-102. The water concentration in the resultant stream, S-103 is increased to 50% by weight. This specification is required for the electrodeionization process.

EDI (Electrodionization Equipment)

Bare-Module Cost: \$4,483,200

The EDI machine strips the salts from the glycerol/water mixture and transports the ions to a concentrate stream (S-104) for disposal into the sewer system. It uses approximately 200 kW of electricity.

P-101 (Pump)

Bare-Module Cost: \$13,993

The pump P-101 is used to raise the pressure of output stream from the EDI by 22 psia.

The break horsepower is 1.63 HP to process 471 cuft/hr. Stainless steel is used for construction.

HX-101 (Heat Exchanger)

Bare-Module Cost: \$77,519

This heat exchanger is used to cool the distillate (S-109) back to room temperature, as well as to preheat the 33230 lb/hr feed to column D-101. It exchanges 1990000 Btu/hr with a surface area of 2052 sq ft. Carbon steel is used on both sides.

H-101 (Heater)

Bare-Module Cost: \$222,840

The heater H-101 is used to give additional heating to stream S-107 in order to reduce the duty requirements on the reboiler in D-101. It uses medium pressure steam to raise the temperature to 242 F, and has a duty requirement of 2410000 Btu/hr. Carbon steel is used on both sides.

D-101 (Distillation Column)

Bare-Module Cost: \$1,166,781

The feed stream into distillation column D-101 is S-108, which is composed of water, methanol, and glycerol. D-102 has 4 theoretical trays and an O'Connell efficiency of 0.093, giving 43 trays. The height of the column is 100 ft and the diameter is 3.9 ft. Carbon steel is used

in construction. The molar reflux ratio is 0.2 and the feed enters halfway up the column. The column operates at 20 psia and the temperature of the reboiler and total condenser are 248 F and 221 F respectively. The purpose of this separation is to remove 70% of the water and all of the methanol from the glycerol stream. Since methanol is lighter than water, both of these requirements can be accomplished at once. The condenser and reboiler duties are 16480000 and 15740000 Btu/hr respectively.

P-102 (Pump)

Bare-Module Cost: \$14,847

P-102 is used to raise the pressure of the water recycle stream to compensate for pressure losses through the machinery. It is made of stainless steel and has a brake horsepower of 0.38 hp.

P-103 (Pump)

Bare-Module Cost: \$15,097

Pump P-103 is used to raise the pressure of the refined glycerol to the reactor operating pressure of 581 psia. It is made of stainless steel and moves 267.0 cu ft of liquid with a brake power of 29.01 hp.

HX-102 (Heat Exchanger)

Bare-Module Cost: \$67,135

HX-102 raises the temperature of the refined glycerol to the reactor operating temperature of 410 F by cooling and condensing the hot vapor effluent from the reactor. ASPEN suggested that the surface area of this heat exchanger is 73 sq ft. The reason why this is low is because of the very high heat capacity of hydrogen and the additional heating that occurs due to condensation of the propylene glycol. It is made of stainless steel due to the presence of hydrogen.

TR-100 (Turbine)

Bare-Module Cost: \$57,089

Turbine TR-100 is an attempt to recover some of the energy as the hydrogen feed pressure is lowered to the reactor operating pressure of 580 psia. It is made of stainless steel and produces 28 hp of power.

CP-101 (Compressor)

Bare-Module Cost: \$258,602

Compressor CP-101 repressurizes the hydrogen recycle to the reactor operating pressure of 580 psia. It is made of stainless steel, moves 6947 cu ft/hr and requires a brake power of 28.67 hp.

M-101 (Mixer)

Bare-Module Cost: \$20,233

M-101 is a mixer that brings the hydrogen recycle and the hydrogen feed together in a small, stainless steel vessel.

HX-103 (Heat Exchanger)

Bare-Module Cost: \$67,421

Heat exchanger HX-103 heats the combined hydrogen in stream S-119 to the reactor operating temperature of 410 F. It uses the vapor from the second flash vessel (F-102) as the hot stream. It exchanges 1130000 Btu/hr and is made of stainless steel. It has a surface area of 165 sq ft.

R-101 (Reactor)

Bare-Module Cost: \$796,642

Reactor R-101 is a trickle bed reactor made out of stainless steel and is 5.2 ft x 11.5 ft (DxL). It contains 4800 kg of Cu-ZnO-Al₂O₃ catalyst and converts 100% of the glycerol to propylene glycol and propanol. The reactor operates adiabatically since the heat of reaction can be effectively removed solely through bulk transport of the effluent. See the reactor section for more information.

F-101 (Flash Vessel)

Bare-Module Cost: \$111,058

Flash vessels were found to be the most effective way to separate the hydrogen from the propylene glycol product. F-101 is placed directly after the reactor to collect the hot liquid product. The vessel is 6.75 ft x 1.7 ft. The vapor outflow contains 52% propylene glycol by mass.

F-102 (Flash Vessel)

Bare-Module Cost: \$89,031

The vapor from F-101 is cooled in HX-102 to 429F, which allows for liquid to form. This stream is separated in F-102, a stainless steel 1.51 ft x 5.36 ft vessel. The vapor outflow from here contains 39.6% propylene glycol by mass.

C-102 (Cooler)

Bare-Module Cost: \$85,504

Cooler C-101 is designed to cool the vapor from F-102 to 225 F, which at the design pressure of 553 psia, results in a vapor-liquid equilibrium that has virtually all of the propylene glycol in the liquid phase. The cooler is made of stainless steel and uses cooling water with a required duty of 7497000 Btu/hr.

F-103 (Flash Vessel)

Bare-Module Cost: \$31,808

F-103 separates the effectively pure hydrogen vapor from the 10398 lb/hr propylene glycol and water liquid. It is made of stainless steel and is 0.74×1.5 ft. The vapor is sent to the recycle compressor.

M-103 (Mixer)

Bare-Module Cost: \$3,699

M-103 combines the three liquid effluents from the flash vessels and then depressurizes the mixture to 30 psia, which is the feed to column D-102. The total flow rate is 19591 lb/hr.

D-102 (Distillation Column)

Bare-Module Cost: \$2,477,162

The feed stream into distillation column D-121 is S-133, which is composed of water, propanol, and glycerol at 34.7%, 0.7%, and 64.5% propylene glycol by mass. D-102 has 14 theoretical trays and an O'Connell efficiency of 0.198, giving 70 trays. The height of the column is 154 ft and the diameter is 2.8 ft. Carbon steel is used in construction. The molar reflux ratio is 0.13 and the feed enters halfway up the column. The column operates at 20 psia and the temperature of the reboiler and total condenser are 364 F and 130 F respectively. The reboiler operates at a temperature that is low enough to use medium pressure steam. The condenser and reboiler duties are 8590000 and 7110000 Btu/hr respectively. The condenser is a partial condenser. The liquid condensate is sent to wastewater treatment while the vapor is hydrogen which is combusted in a pilot flame.

C-103 (Cooler)

Bare-Module Cost: \$75,539

Cooler C-103 reduces the temperature of the final product to a safe handling temperature of 90 F. It is made of carbon steel and the duty is 2580000 Btu/hr. The product has 99.7 % purity.

82

Section 14

SPECIFICATION SHEETS

| | CO | OLER | | |
|------------------------------|----------------------------------|----------------------|--------------|-----------------|
| Identification: | Item Item No. No. required | Cooler C-103 1 | Date: By: | 4-Apr-11 KMH |
| Function: To cool the p | ropylene glycol p | product for storage | e. | |
| Operation: Continuous | | | | |
| Materials handled: | Inlet | Outlet | | |
| | S-134 | S-135 | | |
| Quantity (lb/hr): | 12380 | 12380 | | |
| Composition: (mass fraction) | | | | |
| Water | 0.003 | 0.003 | | |
| Hydrogen | 0 | 0 | | |
| Propylene glycol | 0.997 | 0.997 | | |
| N-propanol | 0 | 0 | | |
| Isopropanol | 0 | 0 | | |
| Methanol | 0 | 0 | | |

| Design Data: | Outlet temperature: 90 F | |
|--------------|--------------------------|--|
| | Outlet pressure: 15 psia | |
| | Vapor fraction: 0 | |
| | Tube length: 20 ft | |
| | Surface area: 183 sqft. | |
| | | |
| | | |

Utilities: Heat duty: -2581912 Btu/hr Controls: Tolerances: Comments and drawings: See process flow sheet, separation section

Equipment Base f.o.b. Cost: \$18,252 Bare Module Factor: 1.0 Bare Module Cost: \$85,504 N-Propanol Isopropanol

Methanol

COMPRESSOR

| Identification: | Item Item No. No. required | Isentropic Compressor CP-101 1 | Date: By: | 11-Apr-11 KMH |
|-----------------------------|-----------------------------------------|--------------------------------------|--------------|------------------|
| Function: To increase | the pressure of | stream S130. | | |
| Operation: Continuous | | | | |
| Materials handled: | Inle | Outlet | | |
| | S-130 | S-131 | | |
| Quantity (lb/hr): | 1600 | 5 1606 | | |
| Composition: (mass fraction |) | | | |
| Water | 0.192 | 0.192 | | |
| Hydrogen | 0.620 | 0.620 | | |
| Propylene Glycol | 0.01 | 0.011 | | |

0.037

0.139

trace

0.037

0.139

trace

| Design Data: | Net work required: 28.67 hp Efficiency: 0.80 |
|--------------|-------------------------------------------------|
| | Discharge pressure: 600 psia |
| | Outlet temperature: 244 F |
| | Isentropic outlet temperature: 241 F |
| | Vapor fraction: 1.0 |

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, pretreatment section.

Equipment base f.o.b. cost: \$92,292 Bare Module Factor: 1 Bare Module Cost: \$258,603

DISTILLATION COLUMN

| Identification: | Item | Distillation Column | | |
|-----------------|--------------|---------------------|-------|-----------|
| | Item No. | D-101 | Date: | 30-Mar-11 |
| | No. required | 1 | By: | KMH |

Function: To separate glycerol from water to purify reactor feed stream.

| Operation: Continuous | | | |
|-----------------------------|-------|------------|---------|
| Materials handled: | Feed | Vapor Dist | Bottoms |
| | S-108 | S-109 | S-114 |
| Quantity (lb/hr): | 33230 | 13985 | 19245 |
| Composition: (mass fraction | | | |
| Glycerol | 0.467 | 0.960 | 0.806 |
| Water | 0.516 | 0 | 0.194 |
| Methanol | 0.017 | 0.040 | trace |

| Design Data: | Number of theoretical trays: 4 |
|--------------|----------------------------------------|
| | Pressure: 20 psia |
| | Functional height: 100 ft |
| | Material of construction: Carbon steel |
| | Recommended inside diameter: 3.8 ft |
| | Tray efficiency: 0.093 |
| | Feed stage: 22 |

Molar reflux ratio: 0.2 Tray spacing: 2.0 ft Skirt height: 14.5 ft Vessel weight: 8507 lb Wall thickness: 1/4 in

Utilities: Condenser duty: -16479327 Btu/hr, Reboiler duty: 15739914 Btu/hr Controls: Tolerances: Comments and drawings: See pretreatment section process flow sheet

| Equipment base f.o.b. cost: | \$125,546 |
|-----------------------------|-----------|
| Bare-Module Factor: | 4.16 |
| Bare Module Cost: | \$522,271 |

DISTILLATION COLUMN

| Identification: | Item | Distillation Column | | |
|-----------------|--------------|---------------------|-------|-----------|
| | Item No. | D-102 | Date: | 30-Mar-11 |
| | No. required | 1 | By: | KMH |

| Function: | Purify propylene glycol | product stream b | ov separating water | , PG, and propanol. |
|-----------|--------------------------|------------------|---------------------|---------------------|
| r uncuon. | i unity propytene grycol | produce stream c | y separating water | , 1 O, and propano |

| Operation: Continuor | us | | | |
|--------------------------|-------|-------------|--------------|---------|
| Materials handled: | Feed | Vapor Dist. | Liquid Dist. | Bottoms |
| | S-133 | S-137 | S-136 | S-135 |
| Quantity (lb/hr): | 19591 | 136.1 | 7075 | 12380 |
| Composition: (mass frac) | | | | |
| Glycerol | 0 | 0 | 0 | 0 |
| Water | 0.348 | 0.043 | 0.956 | 0.003 |
| Hydrogen | trace | 0.012 | trace | 0 |
| Propylene Glycol | 0.644 | trace | 0.040 | 0.997 |
| N-propanol | 0.004 | 0.404 | 0.003 | trace |
| Isopropanol | 0.004 | 0.540 | trace | trace |
| Methanol | 0 | trace | trace | trace |

| Design Data: | Number of theoretical trays: 14 | Molar reflux ratio: 0.13 |
|--------------|----------------------------------------|--------------------------|
| | Pressure: 20 psia | Tray spacing: 2.0 ft |
| | Functional height: 156 ft | Skirt height: 14.5 ft |
| | Material of construction: Carbon steel | Vessel weight: 6025 lb |
| | Recommended inside diameter: 2.8 ft | Wall thickness: 1/4 in |
| | Tray efficiency: 0.198 | |
| | Feed stage: 46 | |

| Utilities: | Condenser duty: -8588644 Btu/hr, Reboiler duty: 7106169 Btu/hr |
|--------------|----------------------------------------------------------------|
| Controls: | |
| Tolerances: | |
| Comments and | drawings: See process flow sheet, separation section |

Equipment base f.o.b. cost: \$112,587 Bare-Module Factor: 4.16 Bare Module Cost: \$468,362

FLASH VESSEL

| Identification: | Item | Vertical Pressure Vessel | | |
|-----------------|--------------|--------------------------|-------|----------|
| | Item No. | F-101 | Date: | 4-Apr-11 |
| | No. required | 1 | By: | KMH |

Function: To purify propylene glycol reactor product.

| Operation: Continuous | | | | |
|------------------------------|-------|------------|---------|--|
| Materials handled: | Feed | Distillate | Bottoms | |
| | S-121 | S-123 | S-122 | |
| Quantity (lb/hr): | 21197 | 16880 | 4317 | |
| Composition: (mass fraction) | | | | |
| Water | 0.336 | 0.393 | 0.114 | |
| Hydrogen | 0.047 | 0.059 | trace | |
| Propylene glycol | 0.597 | 0.524 | 0.881 | |
| N-propanol | 0.006 | 0.008 | 0.002 | |
| Isopropanol | 0.014 | 0.017 | 0.003 | |
| Methanol | trace | trace | trace | |

| Design Data: | Pressure: 560 psia |
|--------------|----------------------------------------|
| | Vapor fraction: 0.927 |
| | Functional height: 6.8 ft |
| | Inside diameter: 1.7 ft |
| | Material of construction: Carbon steel |
| | Vessel weight: 1188 lb |
| | Wall thickness: 0.65 in |

| Utilities: | Heat duty: 0 |
|--------------|---------------------------------------------------|
| Controls: | |
| Tolerances: | |
| Comments and | drawings: See process flow sheet, reactor section |

Equipment base f.o.b. cost: \$26,697 Bare-Module Factor: 4.16 Bare Module Cost: \$111,058

| FLASH VESSEL | | | | |
|---------------------|-----------------------------------------|----------------------------------------|---------------------------|---|
| Identification: | Item Item No. No. required | Vertical Pressure Vessel F-102 1 | Date: 4-Apr-11 By: KMH | |
| Function: To purify | propylene glycol | reactor product. | | _ |

| Operation: Continuous | | | | |
|------------------------------|-------|------------|---------|--|
| Materials handled: | Feed | Distillate | Bottoms | |
| | S-124 | S-126 | S-125 | |
| Quantity (lb/hr): | 16880 | 12004 | 4876 | |
| Composition: (mass fraction) | | | | |
| Water | 0.393 | 0.489 | 0.156 | |
| Hydrogen | 0.059 | 0.083 | trace | |
| Propylene glycol | 0.524 | 0.396 | 0.838 | |
| N-propanol | 0.008 | 0.01 | 0.002 | |
| Isopropanol | 0.017 | 0.023 | 0.003 | |
| Methanol | trace | trace | trace | |

Design Data: Pressure: 560 psia Vapor fraction: 0.902 Functional height: 5.4 ft Inside diameter: 1.5 ft Material of construction: Carbon steel Vessel weight: 766 lb Wall thickness: 0.583 in

Utilities: Heat duty: 0 Controls: Tolerances: Comments and drawings: See process flow sheet, reactor section

Equipment base f.o.b. cost: \$21,402 Bare Module Factor: 4.16 Bare Module Cost: \$89,031

| | FLA | SH VESSI | EL | | |
|------------------------------|-----------------------------------------|-------------------------------|-----------|--------------|-----------------|
| Identification: | Item Item No. No. required | Vertical Pressu F-103 1 | re Vessel | Date: By: | 4-Apr-11 KMH |
| Function: To purify pro | opylene glycol r | eactor product. | | | |
| Operation: Continuous | | | | | |
| Materials handled: | Feed | Distillate | Bottoms | | |
| | S-128 | S-130 | S-129 | | |
| Quantity (lb/hr): | 12004 | 1606 | 10398 | | |
| Composition: (mass fraction) | | | | | |
| Water | 0.489 | 0.192 | 0.535 | | |
| Hydrogen | 0.083 | 0.619 | trace | | |
| Propylene glycol | 0.396 | 0.011 | 0.455 | | |
| N-propanol | 0.008 | 0.037 | 0.005 | | |
| Isopropanol | 0.023 | 0.139 | 0.005 | | |

trace

trace

trace

| Design Data: | Pressure: 553 psia | |
|--------------|----------------------------------------|--|
| | Vapor fraction: 0.581 | |
| | Functional height: 1.5 ft | |
| | Inside diameter: 0.8 ft | |
| | Material of construction: Carbon steel | |
| | Vessel weight: 77 lb | |
| | Wall thickness: 0.375 in | |

Utilities: Heat duty: 0

Controls:

Methanol

Tolerances:

Comments and drawings: See process flow sheet, reactor section

Equipment base f.o.b. cost: \$7641 Bare Module Factor: 4.16 Bare Module Cost: \$31,786

| | | HEAT | ER | | | |
|-----------------|-----------------------------|-----------------------------------------|-----------------|-------------|------------------|-----------------|
| Identifi | ication: | Item Item No. No. required | Heater H-101 | 1 | Date: By: | 4-Apr-11 KMH |
| Function: | To increase prior to dis | e the temperature tillation. | of the gl | ycerol/wate | er outlet stream | from the EDI |
| Operation: | Continuous | 5 | | | | |
| Materials | handled: | Inlet | | Outlet | | |
| | | S-107 | | S-108 | | |
| Quantity (lb/hr |): | 34007 | | 34007 | | |
| Composition: (| mass fraction | ı) | | | | |
| Glycerol | | 0.467 | | 0.467 | | |
| Water | | 0.516 | | 0.516 | | |
| Methanol | | 0.017 | | 0.017 | | |
| Design Data: | Outlet temp | perature: 242 F | | | | |
| | Outlet pres | sure: 36.8 psia | | | | |
| | Vapor fract | tion: 0 | | | | |
| | | | | | | |
| | | | | | | |

| Utilities: | Heat duty: 2410557 Btu/hr |
|---------------|--------------------------------------------------------|
| Controls: | |
| Tolerances: | |
| Comments and | drawings: See process flow sheet, pretreatment section |
| Equipment bas | se f.o.b. cost: \$106,893 |
| Bare Module H | actor: 1.86 |

Bare Module Cost: \$222,839

HEAT EXCHANGER

| Identification: | Item | Heat exchanger | | |
|-----------------|--------------|----------------|-------|----------|
| | Item No. | HX-101 | Date: | 4-Apr-11 |
| | No. required | 1 | By: | KMH |

0.017

Function: To integrate heat between streams.

| Operation: Continu | ious | |
|---------------------------|--------------|--------------|
| Materials handled: | Hot Stream | Cold Stream |
| | S-109, S-110 | S-106, S-107 |
| Quantity (lb/hr): | 13985 | 33231 |
| Composition: (mass frac | | |
| Glycerol | trace | 0.467 |
| Water | 0.960 | 0.516 |

Methanol

Design Data: Hot side: Tube side, Cold side: shell side Hot stream temperature, Inlet: 221.4 F Outlet: 80.0 F Cold stream temperature, Inlet: 69.9 F Outlet: 151.8 F Material of construction: Carbon steel Heat duty: 2136340 Btu/hr Heat transfer coefficient (U): 33.97 Area: 2052 sqft

0.040

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, pretreatment section.

Equipment base f.o.b. cost: \$21,818 Bare Module Factor: 3.17 Bare Module Cost: \$77,519

HEAT EXCHANGER

| Identification: | Item | Heat exchang | zer | | |
|-----------------|--------------|--------------|-----|-------|----------|
| | Item No. | HX-102 | - | Date: | 4-Apr-11 |
| | No. required | | 1 | By: | KMH |
| | | | | | |

| Function: | To integrate l | heat | between | streams. |
|-----------|----------------|------|---------|----------|
|-----------|----------------|------|---------|----------|

| Operation: Continu | uous | |
|-------------------------|--------------|--------------|
| Materials handled: | Hot Stream | Cold Stream |
| | S-123, S-124 | S-115, S-116 |
| Quantity (lb/hr): | 16880 | 19245 |
| Composition: (mass frac | | |
| Glycerol | 0 | 0.806 |
| Water | 0.393 | 0.194 |
| Hydrogen | 0.059 | 0 |
| Propylene glycol | 0.524 | 0 |
| N-propanol | 0.008 | 0 |
| Isopropanol | 0.017 | 0 |
| Methanol | trace | trace |

Design Data: Hot side: Tube side, Cold side: shell side Hot stream temperature, Inlet: 455 F Outlet: 429 F Cold stream temperature, Inlet: 252 F Outlet: 410 F Material of construction: Carbon steel Heat duty: 1990194 Btu/hr Heat transfer coefficient (U): 344.8 Area: 72.8 sqft

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, reactor section.

Equipment base f.o.b. cost: \$9404 Bare Module Factor: 3.17 Bare Module Cost: \$67,136

HEAT EXCHANGER

| Identification: | Item Item No. No. required | Heat exchanger HX-103 1 | Date: By: | 4-Apr-11 KMH |
|------------------------------|-----------------------------------------|-------------------------------|--------------|-----------------|
| Function: To integrate | heat between stre | eams. | | |
| Operation: Continuous | | | | |
| Materials handled: | Hot Stream | Cold Stream | | |
| | S-126, S-127 | S-119, S-120 | | |
| Quantity (lb/hr): | 1952 | 12004 | | |
| Composition: (mass fraction) | | | | |
| Water | 0.158 | 0.489 | | |
| Hydrogen | 0.687 | 0.083 | | |
| Propylene glycol | 0.009 | 0.396 | | |
| N-propanol | 0.031 | 0.010 | | |
| Isopropanol | 0.115 | 0.023 | | |
| Methanol | trace | trace | | |

| Design Data: | Hot side: Tube side, Cold side: shell side |
|--------------|-----------------------------------------------------|
| | Hot stream temperature, Inlet: 429 F Outlet: 409 F |
| | Cold stream temperature, Inlet: 202 F Outlet: 416 F |
| | Material of construction: Carbon steel |
| | Heat duty: 11132628 Btu/hr |
| | Heat transfer coefficient (U): 145.3 Btu/hr*sqft*R |
| | Area: 164.4 sqft |

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, reactor section.

Equipment Base f.o.b. cost: \$9428 Bare Module Factor: 3.17 Bare Module Cost: \$67,420

| | | MIXER | ł | | | |
|-----------------|-----------------------------------------|----------------|---|--------------|------------------|--|
| Identification: | Item Item No. No. required | Mixer M-101 | 1 | Date: By: | 30-Mar-11 KMH | |

| | Function: | Combine recycled and fresh hydrogen streams for feed to reactor. | |
|--|-----------|------------------------------------------------------------------|--|
|--|-----------|------------------------------------------------------------------|--|

| Operation: Continuous | | | |
|------------------------------|--------|--------|--------|
| Materials handled: | Feed 1 | Feed 2 | Outlet |
| | S-118 | S-131 | S-119 |
| Quantity (lb/hr): | 346.4 | 1605.7 | 1952 |
| Composition: (mass fraction) | | | |
| Water | 0 | 0.192 | 0.158 |
| Hydrogen | 1 | 0.620 | 0.687 |
| Propylene glycol | 0 | 0.011 | 0.009 |
| N-propanol | 0 | 0.037 | 0.031 |
| Isopropanol | 0 | 0.139 | 0.115 |
| Methanol | 0 | trace | trace |

Design Data: Pressure: 600 psia Outlet temperature: 202.4 F Vapor fraction: 0.994 Material of construction: Carbon steel

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, reactor section

Equipment base f.o.b. cost: \$3300 Bare Module Factor: 1.0 Bare Module Cost: \$3699

| | | | MIXE | R | | |
|------------|-----------|----------------------------------------|-----------------------------|-------------|------------------|------------------|
| Identi | fication: | Item Item No. No. require | <i>Mixer</i> M-102 ed | 1 | Date: By: | 30-Mar-11 KMH |
| Function: | Combine | e crude glycerol | , fresh wat | er, and rec | yeled water stre | ams. |
| Operation: | Continue | 0115 | | | | |

| operation: continuous | | | | |
|------------------------------|--------|--------|--------|----------|
| Materials handled: | Feed 1 | Feed 2 | Feed 3 | Outlet 1 |
| | S-101 | S-102 | S-112 | S-103 |
| Quantity (lb/hr): | 19392 | 5384 | 9230 | 34007 |
| Composition: (mass fraction) | | | | |
| Glycerol | 0.800 | 0 | trace | 0.456 |
| Water | 0.150 | 1.000 | 0.960 | 0.504 |
| Methanol | 0.010 | 0 | 0.040 | 0.017 |
| NaCl | 0.040 | 0 | 0 | 0.023 |
| | | | | |

| Design Data: | Pressure: 25 psia |
|--------------|----------------------------------------|
| | Outlet temperature: 54.8 F |
| | Vapor fraction: 0 |
| | Material of construction: Carbon steel |

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, pretreatment section

Equipment base f.o.b. cost: \$3300 Bare Module Factor: 1.0 Bare Module Cost: \$3699

MIXER

| Identification: | Item | Mixer | | | |
|-----------------|-------------|-------|---|-------|-----------|
| | Item No. | M-103 | | Date: | 30-Mar-11 |
| | No. require | d | 1 | By: | KMH |

Function: Combine propylene glycol product streams from three separators for purification.

| Oneration | Continuous |
|------------|------------|
| Operation: | Continuous |

| Materials handled: | Feed 1 | Feed 2 | Feed 3 | Outlet 1 | |
|------------------------------|--------|--------|--------|----------|--|
| | S-122 | S-129 | S-125 | S-132 | |
| Quantity (lb/hr): | 4317 | 10398 | 4876 | 19591 | |
| Composition: (mass fraction) | | | | | |
| Glycerol | 0 | 0 | 0 | 0 | |
| Water | 0.114 | 0.535 | 0.156 | 0.348 | |
| Methanol | trace | 0.005 | 0.003 | 0.004 | |
| Hydrogen | trace | trace | trace | trace | |
| Propylene glycol | 0.881 | 0.455 | 0.838 | 0.644 | |
| N-propanol | 0.002 | 0.004 | 0.002 | 0.004 | |
| Isopropanol | 0.003 | 0.005 | 0.003 | 0.004 | |

| Design Data: | Pressure: 548 psia |
|--------------|----------------------------------------|
| | Outlet temperature: 317 F |
| | Vapor fraction: 0 |
| | Material of construction: Carbon steel |

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, separation section

Equipment base f.o.b. cost: \$3300 Bare Module Factor: 1 Bare Module Cost: \$3699

| PUMP | | | | | | |
|-----------------|-----------------------------------------|----------------------|---|--------------|-----------------|--|
| Identification: | Item Item No. No. required | <i>Pump</i> P-101 | 1 | Date: By: | 4-Apr-11 KMH | |

Function: To increase the pressure of the glycerol/water stream exiting EDI prior to HX.

| Operation: Continuous | | |
|------------------------------|-------|--------|
| Materials handled: | Inlet | Outlet |
| | S-105 | S-106 |
| Quantity (lb/hr): | 33231 | 33231 |
| Composition: (mass fraction) | | |
| Glycerol | 0.467 | 0.467 |
| Water | 0.516 | 0.516 |
| Methanol | 0.017 | 0.017 |

| Design Data: | Discharge pressure: 42 psia | | | |
|----------------------------------------|-----------------------------------|--|--|--|
| | Pressure change: 22 psi | | | |
| | Volumetric flow rate: 472 cuft/hr | | | |
| Material of construction: Stainless st | | | | |
| | Pump speed: 3600 RPM | | | |
| | Pump efficiency: 0.463 | | | |
| | Net work required: 1.629 hp | | | |
| | | | | |

Utilities: Electricity requirement: 0.283 kW

Controls:

Tolerances:

Comments and drawings: See process flow sheet, pretreatment section

Equipment base f.o.b. cost: \$3133 Bare Module Factor: 3.3 Bare Module Cost: \$15,097

| PUMP | | | | | | |
|-----------------|-----------------------------------------|---------------|---|--------------|------------------|--|
| Identification: | Item Item No. No. required | Pump P-102 | 1 | Date: By: | 30-Mar-11 KMH | |

Function: To increase the pressure of the water/methanol stream for feed to the EDI process.

| Operation: Continuous | | |
|------------------------------|-------|--------|
| Materials handled: | Inlet | Outlet |
| | S-111 | S-112 |
| Quantity (lb/hr): | 9230 | 9230 |
| Composition: (mass fraction |) | |
| Glycerol | trace | trace |
| Water | 0.960 | 0.960 |
| Methanol | 0.040 | 0.040 |

| Design Data: | Discharge pressure: 30 psia | | | |
|-------------------------------------|-----------------------------------|--|--|--|
| | Pressure change: 10.3 psi | | | |
| | Volumetric flow rate: 151 cuft/hr | | | |
| Material of construction: Stainless | | | | |
| | Pump efficiency: 0.296 | | | |
| | Net work required: 0.380 hp | | | |

Utilities:Electricity requirement: 0.283 kWControls:Tolerances:Comments and drawings: See process flow sheet, pretreatment section

Equipment base f.o.b. cost: \$3081 Bare Module Factor: 3.3 Bare Module Cost: \$14848

| PUMP | | | | | | |
|-----------------|-----------------------------------------|---------------|---|--------------|-----------------|--|
| Identification: | Item Item No. No. required | Pump P-103 | 1 | Date: By: | 4-Apr-11 KMH | |

| Function: | To increase the pressure of the glycero | ol/water stream for feed to the reactor. |
|-----------|-----------------------------------------|------------------------------------------|
|-----------|-----------------------------------------|------------------------------------------|

| Operation: Continuous | | |
|------------------------------|-------|--------|
| Materials handled: | Inlet | Outlet |
| | S-111 | S-112 |
| Quantity (lb/hr): | 19245 | 9230 |
| Composition: (mass fraction) | | |
| Glycerol | 0.806 | 0.806 |
| Water | 0.194 | 0.194 |
| Methanol | trace | trace |

| Design Data: | Discharge pressure: 587 psia | |
|--------------|-------------------------------------------|--|
| | Pressure change: 567 psi | |
| | Volumetric flow rate: 267 cuft/hr | |
| | Material of construction: Stainless steel | |
| | Pump efficiency: 0.378 | |
| | Net work required: 29.1 hp | |

Utilities: Electricity requirement: 21.7 kW Controls: Tolerances: Comments and drawings: See process flow sheet, reactor section

Equipment base f.o.b. cost: \$3100 Bare Module Factor: 3.3 Bare Module Cost: \$14,969
TURBINE

| Identification: | Item | Isentropic Turbine | | |
|-----------------|--------------|--------------------|-------|-----------|
| | Item No. | T-101 | Date: | 11-Apr-11 |
| | No. required | 1 | By: | KMH |

Function: To decrease the pressure of the inlet hydrogen to acceptable process pressure.

| Operation: Continuous | | |
|------------------------------|-------|--------|
| Materials handled: | Inlet | Outlet |
| | S-117 | S-118 |
| Quantity (lb/hr): | 346 | 346 |
| Composition: (mass fraction) | | |
| Hydrogen | 1 | 1 |

| Design Data: | Net work required: -27.7 hp | | |
|--------------|---------------------------------------|--|--|
| | Efficiency: 0.80 | | |
| | Discharge pressure: 600 psia | | |
| | Outlet temperature: 11.7 F | | |
| | Isentropic outlet temperature: -3.1 F | | |
| | Vapor fraction: 1.0 | | |
| | | | |

Utilities: Controls: Tolerances: Comments and drawings: See process flow sheet, pretreatment section.

Equipment base f.o.b. cost: \$7810 Bare Module Factor: 1 Bare Module Cost: \$8754 Section 15

FIXED CAPITAL INVESTMENT SUMMARY

Discussion of Total Permanent Investment (CTPI)

The total permanent investment for this project was calculated by summing a number of costs, including the bare-module investment, direct permanent investment and total depreciable capital. The bare-module cost was found to be approximately \$15.3 million, and it accounts for all on-site fabricated equipment and process machinery. The purchase cost for each piece of equipment was calculated using correlations provided in the text <u>Product and Process Design</u> <u>Principals</u> while the installed cost was determined using the factor-cost method. The cost of the catalyst was also included accounting for catalyst replacement over the course of the plant's projected life. It was found to be \$1.97 million.

The total permanent investment accounts for site preparation and service facilities like installing utility lines, building plant facilities and other similar tasks. In total, direct permanent investment was assumed to be 10% of the bare-module cost for equipment. After adding funds to account for the cost of contingencies and contractor fees (assumed to be 18% of the direct permanent investment), the total depreciable capital was found. This value is used in the financial analysis section to determine tax reductions for the company.

Finally, the cost of land and plant start-up are added to determine the unadjusted total permanent investment. Start-up costs were assumed to be 20% of the total depreciable capital. Since the plant will be located in the Gulf Coast, a site factor of 1.00 was used to determine the final total permanent investment, \$24.2 million.

| Investment Summary | | | | |
|------------------------------------------------------------|----|------------|----|------------|
| | | | | |
| Bare Module Costs | | | | |
| Fabricated Equipment | \$ | 10,412,750 | | |
| Process Machinery | \$ | 1,019,119 | | |
| Spares | \$ | 311,696 | | |
| Storage | \$ | 1,583,445 | | |
| Other Equipment | \$ | - | | |
| Catalysts | \$ | 1,972,383 | | |
| Computers, Software, Etc. | \$ | - | | |
| Total Bare Module Costs: | | | \$ | 15,299,392 |
| Direct Permanent Investment | | | | |
| Cost of Site Preparations: | \$ | 764,970 | | |
| Cost of Service Facilities: | \$ | 764,970 | | |
| Allocated Costs for utility plants and related facilities: | \$ | - | | |
| Direct Permanent Investment | | | \$ | 16,829,331 |
| | | | | |
| Total Depreciable Capital | | | | |
| Cast of Contingoncies & Contractor Foos | ¢ | 2 020 200 | | |
| Cost of Contingencies & Contractor Fees | φ | 3,029,200 | | |
| Total Depreciable Capital | | | ¢ | 10 050 611 |
| | | | φ | 17,030,011 |
| Total Permanent Investment | | | | |
| Cost of Land: | \$ | 397,172 | | |
| Cost of Royalties: | \$ | - | | |
| Cost of Plant Start-Up: | \$ | 3,971,722 | | |
| Total Permanent Investment - Unadjusted | | | \$ | 24,227,505 |
| Site Factor | | | | 1.00 |
| Total Permanent Investment | | | \$ | 24,227,505 |

Section 16

ENVIRONMENTAL CONSIDERATIONS

Environmental Considerations

The use of glycerol in propylene glycol manufacture will drastically decrease the negative environmental impact that is currently associated with the conventional methods of propylene glycol production. These conventional methods require propylene oxide as a feedstock. Propylene oxide is manufactured during the process of petroleum refinement; the manufacture of propylene oxide from crude oil reduces the availability of crude oil for refinement into gasoline and depletes natural resources, such as crude oil wells, that are already scarce. In contrast, glycerol is abundantly available as a byproduct from biodiesel manufacture. The amount of crude glycerol available is growing with the expansion of the biodiesel industry. By switching to this new method of propylene glycol production, the world's crude oil reserves will be conserved, and waste associated with biodiesel manufacture will be reduced.

According to their material safety data sheets, neither pure glycerol nor propylene glycol has significant environmental or health hazards, and both have a very low risk of flammability. As such, there are essentially no regulations on the transport and storage of industrial grade propylene glycol products. The large volume of hydrogen in the process is also essentially harmless to the environment, unless ignited.

The main environmental concerns in the glycerol to propylene glycol manufacturing process involve the disposal of the wastewater streams and the use of electricity for equipment operations. The entire plant will be drawing between 250 and 300 kW of power from the grid in operation. On the Gulf Coast, power is mainly supplied by oil power plants, which cause significant pollution and waste valuable resources. This negative impact could be offset by building and using a local power source such as solar panels on the roof of the plant. Government grants can be acquired to fund this endeavor. For example, Ford Motor Company

recently built a 500 kW solar plant on the roof of one of its manufacturing facilities in Michigan. This cost Ford less than \$3 million due to the government subsidies that it received.

The wastewater from the electrodeionization machine contains over 13% by mass of salt, which is approximately 4 times the salinity of the Gulf of Mexico. While not considered to be a pollutant, its disposal is non-negligible. The coastland on the Gulf of Mexico is mainly marshland, which is very sensitive to salinity changes. Disposing of such a highly concentrated saline solution in such an environment may have an enormous negative impact on the plant and animal species native to the area. The existing sewage treatment facilities are not equipped to remove these inorganic salts in large concentrations. Thus, this waste stream must be piped into the ocean far offshore, which is essentially a salt sink. As such, it is recommended that the construction of a pipeline to move the waste stream a safe distance from shore to dispose of the salt water should be strongly considered.

Two additional wastewater streams containing methanol and propanol must be considered. The methanol and propanol in these streams cause them to be toxic to the environment. These streams will be sent to the sewer authority for digestion. Since this is a commercial facility, sewage fees will be assessed by the sewer authority based on the quantities of water and organic compounds.

Finally, considerations must be made for the small pilot flame that will be used to burn excess hydrogen and propanol dissolved in the feed to the refinement column. Since propanol is easily combusted, the only expected products are water and carbon dioxide. With routine monitoring by plant personnel, it is not expected to pose significant environmental risks.

Section 17

HEALTH AND SAFETY CONSIDERATIONS

Health and Safety Considerations

The main safety issue in the plant's operation is the presence of hydrogen gas at high pressures. Hydrogen gas, a very small molecule, diffuses rapidly through many materials, so extreme caution must be exercised to avoid leakage into the plant. Hydrogen gas has an explosion limit between 4% and 47%. This is relatively low, and there is a risk of combustion in the presence of atmospheric air as the pressurized hydrogen gas escapes from the vessel.

Hydrogen embrittlement occurs when hydrogen adsorbs on certain metals and alloys causing a loss of ductility in the metal, making the equipment more brittle. This increases the likelihood of hydrogen leakage or large-scale breaks in the system, which could be catastrophic. To minimize the possibility of leakage and prevent embrittlement, process equipment that contains pressurized hydrogen uses stainless steel alloy, which is not susceptible to embrittlement.

To ensure that no dangerous buildup of hydrogen gas occurs, sensors will be installed in critical areas of the plant to monitor the presence of hydrogen in the air and alert operators of significant leaks. One target area for hydrogen leakage control is directly around the reactor vessel, which houses a large amount of pressurized hydrogen along with the liquid feed and products. Tight hydraulic fittings will be used throughout the process to minimize hydrogen leaks from the pipelines.

The heat of reaction of glycerol to propylene glycol varies slightly with temperature. It has a negligible effect due to the large volume of hydrogen gas that absorbs heat generated. The reactor is therefore assumed to operate adiabatically.

The large amount of electricity that will be drawn by the EDI machine requires appropriate insulation and warning systems to avoid the risk of electric shock, short circuiting, or fire. Sufficient insulation on all electric cables is required and close monitoring of electrical systems is required to swiftly detect any anomilies and possible short circuits. Any electric cable that is used must be checked periodically to prevent fraying or wire exposure, and in the event of damage, the cable must be replaced immediately. Smoke alarms must be installed throughout the plant to ensure rapid detection of and emergency response to any fires, which could arise from the electrical connectivity, hydrogen leakage or other accidental combustion. Fire extinguishers must be easily accessible by personnel as mandated by the municipality's fire safety inspector.

To ensure that there are no interruptions in the EDI function, a generator that can provide the power required for 12 hours of plant operation should be available in case of power outage. Such an interruption could cause immense damage to the propylene glycol product's purity and quality, imposing a large financial cost and potentially endangering consumers.

Product safety is a major concern with regard to USP grade propylene glycol, which is used widely in pharmaceutical and food products. The propylene glycol from this plant will be 99.99% pure, which must be guaranteed to avoid the risk of poisoning consumers. There is a chance that trace amounts of ethylene glycol could be produced in a side reaction during production. Consumption of ethylene glycol is hazardous to the kidneys, liver, and central nervous system of humans and is potentially fatal, according to its Material Safety Data Sheet. It also has mutagenic effects on some types of mammalian cells and is a known carcinogen. Therefore, it is very important that the product not be sold containing any ethylene glycol. Samples will be analyzed every 30 minutes with a gas chromatograph to ensure that trace

quantities of ethylene glycol are not present in the USP grade product. This is especially

important during startup (and in the early stages of the process).

Section 18

STARTUP CONSIDERATIONS

Startup Considerations

The majority of the analysis done on this process considers only steady state operation of the plant. However, certain considerations must be made for the startup period prior to reaching steady state. Specific startup considerations must be considered for the major equipment units, including both distillation columns, the reactor, and the EDI machine, as well as the hydrogen recycle loop and its integration into the system.

Initially, process-clean water will be run through all of the major process units, including the EDI machine, in order to clean out the system. Once all process units except for the EDI machine have been emptied of water, crude glycerol will be slowly introduced into the process and the EDI machine will begin to desalt the glycerol. Once the glycerol attains an acceptable level of salt content to remove the risk of corrosion, it will be sent to column D-101, which will operate at total reflux and use excess steam utility in the reboiler until the column temperature profile is achieved. The reflux ratio will then be adjusted to reach the process specification of a molar reflux ratio of 0.2, and the distillate will then start to heat the column feed in HX-101. This will reduce utility usage in the reboiler until the steady state value of 15700000 Btu/hr is achieved. The bottoms product of refined glycerol will be collected and analyzed until it meets the required specifications of 10 ppm salt content and 15% water by mass. Once these specifications are reached, this stream will be connected to the reactor section.

To avoid combustion when the pressurized hydrogen is introduced to the process, pipelines that will contain hydrogen must be flushed with nitrogen gas to remove oxygen. Once the hydrogen has been introduced and begins circulating through the reactor, the refined glycerol will slowly be introduced. Additionally, due to the extensive heat integration, startup heaters

will be required to raise the temperature of the hydrogen and glycerol streams that enter the reactor to the reactor inlet temperature of 410 F. Additional cooling will be provided by C-102 until the heat exchanger network begins to operate.

Once the liquid reactor effluent is collected, it will be sent to column D-102, which will initially operate at total reflux in a manner similar to D-101. Once the temperature profile is achieved, product will start to be collected and analyzed until it meets specifications.

Section 19 PLANT LAYOUT

Plant Layout

Because the process requires only one tanker truck per hour of both product and feed, the plant does not necessarily need to have a rail connection, although it would increase efficiency. The only requirement for plant layout is that D-102 needs to not be covered by the factory roof since there will be a hydrogen flame coming off the condenser. Additionally, the EDI machine must be protected from the elements due to the sensitive nature of the electrodes. Aside from that, this layout was chosen to reduce pipeline length and to ensure that utilities are in proximity of process equipment that requires them (i.e. electricity connection near EDI, hydrogen near the reactor, and heating and cooling utilities near the distillation columns).



Figure 8 Plant layout diagram

Section 20

OPERATING COSTS AND ECONOMIC ANALYSIS

Operating Costs and Economic Analysis

A number of assumptions were made during the financial analysis process that led to the values presented above. The plant will be located in the Gulf Coast and will operate 330 days per year resulting in an overall operating factor of 0.9041. Two years are allocated for design and construction of the plant (2012, 2013) and operation is expected to begin in 2014. To account for a typical efficiency lag, production will begin at 50% efficiency and ultimately reach 90% efficiency in the third year of production (2016). A 5 year MACRS depreciation schedule was chosen for this project to remain consistent with financial analysis of projects of a similar scale.

A number of utilities will be required to run the plant, including medium pressure steam, process water, cooling water, electricity and treatment of waste streams. Of these utilities, electricity and organic waste water treatment are the most significant costs at \$0.06/kWh and \$0.048/lb respectively. In addition to the crude glycerol feed, the hydrogen feed will also be purchased at \$0.50/lb. Although n-propanol and isopropanol are being created as byproducts of the reaction of glycerol to propylene glycol, they are produced in such small quantities that it is more profitable to simply dispose of them as waste instead of investing in additional purification equipment. Summing the raw material costs and utilities with the general expenses yields a total variable cost of \$45.1 million.

Based on the proposed design, the process requires 4 operators per shift, while assuming each operator will have 5 shifts per week. The EDI pretreatment section, reactor section and separation section will each need one dedicated operator while a fourth operator will be charged with managing the hydrogen recycle system and storage systems for the waste and product

steams. The EDI equipment is expected to be licensed from Dow, formerly Rohm and Haas, for \$2.00 million per year. In total, the fixed costs for per year will equal \$7.16 million.

The total bare module cost is projected to be \$15.3 million, which includes the process equipment, catalyst, storage equipment and spares. Accounting for multiple recycle streams EDI integration, 20% of total depreciable capital was allocated to plant start-up costs. The total permanent investment is found to be \$22.2 million and will be completely accounted for in the cash flow during the year of construction, 2013. Working capital is projected based on 14 days in inventory for the raw materials and final product as well as 30 days of accounts receivable and cash reserves. When summed with the \$11.8 million in working capital, the total capital investment was found to be \$34.0 million.

Analysis of the resulting cash flows shows that the first year of cumulative positive net present value occurs in 2016. Variable costs for the year total \$40.5 million while fix costs total \$6.60 million. Clearly, the calculated NPV value is heavily weighted by variable costs such as raw material prices and utilities. Thus, a sensitivity analysis will focus on these factors to understand the overall volatility of the process profitability.

As discussed in the Market Analysis section of the report, the production of propylene glycol from renewable biodiesel represents a rapidly growing market that has the potential for high profitability. Using techniques and correlations presented in <u>Product and Process Design</u> <u>Principles</u> (Seider *et al.* 2009), the proposed process was found to be highly profitable, as expected. The base case scenario projected a net present value (NPV) of \$88.4 million in 2012 with an investor's rate of return (IRR) of 58.54%. These high values are primarily driven by the

large margin between the selling price of propylene glycol and the cost of the crude glycerol feed, \$1.00/lb and \$0.22/lb respectively.

Sensitivity Analysis

As previously mentioned, a number of key factors were identified that could have a significant impact on the overall profitability. The factors include commodity costs such as the price of propylene glycol, glycerol and hydrogen feed prices and electricity costs in addition to variable costs such as organic waste water treatment and EDI licensing fees. The sensitivity-plot shown in Figure 9 is a graphical representation of the sensitivity analysis on these variables. This graph plots the percent change in a given factor against the change in IRR relative to the base case, 58.5%. Thus, a ten unit increase on the y-axis corresponds to an IRR of 68.5%.



Figure 9 Sensitivity Graph

The commodity costs were varied by plus and minus 15% to understand the profitability relationship to tangential markets. In spite of our best efforts, we were not able to confidently estimate the EDI licensing costs; therefore, this factor was varies by plus or minus 30% to account for the increased uncertainty.

Propylene Glycol (PG) Price Sensitivity

The plus or minus 15% uncertainly in spot propylene glycol prices appears to be an accurate representation of the uncertainty in the market. According to a recent ICIS, a subsidiary of Reed Business Information and trusted information provider for the chemical and oil industry, propylene glycol prices in the US were \$0.93-\$1.05/lb in January 2011, up from \$0.88-\$0.96/lb in November 2010. The unusually harsh winters of the past 3 years has increased the demand for antifreeze and plane de-icing fluid, and as global mean temperatures continue to rise, the increased moisture in the atmosphere will result in greater snowfall. Prices are projected to potential increase another \$0.15/lb in the first half of 2011 due to a seasonal spike in demand (ICIS 2011) Furthermore, an increase in natural gas prices increased the price of the percent of propylene glycol market derived propylene, a product of natural gas. However, these price points could easily relax as the artificial boost due to seasonal volatility will not continue into the later spring and early summer months.

In general, the demand for propylene glycol has been steady at 2%, with a yearly growth of about 1.5% in the unsaturated polyester resins (UPRs) and about 2.5% in the cosmetics and consumer products industry. In Asia, demand for propylene glycol has been estimated as growing about 6% per year (ICIC 2011). Coupled with the expected increase in supply of crude glycerol in the coming years, and thus lower feed prices, it is reasonable to expect that the

increased demand for propylene glycol will be offset by increased supply of feed. Thus, the price of propylene glycol likely will remain relatively stable within a plus or minus 15% swing.

As shown in Figure 9, the IRR is the most sensitive to volatility in PG prices relative to the other six factors studied. Figure 10 shows that a 15% decrease in PG prices to \$0.85/lb could decrease the IRR as much as 15% to 43.5%. However, the current process is still exceedingly profitable and is able to absorb a commodity price shock of this magnitude. In fact, assuming constant variable costs and working capital, the price of propylene glycol would have to fall to \$0.68/lb to make the project just barely viable at 20.67% IRR. With the expected increase it PG production in the coming years as well as a potential decrease in price due to competition in a number of the chemical's largest markets, this could be a very possible scenario in the coming decade; however, it is unlikely to happen in the short-term five year horizon.



Figure 10 Sensitivity of PG price.

Glycerol (feed) Price Sensitivity

As previously mentioned, crude glycerol is a natural by-product of biodiesel production. For every 9 kg of biodiesel produced, about 1 kg of glycerol is formed (Suppes 2006). With US production of glycerol exclusively from biodiesel projected to be 1.92 billion pounds in 2012, up from 281 million lbs in 2007, the price of crude glycerol is expected to be significantly depressed from the estimate of \$0.22/lb used in this financial assessment (Gobina 2007). Gobina goes on to project crude glycerol prices to fall as low as \$0.05/lb, citing the fact that some biodiesel producers find it more profitable to burn it as waste than attempt to sell it on the open market.

However, recent technologies have open up alternative markets for excess glycerol that may help curb over-supply. Solvay Chemicals commercialized a process to reverse the conventional route of making synthetic glycerol from epichlorohydrin. After the success of their pilot plant in 2007, a full scale commercial plant is scheduled to be built in Map Tha Phut, Thailand. Alkylene carbonate (reactive intermediate or solvent), biomethanol, and acrylic acid are additional examples of potential products produced from glycerol feed material. (Chemical Business 2010). Furthermore, new purification systems, such as EDI, offer alternatives to cost and labor intensive vacuum distillation, which could further equalize glycerol supply and demand.

The price of glycerol feed, being one of the two primary inputs to the process, obviously has a significant effect on the overall profitability. Using a very conservative estimate of a 15% decrease in the price of crude glycerol to \$0.187/lb, the IRR increases 5.3% to 63.8% as shown in Figure 11. If the current spot price of glycerol, \$0.10/lb, is used, the IRR increases to 76.5%. The greater the drop in crude glycerol price, the larger the margin for propylene glycol and thus the greater the overall profitability for the process.



Figure 11 Sensitivity of glycerol feed price.

Organic Waste Water Treatment Sensitivity

Waste water from this process will contain methanol, n-propanol and iso-propanol. The cost of water treatment was calculated using a correlation provided by Bruce Vrana from DuPont:

$$Cost \left[\frac{\$}{lb}\right] = 0.044 + 0.048 * (\% \text{ organic impurities by weig }ht)$$

It is understood that this correlation is an estimate that can be used to quantify this cost; however, the actual value could vary based on geography, market conditions and quantity of wastewater. A 10% increase in waste water treatment costs corresponds to a 0.64% increase in IRR. However, this variable represents a moderate contribution to variable costs and thus is not a critical concern based on the economic analysis.



Figure 12 Sensitivity of waste water treatment price.

EDI Licensing

EDI Licensing is assumed to be \$2.00 million per year. This estimate has significant uncertainty as we were unable to obtain an exact quote for a process of this scale. As a result, a larger sensitivity analysis was performed, varying the licensing fee by 30% in both directions. Relative to the commodity costs of PG and glycerol, the licensing fee does not have as critical of an impact on the overall profitability, providing confidence that this is an economically viable
technology to incorporate into the process. A 30% increase in licensing fees for a total of \$2.6 million per year would decrease the IRR by 1.17% to 56.9%.



Figure 13 Sensitivity of EDI licensing cost.

Hydrogen Sensitivity

Hydrogen will be supplied to the plant via direct pipeline from the supplier at a cost of \$0.50/lb. Although hydrogen will be conserved via recycle streams to reduce utility usage, a significant amount will be needed during the start-up process to fill the system. The price of hydrogen is historically constant, and our analysis shows that a plus or minus 15% fluctuation would minimally affect IRR.



Figure 14 Sensitivity of hydrogen price.

Electricity Sensitivity

Electricity was projected to cost \$0.06/kWh, based on information provided in Process Design Principles. The EDI pre-treatment section adds a significant demand for electricity for the process, approximately 200kW, resulting in an overall electricity need of 241kW. Since the plant is designed to be environmentally friendly with a minimal carbon footprint, it is possible that electricity will be sought from renewable sources for a higher fee. There already exists a significant network of wind energy generation facilities in the local area, and the recent British Petroleum Oil spill will likely encourage further development of renewable energy sources in the region. (Save Energy 2004) Assuming a potential surcharge of up to 15% would cause a decrease in IRR by 0.28%. The high profitability of the current process would justify a marginal surcharge in sourcing electricity from green sources.



Figure 15 Sensitivity of Electricity price.

Summary of Sensitivity Analysis

The slope of the linear regressions found for each line indicates the relative influence on the IRR of the project. As shown in Figure 9, the price of PG and glycerol are by far the most significant levers that determine the overall profitability. Based on market analysis, the price of PG is expected to seasonally fluctuate but trend downward over the next decade. However, the price of crude glycerol is also expected to decrease significantly ensuring the next margin will remain. From a financial point of view, this process warrants serious consideration for implementation.

| Rank | Factor | Slope | Absolute Value of Slope |
|------|---------------------|---------|-------------------------|
| 1 | PG Price | 0.9514 | 0.9514 |
| 2 | Glycerol Feed Price | -0.3533 | 0.3533 |
| 3 | Organic Waste Water | -0.0626 | 0.0626 |
| 4 | EDI Licensing | -0.0385 | 0.0385 |
| 5 | Hydrogen Price | -0.0203 | 0.0203 |
| 6 | Electricity | -0.0017 | 0.0017 |

Figure 16 Ranked factors based on sensitivity.

General Information

Process Title:Glycerol to Propylene Glycol Product:Propylene Glycol Plant Site Location:Gulf Coast Site Factor:1.00 Operating Hours per Year:7920 Operating Days Per Year:330 Operating Factor:0.9041

Product Information

This Process will Yield

12,379lb of Propylene Glycol per hour 297,096lb of Propylene Glycol per day 98,041,680lb of Propylene Glycol per year

Price

\$1.00/lb

Chronology

| | | Distribution of | Production | Depreciation | Product Price |
|-----------------|--------|----------------------|------------|---------------------|---------------|
| <u>Year</u> | Action | Permanent Investment | Capacity | 5 year MACRS | |
| 2012Design | | | 0.0% | | |
| 2013Constructio | n | 100% | 0.0% | | |
| 2014Production | | 0% | 45.0% | 20.00% | \$1.00 |
| 2015Production | | 0% | 67.5% | 32.00% | \$1.00 |
| 2016Production | | 0% | 90.0% | 19.20% | \$1.00 |
| 2017Production | | | 90.0% | 11.52% | \$1.00 |
| 2018Production | | | 90.0% | 11.52% | \$1.00 |
| 2019Production | | | 90.0% | 5.76% | \$1.00 |
| 2020Production | | | 90.0% | | \$1.00 |
| 2021Production | | | 90.0% | | \$1.00 |
| 2022Production | | | 90.0% | | \$1.00 |
| 2023Production | | | 90.0% | | \$1.00 |
| 2024Production | | | 90.0% | | \$1.00 |
| 2025Production | | | 90.0% | | \$1.00 |
| 2026Production | | | 90.0% | | \$1.00 |
| 2027Production | | | 90.0% | | \$1.00 |
| 2028Production | | | 90.0% | | \$1.00 |

Equipment Costs

| Equipment Description | | Bare Module Cost |
|-----------------------|----------------------|------------------|
| M-102 | Process Machinery | \$3,699 |
| S-101 | Fabricated Equipment | \$4,483,200 |
| P-101 | Process Machinery | \$13,993 |
| HX-101 | Fabricated Equipment | \$77,519 |
| H-101 | Fabricated Equipment | \$222,840 |
| D-101 | Fabricated Equipment | \$1,166,781 |
| D-101a | Process Machinery | \$81,770 |
| D-101b | Process Machinery | \$37,593 |
| D-101c | Process Machinery | \$37,003 |
| D-101d | Process Machinery | \$141,198 |
| P-103 | Process Machinery | \$15,097 |
| Catalyst | Catalysts | \$1,972,383 |
| P-102 | Process Machinery | \$14,847 |
| R-101a | Fabricated Equipment | \$796,642 |
| R-101b | Fabricated Equipment | \$796,642 |
| F-101 | Process Machinery | \$111,058 |
| HX-102 | Fabricated Equipment | \$67,135 |
| F-102 | Process Machinery | \$89,031 |
| HX-103 | Fabricated Equipment | \$67,421 |
| C-102 | Fabricated Equipment | \$85,504 |
| F-103 | Process Machinery | \$31,808 |
| CP-101 | Process Machinery | \$258,602 |
| TR-100 | Process Machinery | \$57,089 |
| M-101 | Process Machinery | \$20,233 |
| M-103 | Process Machinery | \$3,699 |
| V-101 | Process Machinery | \$11,208 |
| D-102 | Fabricated Equipment | \$2,477,126 |
| D-102a | Fabricated Equipment | \$23,458 |
| D-102b | Process Machinery | \$37,258 |
| D-102c | Process Machinery | \$45,179 |
| D-102d | Fabricated Equipment | \$72,944 |
| C-103 | Fabricated Equipment | \$75,539 |
| T-101 | Storage | \$542,691 |
| T-102 | Storage | \$313,824 |
| T-103 | Storage | \$100,029 |
| T-103 | Storage | \$56,322 |
| T-104/T-105 | Storage | \$125,967 |
| T-106 | Storage | \$444,611 |
| Additional Equipment | | \$320,449 |

<u>Total</u>

\$15,299,392

| Raw | Materials | | | | | | |
|--------|-------------------------------------------------------------|--------------------------------|-------------------------------------|--------------------------------------------|-------------------------------------------|----------------|-----------------------------------------------------------------------------------------|
| | <u>Raw Material:</u> Crude Glycerol Water Hydrogen | <u>Unit:</u> Ib Ib Ib | Required Ratio: | 1.2287lb p 0.052407lb p 0.027948lb p | er lb of PG er lb of PG er lb of PG | | <u>Cost of Raw</u> <u>Material:</u> \$0.220per lb \$0.00per lb \$0.50per lb |
| | Total Weighted Average: | | | | | | \$0.284per lb of PG |
| Utilit | ie | | | | | | |
| S | | | | | | | |
| | <u>Utility:</u> Med Pressure | <u>Unit:</u> | Required Ratio | | | - | <u>Utility</u> <u>Cost</u> \$3,000E- |
| | Steam | lb | | 2.515464lb p | er lb of PG | | 03per lb \$7.500E- |
| | Process Water | gal | | 0.049633gal | per lb of PG | | 04per gal \$7.500E- |
| | Cooling Water | lb | | 71.1lb p | er lb of PG | | 05per lb per |
| | Electricity Organic Waste | kWh | | 0.0194743kWl | n per lb of PG | i | \$0.060kWh |
| | water Salt Waste Water | di Ib | | 0.9556lb p 0.474637lb p | er ib of PG er ib of PG | | \$0.048per lb \$0.00per lb |
| | Total Weighted Average: | | | | | | \$0.060per lb of PG |
| Varia | able Costs | | | | | | |
| | General Expenses: | <u>.</u> | | _ | | | |
| | | | Selling / Transfe | er Expenses: | 3.00%of | Sales | |
| | | | Dire | ct Research: | 4.80%of | Sales | |
| | | | Allocate Administrati Managom | ed Research: ive Expense: | 0.50%of 2.00%of | Sales Sales | |
| | | | CC | mpensation: | 1.25%of | Sales | |
| Worl | king Capital | | | | | | |
| | Accounts Receivable Cash Reserves (exc | e Iuding Raw | | ₽ | 30 | Days | |
| | Materials) | | | ₽ | 30 | Davs | |
| | , Accounts Payable | | | ₽ | 0 | Davs | |
| | Propylene Glycol Inv | ventory | | ₽ | 14 | Davs | |
| | Raw Materials | 5 | | ₽ | 14 | Days | |

| Total Permanent Investmer | nt | |
|-----------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| | Cost of Site Preparations: Cost of Service Facilities: Allocated Costs for utility plants and related facilities: Cost of Contingencies and Contractor Fees: Cost of Land: Cost of Royalties: Cost of Plant Start-Up: | 5.00% of Total Bare Module Costs 5.00% of Total Bare Module Costs \$0 18.00% of Direct Permanent Investment 2.00% of Total Depreciable Capital \$0 20.00% of Total Depreciable Capital |
| Fixed Costs Operations | Operators per Shift: Direct Wages and Benefits: Direct Salaries and Benefits: Operating Supplies and Services: Technical Assistance to Manufacturing: Control Laboratory: | 4(assuming 5 shifts) \$40/operator hour 15%of Direct Wages and Benefits 6%of Direct Wages and Benefits \$0.00per year \$0.00per year |
| <u>Maintenance</u> | Wages and Benefits: Salaries and Benefits: Materials and Services: Maintenance Overhead: | 4.50%of Total Depreciable Capital 25%of Maint. Wages + Benefits 100%of Maint. Wages + Benefits 5%of Maint. Wages + Benefits |
| Operating Overhead | General Plant Overhead: Mechanical Department Services: Employee Relations Department: Business Services: | 7.10%of Maint and Ops Wages and Benefits 2.40%of Maint and Ops Wages and Benefits 5.90%of Maint and Ops Wages and Benefits 7.40%of Maint and Ops Wages and Benefits |
| Property Taxes and Insura | nce Property Taxes and Insurance: | 2% of Total Depreciable Capital |
| Straight Line Depreciation Direct Plant: Allocated Plant: | 8.00%of Total Depreciable Capital, less 1.18 times the All 6.00%of 1.18 times the Allocated Costs for Utility Plants | located Costs for Utility Plants and Related Facilities and Related Facilities |
| Other Annual Expenses | Rental Fees (Office and Laboratory Space): Licensing Fees:\$ Miscellaneous: | \$0 2,000,000 \$0 |
| Depletion Allowance | Annual Depletion Allowance: | \$0 |

Variable Cost Summary

Variable Costs at 100% Capacity:

General Expenses

| | Selling / Transfer Expenses: | \$ | 2,941,250 |
|------------------------|--------------------------------------|-------|---------------|
| | Direct Research: | \$ | 4,706,001 |
| | Allocated Research: | \$ | 490,208 |
| | Administrative Expense: | \$ | 1,960,834 |
| | Management Incentive Compensation: | \$ | 1,225,521 |
| Total General Expenses | | \$ | 11,323,814 |
| Raw Materials | \$0.284293per lb of Propylene Glycol | | \$27,872,536 |
| Byproducts | \$0.000000per lb of Propylene Glycol | | \$0 |
| <u>Utilities</u> | \$0.060025per lb of Propylene Glycol | | \$5,884,956 |
| Total Variable Costs | <u>_\$</u> | 45,08 | <u>31,306</u> |
| Fixed Cost Summary | | | |

Operations

| | Direct Wages and Benefits | \$ 1,664,000 |
|--------------------|---------------------------------------|-----------------|
| | Direct Salaries and Benefits | \$ 249,600 |
| | Operating Supplies and Services | \$ 99,840 |
| | Technical Assistance to Manufacturing | \$ - |
| | Control Laboratory | \$ - |
| | Total Operations | \$ 2,013,440 |
| Maintenance | | |
| | Wages and Benefits | \$ 893,637 |
| | Salaries and Benefits | \$ 223,409 |
| | Materials and Services | \$ 893,637 |
| | Maintenance Overhead | \$ 44,682 |
| | Total Maintenance | \$ 2,055,366 |
| Operating Overhead | | |
| | General Plant Overhead: | \$ 215,176 |
| | Mechanical Department Services: | \$ 72,736 |
| | Employee Relations Department: | \$ 178,808 |
| | Business Services: | \$ 224,268 |
| | Total Operating Overhead | \$ 690,987 |

| Property Taxes and Insurance | | | | | |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------|----------------------|---------------------------------------------------------------|-------------------------------------------------------------------------------|--------------------|
| | Property Taxes and Insurance: | | \$ | 397,172 | |
| Other Annual Expenses | | | | | |
| | Rental Fees (Office and Laboratory Space): Licensing Fees: Miscellaneous: | | \$ \$ \$ | - 2,000,000 - | |
| | Total Other Annual Expenses | | \$ | 2,000,000 | |
| Total Fixed Costs | | | \$ | 7,156,966 | |
| Investment Summary | | | | | |
| Bare Module Costs | | | | | |
| Fabricated Equipment Process Machinery Spares Storage Other Equipment Catalysts Computers, Software, Etc. <u>Total Bare Module Costs:</u> <u>Direct Permanent Investment</u> Cost of Site Preparations: Cost of Service Facilities: Allocated Costs for utility plan | | \$ \$ \$ \$ \$ \$ \$ | 10,412, 1,019, 311, 1,583, 1,972, 764, 764, | 750 119 696 445 - 383 - <u>\$</u> 970 970 | <u>15,299,392</u> |
| Direct Permanent Investment | | Φ | | <u></u> | 16,829,331 |
| Total Depreciable Capital | | | | | |
| Cost of Contingencies & Contr | ractor Fees | \$ | 3,029, | 280 | |
| Total Depreciable Capital | | | | \$ | <u> 19,858,611</u> |
| Total Permanent Investment | | | | | |
| Cost of Land: Cost of Royalties: Cost of Plant Start-Up: | | \$ \$ \$ | 397, 3,971, | 172 - 722 | |
| Total Permanent Investment - Site Factor | Unadjusted | | | \$ | 24,227,505 1.00 |
| Total Permanent Investment | | | | \$ | 24,227,505 |

Working Capital

| | ¢ <u>20</u> | <u>13</u> | <u>2014</u> | <u>2015</u> |
|----------------------------|----------------------|--------------------------|--------------|-------------|
| Accounts Receivable | ₃ 3,626,199 \$ | ↓ 1,813, \$ | 100 \$ | 1,813,100 |
| Cash Reserves | 408,400 | 204,20 | 00 \$ | 204,200 |
| Accounts Payable | \$ \$ | - \$ \$ | - \$ | - |
| Propylene Glycol Inventory | 1,692,226 \$ | 846,1 ⁻ \$ | 13 \$ | 846,113 |
| Raw Materials | 481,088 | 240,54 | 44 \$ | 240,544 |
| Total | \$ 6,207,913 | \$ 3,103, | 956 \$ | 3,103,956 |
| Present Value at 15% | \$ | 5,398,185\$ | 2,347,037 \$ | 2,040,902 |
| Total Capital Investment | | <u>\$</u> 34,013 | 3,629 | |

| Cash Flow Summary | Depletion Cumulative Net | osts Working Capital Var Costs Fixed Costs Depreciation Allowance Taxible Income Taxes Net Earnings Cash Flow Present Value at 15% | 27,500 (6,207,900) (30,435,400) (26,465,600) | (3,104,000) (20,286,600) (7,157,000) (3,971,700) - 12,703,500 (4,700,300) 8,003,200 8,871,000 (19,757,900) | - (3,104,000) (30,429,900) (7,157,000) (6,354,800) - 22,236,500 (8,227,500) 14,009,000 17,259,800 (8,409,300) | - (40,573,200) (7,157,000) (3,812,900) - 36,694,500 (13,577,000) 23,117,500 26,930,400 6,988,300 | - (40,573,200) (7,157,000) (2,287,700) - 38,219,700 (14,141,300) 24,078,400 26,366,100 20,096,900 | (40,573,200) (7,157,000) (2,287,700) - 38,219,700 (14,141,300) 24,078,400 26,366,100 31,495,700 | - (40,573,200) (7,157,000) (1,143,900) - 39,363,500 (14,564,500) 24,799,000 25,942,900 41,248,600 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,519,600 49,591,000 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,548,300 56,845,300 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,519,600 63,153,300 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,519,600 68,638,600 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,519,600 73,408,400 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,519,600 77,556,100 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,519,600 81,162,700 | - (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 25,519,600 84,298,900 | - 12,415,800 (40,573,200) (7,157,000) - 40,507,400 (14,987,700) 25,519,600 37,935,500 88,352,900 |
|-------------------|--------------------------|------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------|------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Cash Flow 5 | | king Capital Var Costs Fixed Cost | (6,207,900) | (3,104,000) (20,286,600) (7,157,0 | (3,104,000) (30,429,900) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | - (40,573,200) (7,157,0 | 12,415,800 (40,573,200) (7,157,0 |
| | | Capital Costs Working Car | - (24,227,500) (6,20] | ,800 - (3,10, | ,100 - (3,10 | ,500 - | .,500 | . 500 | ,500 - | ,500 - | .500 | .,500 | ,500 - | ,500 - | .500 | .500 | ,500 - | .500 - 12,415 |
| | Product Unit | <u>1y Price Sales</u> | | \$1.00 44,118, | \$1.00 66,178, | \$1.00 88,237, | \$1.00 88,237, | \$1.00 88,237, | \$1.00 88,237, | \$1.00 88,237, | \$1.00 \$8,237, | \$1.00 88,237, | \$1.00 \$8,237, | \$1.00 88,237, | \$1.00 88,237, | \$1.00 88,237, | \$1.00 88,237, | \$1.00 88,237, |
| | Percentage of | Year Design Capacit | 2013 0% | 2014 45% | 2015 68% | 2016 00% | 2017 🕄 | 2018 90% | 2019 90% | 2020 90% | 2021 90% | 2022 90% | 2023 90% | 2024 90% | 2025 90% | 2026 90% | 2027 90% | 2028 90% |

| Touriadinity Vincestates S& 45% The Internal Rate of Return (RR) for this project in 2012 is \$ 88.332,000 The Net Present Value (VPO) of this project in 2012 is \$ 88.332,000 Amaud Sales \$ 83.33,512 Amaud Sales \$ 83.33,512 Amuad Casis \$ 83.33,512 Amuad Sales \$ 84.33,311 Amuad Sales \$ 84.33,311 Amuad Sales \$ 84.33,31 Amuad Sales \$ 84.33,31 Amuad Sales \$ 84.33,31 <td< th=""><th>Drof</th><th>Hobility Moscur</th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th></td<> | Drof | Hobility Moscur | | | | | | | | | | |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------|-------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|------------------------------|------------------------------|------------------------------|------------------------------|------------------------------|------------------------------|------------------------------|------------------------------|------------------------------|
| The Net Present Value (NPV) of this project in 2012 is S 8.332,000 COI Analysis (Third Production Year) COI Analysis (Third Production Year) Annual Sales 88.2375(1) Annual Coles 48.431 | The Ir | Internal Rate of Return | n (IRR) for this | project is | | | 58.45% | | | | | |
| Col Analysis (Third Production Year) Col Analysis (Third Production Year) Connal Sials 83335121 Amual Cosis (1732014) Amual Cosis (1732014) Depreciation (1382047) Colori Capital Investment 24.245517 Astribit 24.245517 Depreciation (1382047) Colori Capital Investment 24.245517 Colori Capital Investment 24.245517 Colori Capital Investment 43.175 Depreciation (1382046) Colori Capital Investment 54.1975 Colori Capital Investment 54.1975 Statis 20.956145 54.0573.175 Statis 21.956 54.0573.175 Variation 20.556 54.1073.567 56.4054.608 54.1073.667 Statin Statin Statis 10.756 <td< td=""><td>The N</td><td>vet Present Value (NF</td><td>vV) of this proj</td><td>ect in 2012 is</td><td></td><td></td><td>\$ 88,352,900</td><td></td><td></td><td></td><td></td><td></td></td<> | The N | vet Present Value (NF | vV) of this proj | ect in 2012 is | | | \$ 88,352,900 | | | | | |
| Amual Sales 88 237,512 Amual Casts (1730,141) (1720,141) Deprectation forome fait be Lemings (1932,014) (1920,012) (1932,014) (1920,012) Deprectation for the Lemings (1932,014) (1920,012) (1932,014) (1920,012) Deprectation for the Lemings (1920,014) (1920,012) (1920,014) (1920,012) Sensitivity Analy Coll Capital Investment (1920,014) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) Sensitivity Analy (1920,012) 6433,31 (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) Sensitivity Analy (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) (1920,012) Deprecision (111,112) (192,015) (112,012) (1920,012) (1920,012) (1920,012) Deprecision (111,112) (192,015) (112,012) (112,012) (112,012) (112,012) (112,012) Deprecision (111,112) (192,016) (112,012) (1920,016) (112,012) (112,012) (112,012) (112,012) Deprecision (111,112) (112,012) (112,012) | ROI A | Analysis (Third Produ | iction Year) | | | | | | | | | |
| Sensitivity Analyses Analyses Analyses Sector below takes quite a bit of memory to update each time a cell is changed: therefore, automatic calculations are turned off. After making your axis selections, press "FPT" Note: The Sensitivity Manalyses Uvary Initial Value by 4t. Vary Initial Value by 4t. Variable Costs Variable 50% 50% Variable Variable Costs Variable Costs statis 50% 731,556,914 330,65,045 540,57,3175 540,591,357 540,579,357 561,075,57 558,065,098 563,113,828 561,075 563,065,098 563,113,828 561,075,57 558,065,098 563,113,828 567,67 70,075,57 558,065,098 563,113,828 567,67 70,075,57 558,065,098 563,113,828 567,67 70,075,57 558,065,098 563,113,828 567,67 563,113,828 567,67 563,113,828 567,675 561,075,57 558,065,098 563,113,828 561,075,57 558,065,098 563,113,828 561,075,57 558,065,098 563,113,828 561,075,57 558,065,098 563,113,828 561,075,57 558,065,098 563,113,828 561,075,57 | An De Ne RO RO | mual Sales nual Costs prectation come Tax t Earnings tal Capital Investment 1 | 88,237,512 (47,730,141 (1,938,200) (14,270,593, 24,298,577 66,31% | 36,643,331 | | | | | | | | |
| Vary Initial Value by -t. Vary Initial Value by -t. x-axis 50% y-axis 80% x-axis 80% y-axis 80% x-axis 80% | Sen Vote: | Isitivity Analyse: The Sensitivity Analyses R values. (These two lin | S section below ta es may be delete | kes quite a bit of m | hemory to update (| each time a cell is | changed; therefor | re, automatic calcu | ulations are turned | l off. After making | your axis selection | ns, press "F9" t |
| Variable Costs Solution (55) \$27,048,784 \$31,556,914 \$36,065,045 \$40,531,35 \$49,589,436 \$54,097,567 \$58,605,698 \$63,113,828 \$67,621,959 \$50,046 \$36,065,045 \$40,531,35 \$49,589,436 \$54,097,567 \$58,605,698 \$63,113,828 \$67,621,959 \$50,046 \$30,065,045 \$40,531,35 \$45,081,306 \$54,097,567 \$58,605,698 \$63,113,828 \$67,621,959 \$50,065,045 \$40,5041 \$30,05,045 \$40,5041,361 \$54,097,567 \$58,605,608 \$63,113,828 \$63,113,828 \$63,113,828 \$63,113,828 \$63,113,828 \$63,104,736 \$63,113,828 \$63,104,736 \$63,113,828 \$63,104,836 \$63,113,828 \$63,104,836 \$63,113,828 \$63,104 \$63,104 | | Vary Initic x-axis y-axis | al Value by +/- 50% 80% | | | | | | | | | |
| \$22.540.653 \$27.048.784 \$31.556.914 \$36.065.045 \$45.081.306 \$49.589.436 \$54.097.567 \$58.605.698 \$63.113.828 \$67.621.959 \$5.07.567 5.67% Negative IRR Negative IRR </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>Variable Cos</td> <td>ts</td> <td></td> <td></td> <td></td> <td></td> | | | | | | | Variable Cos | ts | | | | |
| Description State NoLowegative Irck Negative Irc | | \$22,540,653 | \$ \$27,048,784 | \$31,556,914 | \$36,065,045 | \$40,573,175 Macatine IDD | \$45,081,306 | \$49,589,436 | \$54,097,567 | \$58,605,698 | \$63,113,828 | \$67,621,959 |
| \$0.52 31.45% 24.93% 17.86% 9.85% -0.13% Negative IRR Negative | | \$0.36 5.67% | Negative IRK Negative IRR | Negative IKK Negative IRR | Negative IKK Negative IRR | Negative IRR Negative IRR | Negative IRR Negative IRR | Negative IRR Negative IRR | Negative IRR Negative IRR | Negative IRK Negative IRR | Negative IKK Negative IRR | Negative IKK Negative IRR |
| E 5.04d 61.63% 62.52% 57.33% 52.04% 46.65% 41.12% 35.42% 29.48% 16.43% 16.43% 16.43% 16.43% 16.43% 16.75% 16.43% 16.43% 16.43% 187% 19.42% 13.12% 16.63% 11.12% 106.67% 11.12% 106.67% 111.12% 106.67% 111.12% 106.67% 111.12% 106.67% 111.12% 106.67% 111.12% 110.67% 111.01% 106.65% 110.22% 88.64% 88.64% 88.06% 74.12% 64.44% 59.51% 70.09% 65.33% T 112.43 106.67% 110.65% 110.22% 111.01% 110.65% 102.24% 98.99% 84.50% 79.98% S1.4d 136.21% 112.33% 127.83% 110.40% 115.16% 110.09% 102.34% 98.02% 93.68% | e | \$0.52 31.45% \$0.68 50.71% | 24.93% 45.15% | 17.86% 30.43% | 9.85% 33.51% | -0.13% 27 30% | Negative IRR | Negative IRR 13 36% | Negative IRR | Negative IRR | Negative IRR | Negative IRR |
| 31 31.12% 78.29% 73.42% 68.49% 63.50% 58.45% 53.31% 48.07% 42.72% 37.22% 31.51% 31 31.16 97.54% 92.93% 88.28% 88.46% 54.51% 69.51% 54.51% 49.272% 31.22% 31.51% 5 51.16 97.54% 92.93% 88.360% 78.88% 74.12% 69.31% 64.44% 59.51% 54.51% 49.42% 5 51.12 110.6.67% 102.20% 97.71% 93.19% 88.64% 84.06% 79.44% 74.79% 70.09% 65.33% 51.48 123.397% 119.67% 111.01% 106.65% 102.27% 97.87% 93.44% 88.99% 84.50% 79.98% 51.64 136.21% 127.83% 123.62% 119.40% 115.16% 110.90% 106.63% 93.68% | Price | \$0.84 67.63% | 62.52% | 57.33% | 52.04% | 46.65% | 41.12% | 35.42% | 29.48% | 23.21% | 16.43% | 8.78% |
| 2 \$1.16 97.54% 92.93% 88.28% 83.60% 78.88% 74.12% 69.31% 64.44% 59.51% 54.51% 49.42% 2 \$1.13 111.12% 106.67% 102.20% 97.71% 93.19% 88.64% 84.06% 79.44% 74.79% 70.09% 65.33% \$1.14 123.97% 119.67% 111.01% 106.65% 102.27% 97.87% 93.44% 88.99% 84.50% 79.98% \$1.64 136.21% 127.83% 119.40% 115.16% 110.90% 106.63% 93.68% | l tou | \$1.00 83.12% | 78.29% | 73.42% | 68.49% | 63.50% | 58.45% | 53.31% | 48.07% | 42.72% | 37.22% | 31.51% |
| C \$1.32 111.12% 106.67% 102.20% 97.71% 93.19% 88.64% 84.06% 79.44% 74.79% 70.09% 65.33% \$1.48 123.97% 119.67% 111.01% 106.65% 102.27% 97.87% 93.44% 74.79% 70.09% 65.33% \$1.48 123.97% 119.67% 111.01% 106.65% 102.27% 97.87% 93.44% 88.99% 84.50% 79.98% \$1.64 136.21% 132.03% 127.83% 119.40% 115.16% 110.663% 102.34% 98.02% 93.68% | 1poJ | \$1.16 97.54% | 92.93% | 88.28% | 83.60% | 78.88% | 74.12% | 69.31% | 64.44% | 59.51% | 54.51% | 49.42% |
| \$1.48 123.97% 119.67% 115.35% 111.01% 106.65% 102.27% 97.87% 93.44% 88.99% 84.50% 79.98% \$1.64 136.21% 132.03% 127.83% 123.62% 119.40% 115.16% 110.90% 106.63% 102.34% 98.02% 93.68% | Ы | \$1.32 111.12% | 106.67% | 102.20% | 97.71% | 93.19% | 88.64% | 84.06% | 79.44% | 74.79% | 70.09% | 65.33% |
| \$1.64 136.21% 132.03% 127.83% 123.62% 119.40% 115.16% 110.90% 106.63% 102.34% 98.02% 93.68% | | \$1.48 123.97% | 119.67% | 115.35% | 111.01% | 106.65% | 102.27% | 97.87% | 93.44% | 88.99% | 84.50% | 79.98% |
| | | \$1.64 136.21% | 132.03% | 127.83% | 123.62% | 119.40% | 115.16% | 110.90% | 106.63% | 102.34% | 98.02% | 93.68% |

| | | \$63,113,828 | Negative IRR | Negative IRR | Negative IRR | Negative IRR | 16.43% | 37.22% | 54.51% | 70.09% | 84.50% | 98.02% | 110.81% |
|-------|---------------|--------------|---------------------|--------------|--------------|--------------|----------|----------|--------|-----------|-----------|-----------|-----------|
| | | \$58,605,698 | Negative IRR | Negative IRR | Negative IRR | Negative IRR | 23.21% | 42.72% | 59.51% | 74.79% | 88.99% | 102.34% | 114.98% |
| | | \$54,097,567 | Negative IRR | Negative IRR | Negative IRR | 4.80% | 29.48% | 48.07% | 64.44% | 79.44% | 93.44% | 106.63% | 119.14% |
| | S | \$49,589,436 | Negative IRR | Negative IRR | Negative IRR | 13.36% | 35.42% | 53.31% | 69.31% | 84.06% | 97.87% | 110.90% | 123.29% |
| | Variable Cost | \$45,081,306 | Negative IRR | Negative IRR | Negative IRR | 20.67% | 41.12% | 58.45% | 74.12% | 88.64% | 102.27% | 115.16% | 127.42% |
| | | \$40,573,175 | Negative IRR | Negative IRR | -0.13% | 27.30% | 46.65% | 63.50% | 78.88% | 93.19% | 106.65% | 119.40% | 131.53% |
| | | \$36,065,045 | Negative IRR | Negative IRR | 9.85% | 33.51% | 52.04% | 68.49% | 83.60% | 97.71% | 111.01% | 123.62% | 135.64% |
| | | \$31,556,914 | Negative IRR | Negative IRR | 17.86% | 39.43% | 57.33% | 73.42% | 88.28% | 102.20% | 115.35% | 127.83% | 139.73% |
| 80% | | \$27,048,784 | Negative IRR | Negative IRR | 24.93% | 45.15% | 62.52% | 78.29% | 92.93% | 106.67% | 119.67% | 132.03% | 143.81% |
| S | | \$22,540,653 | Negative IRR | 5.67% | 2 31.45% | 3 50.71% | 4 67.63% | 9 83.12% | 97.54% | 2 111.12% | 3 123.97% | 4 136.21% | 0 147.89% |
| y-axi | | | \$0.2(| \$0.36 | \$0.52 | \$0.68 | \$0.8 | \$1.00 | \$1.16 | \$1.32 | \$1.48 | \$1.6⁄ | \$1.8(|

Section 21

CONCLUSION AND RECOMMENDATIONS

Conclusion and Recommendations

The extremely high profitability metrics calculated and presented in this report show that this process is very profitable and should be implemented immediately. However, this analysis is likely optimistic. The equipment costing estimation methods used produced cost values for some of the process units that appear to be lower than the typical cost for such equipment. Discussions with industrial design consultants revealed that the distillation towers particularly should cost approximately 2-3 times more than the calculated prices. Additionally, the annual cost of the electrodeionization equipment leasing was rougly estimated. Due to the proprietary nature of the process, detailed information is not currently available to the public regarding its price or operation; however, conservative estimates were made to account for this uncertainty.

The reaction mechanism remains yet unverified on an industrial scale of our magnitude. The construction of a pilot plant would be an excellent indication of whether such large scale up is feasible. It is expected that the reaction will not reach 100% conversion, but as similar reactions are used in methanol production and hydrodesulfurization, the scale up is expected to be successful. If the large scale reaction produces measurable quantities of ethylene glycol, although this was unobserved in the work of Zhou et.al. (2010), another distillation tower would be required in order to remove this heavier impurity.

Despite these challenges, the difference in purchase price between propylene glycol and crude glycerol is very high at approximately \$0.78/lb, and so there inheriently is potential for profitability. The price of glycerol is inversely correlated with the production of biodiesel, as glycerol is a byproduct of this process. As biodiesel production is predicted to increase over the allotted time period for the project, the price of crude glycerol should decrease. The invention and implementation of new methods such as electrodeionization for refining crude glycerol that

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are currently being developed will make glycerol refinement increasingly profitable, which may cause an increase in the price of the raw material. While the price of propylene glycol varies seasonally with highest demand in the winter, its overall usefulness and superiority over ethylene glycol in almost every specification will ensure that the price of propylene glycol remains high. Additionally, propylene glycol is traditionally manufactured from petroleum products, so there will continue to be a large opportunity cost involved in using propylene oxide, since the propylene oxide could be transformed instead into other useful products if it was not being used for propylene glycol production. EDI is an increasingly more attractive method of purifying glycerol, since the operating costs are extremely low. In the future, it will be increasingly hard for companies like Dow to maintain as much proprietary control over this process.

Overall, the method presented in this report is expected to be profitable and successful. It is recommended that this project be further investigated with the construction of a pilot plant, and then cautiously implemented industrially.

Section 22

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Section 23

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Section 25

APPENDIX

Detailed Equipment Pricing Sample Calculations

The following examples are rigorous calculations examples for every type of unit used during this process. Correlations, calculations and heuristics were taken from the text <u>Product & Process Design Principles</u> by Seader, Seider and Lewin (2009). The accuracy of this method is plus or minus 20%.

The price index for all calculations were based on the CE Index = 560.4 for December 2010, which is the most recent CE Index value posted on www.CHE.com (2011).

<u>Mixer M-102</u>

The mixer uses a turbine, which is the preferred type of agitator for mixing miscible and immiscible liquids. To find the horsepower requirement for mixing miscible liquids, the following correlation was used:

$$\frac{HP}{1000 \ Gallons} = 0.5$$

ASPEN Reports: Q = 3520.33 gal/hr

At a residence time of 10 minutes,

Q = 586.7 gallons

 $HP = 586.7 gallons \times \frac{0.5 HP}{1000 \ gallons} = 0.29 HP$

For an efficiency of $\eta = 0.3$, HP = 1

The cost of a closed vessel mixer is given by:

 $C_p = 3300S^{0.17}$ $C_p = 3300(1)^{0.17} = $3,000$ $C_{BM-2006} = (F_{BM})(C_p)$ $C_{BM-2006} = (1)(3,300)$ $C_{BM-2006} = $3,300$ 2006 CE Index = 500 2011 CE Index = 560.4 $C_{BM-2011} = $3,300 \times 560.4/500$ $C_{BM-2011} = $3,699$

Distillation Column D-101

The total free on board (f.o.b) purchase cost for the unit is calculated using the following relationship:

 $C_P = F_M C_V + C_{PL} + C_T$

Where F_M = the material cost factor C_V = the cost of an empty vessel C_{PL} = the cost for the platforms and ladders C_T = the cost of the trays

Length Calculation

ASPEN Report: After accounting for the condenser and reboiler, Theoretical trays = 4

O'Connell Efficiency = 0.093

 $N(actual) = \frac{4}{0.093} = 43 \ trays$

L = (43)(2) + (14) = 100 feet

Thickness Calculation

The thickness for the column was estimated using the following equation:

 $t_{s} = \frac{P_{d}D_{i}}{2SE - 1.2P_{d}}$ Where P_{d} = Design pressure S = Maximum allowable stress E = Weld efficiency

ASPEN Report: $D_i = 3.9$ ft

 $P_d = 10 \text{ psig.}$ This assumption is made because the column operates at 20 psia (5 psig) S = 13,750 psiCommonly used for carbon steel over a design temperature range of -20F to 650F E=1.0

$$t_s = \frac{(10)(3.9)}{2(13750)(1) - 1.2(10)} = 0.001415 \text{ inches}$$

The theoretically determined thickness is too small for the required rigidity; therefore, a minimum wall thickness of 0.25 inches was used with a corrosion allowance of 0.125 inches.

 $t_s = 0.375 inches = 0.0313 feet$

Weight Calculation

The weight of the column was calculated using the following equation:

$$W = \pi (D_i + t_s)(L + 0.8D_i)t_s\rho$$

Where D_i = diameter of the column (ft) L = the length (ft) t_s = the wall thickness (ft) ρ = the density of carbon steel (490 lb/ft³)

 $W = \pi(3.9 + 0.0313)(100 + 0.8 * 3.9) * 0.0313 * 490$

 $W = 25,315 \ lbs$

Cost of Empty Vessel

The f.o.b cost of the empty vessel was calculated using the following equation:

 $C_{W} = e^{[7.037 + 0.18255(\ln(W) + 0.02297(\ln(W)^{2})]]}$

$C_V = 165794.26$

Cost of Platforms and Ladders

The f.o.b cost of the platforms and ladders was calculated using the following equation:

$$C_{PL} = 361.8(D_i)^{0.7960}(L)^{0.70684}$$

$C_{PL} = 28517.01$

Cost of Trays

The f.o.b cost of the empty vessel was calculated using the following equation: $C_T = N_t F_{nt} F_{tt} F_{tm} C_{bt}$

$$\begin{split} N_t &= number \ of \ trays = 43 \\ F_{nt} &> 1 \ therefore = 1 \\ F_{tt} &= 1.0 \ since \ sieve \ trays \ are \ used \\ F_{tm} &= 1.413 \ since \ stainless \ steel \ is \ used \end{split}$$

 $C_{bt} = 468e^{(0.1739*D_i)}$

 $C_{bt} = base cost = 922.1$

 $C_T = (43)(1)(1)(1)(1)(922.1)$

$C_T = 55935.83$

Cost of Vertical Tower

 $C_P = F_M C_V + C_{PL} + C_T$

 $C_P = (1)(165794.26) + (28517.01) + (39650.3)$

 $C_P = $250,244$

$$\begin{split} & C_{BM\text{-}2006} = (F_{BM})(C_p) \\ & C_{BM\text{-}2006} = (4.16)(250244) \\ & C_{BM\text{-}2006} = \$1,041,027 \end{split}$$

2006 CE Index = 500 2011 CE Index = 560.4

 $C_{BM-2011} =$ \$1,041,027 x 560.4/500 $C_{BM-2011} =$ \$1,166,781

Cost of Reflux Accumulator D-101b

The total free on board (f.o.b) purchase cost for the unit is calculated using the following relationship:

 $C_P = F_M C_V + C_{PL}$

Where F_M = the material cost factor C_V = the cost of an empty horizontal vessel C_{PL} = the cost for the platforms and ladders

Cost of Horizontal Vessel

ASPEN Report: Distillate Rate = 248.65 cuft/hr Reflux Ration = 0.2

Volumetric Flow Rate = $248.65 + 0.2 * 248.65 = 297.6 \frac{cuft}{hr}$

For a residence time of $\tau = 0.08 hrs$, $Volume = \tau * 297.6 = 47.6 cuft$

$$D = \frac{4 * V * \tau^{0.33}}{\pi}$$
$$D = \frac{4 * 47.6 * (.08)^{0.33}}{\pi} = 1.61 \, feet$$

L = 2(D) = 3.21 feet

Using the equations for vessel thickness presented before, the thickness is found to be 0.375 inches. The weight is then found to equal 355 lbs.

The f.o.b cost of the empty vessel was calculated using the following equation:

 $C_V = e^{(8.717 - 0.2330 \cdot \ln(W) + 0.04333 \cdot \ln(W)^2)}$

 $C_V = 8,788.46$

Cost of Platforms and Ladders

The f.o.b cost of the platforms and ladders was calculated using the following equation:

 $C_{PL} = 2005 (D_i)^{0.79600.20294}$

 $C_{PL} = 2208.45$

Cost of Reflux Accumulator

 $C_P = F_M C_V + C_{PL}$

 $C_P = (1)(8788.46) + 2208.45$

$C_p = $10,996.91$

$$\begin{split} &C_{BM\text{-}2006} = (F_{BM})(C_p) \\ &C_{BM\text{-}2006} = (3.05)(10996.91) \\ &C_{BM\text{-}2006} = \$33,540 \end{split}$$

2006 CE Index = 500 2011 CE Index = 560.4

 $C_{BM-2011} = $33,540 \text{ x } 560.4/500$ $C_{BM-2011} = $37,593$

Cost of Centrifugal Pump D-101c

The cost of a centrifugal pump is given by the equation:

$$C_P = (F_t)(F_m)(C_b)$$

Where F_t = pump-type factor F_m = material factor C_b = base cost

The size factor for the centrifugal pump is calculated using the following equation:

 $S = QH^{0.5}$ Q = 31 * 0.2 + 31 = 37.2 gpm $H = \frac{p}{\rho} = \frac{\rho gh}{\rho} = 2085.5 \text{ ft}$ $\rho = 56.25 \text{ lb/cuft}$ $g = 32.0 \text{ ft/s}^2$ $g_c = 3.28 \text{ ft/s}^2$ h = 100 ft $S = (37.2)(2085.2)^{0.5} = 1698.7$ $C_B = \exp[9.7171 - 0.6019[\ln(S)] + 0.519[\ln(S)]^2]$ $C_B = 3332.25$ $C_P = (F_t)(F_m)(C_b)$ $C_P = (1.35)(1)(3332.25) = 4498.54 $C_{BM-2006} = (F_{BM})(C_p)$ $C_{BM-2006} = (3.3)(4498.54)$ $C_{BM-2006} = $12,162.7$

Electric Motor:

The size parameter of the motor, P_c, can be found using the following equation:

$$P_c = \frac{QH\rho}{(33000\eta_p\eta_m)}$$

Where:

$$\begin{split} Q &= \text{flow rate through the pump (gpm)} \\ H &= \text{Pump Head} \\ \rho &= \text{density (lb/gal)} \\ \eta_p &= \text{fractional efficiency} \\ \eta_m &= \text{fractional efficiency of the electric motor} \end{split}$$

ASPEN Report:

Q = 37.2 gpmH = 2085.5 ft $\rho = 7.52 \text{ lb/gal}$ $\eta_P = -0.316 + 0.24015 \ln(Q) - 0.01199 \ln(Q)^2$ $\eta_P = 0.396$ $\eta_m = 0.8 + 0.0319 \ln(P_b) - 0.00182 \ln(P_b)^2$ $\eta_m = 0.895$

Where:

$$P_b = \frac{QH\rho}{(33000\eta_p)} = 44.64$$

Plugging into the equation for P_c above, it was found to equal 373.2 HP.

The cost of the motor was then calculated using the following equation:

 $C_b = \exp[5.8259 + 0.1314 \ln(P_c) + 0.053255 \ln \ln(P_c)^2 + 0.02865 \ln(P_c)^3 C_b = \$22,852$

 $C_P = (F_t)(C_b)$ $C_P = (1)(22,852) = $22,852$

 $C_{BM-2006} = (F_{BM})(C_p)$ $C_{BM-2006} = (1.0)(22852)$ $C_{BM-2006} = $22,852$

Adding the cost of the electric motor,

 $C_{BM-2006} = 12162.7 + 22852.3$ $C_{BM-2006} = $35,015$

2006 CE Index = 500 2011 CE Index = 560.4

 $C_{BM-2011} = $35,015 \text{ x } 560.4/500$ $C_{BM-2011} = $37,003$

Cost of the Fired Heater H-101

The cost of a fired heater is given by the equation:

 $C_P = (F_t)(F_m)(C_b)$

Where

 $C_b = \exp[0.32325 + 0.766 * \ln(Q)]$

ASPEN Report: Q = 106,893 Btu/hr

 $C_p = (1.7)(0.986)(106893) = 179,162$

$$\begin{split} C_{BM\text{-}2006} &= (F_{BM})(C_p) \\ C_{BM\text{-}2006} &= (1.86)(117162) \\ \textbf{C}_{BM\text{-}2006} &= \$333,\!241 \end{split}$$

Cost of Turbine TR-100

The cost of a turbine is given by the equation:

 $C_p = 3620S^{0.57}$ ASPEN Report: S = 27 HP

 $C_p = 3620(27)^{0.57} = \$23,391$

$$\begin{split} C_{BM\text{-}2006} &= (F_{BM})(C_p) \\ C_{BM\text{-}2006} &= (2.15)(23391) \\ \textbf{C_{BM\text{-}2006}} &= \textbf{\$50, \textbf{936}} \end{split}$$
Cost of Storage Tank T-101

The cost of the storage tank T-101 is given by the equation:

 $C_P = 475V^{0.51}$

ASPEN Report Q = 1181.02 gal/hr

For two weeks of storage, the required volume is:

 $1811.02 \frac{gal}{hr} \times \frac{24hr}{day} \times 14days = 608503 \ gal$ $C_P = 475(608503)^{0.51}$ $C_P = \$423,319$ $C_{BM-2006} = (F_{BM})(C_p)$ $C_{BM-2006} = (1.0)(423319)$ $C_{BM-2006} = \$423,319$

Electricity Requirement of EDI Machine

$$\xi = \frac{zFQ_f(C_{inlet}^d - C_{outlet}^d)}{NI}$$

Efficiency is about 0.8 for USP applications. (EETCorp, 2011)

Charge magnitude (z) = 1 for NaCl

Faraday's Constant = 96485 Amp-s/mol

Diluent volumetric flow rate (Q_f) = 1.63 L/s

Salt inlet concentration = 0.82 mol/L for each sodium and chloride ions. (assuming sulfate and nitrates are negligible).

Salt outlet concentration is $10ppm = 2.05 \times 10^{-5}$ for both ions\

N = 800 (Datta, 2002)

So I = 400 Amps

Assume 0.5 V/cell (Ibid.)

With 400 V (.5 V/cell * 800 cells), power requirements are 160 kW. Actual power draw requirements should be 200 kW to account for scale buildup on the membranes. (Ibid.)

Calculation for Mass of Catalyst in Reactor

| | lb/hr | MW | lbmol/hr | gram-mol/hr | gram-mol/s | | | |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--|--|--|
| Glycerol In | 20075.16 | 92.09 | 217.995 | 98902.15155 | 27.47281988 | | | |
| H2 In (7:1 Mole ratio w/ Glycerol) | 3051.93 | 2 | 1525.965 | 692315.0609 | 192.3097391 | | | |
| H2O In | 3764.092 | 18 | 209.1162 | 94873.94934 | 26.35387482 | | | |
| Glycerol Out | 3713.905 | 92.09 | 40.32908 | 18296.89804 | 5.082471677 | | | |
| PG Out | 12626.26 | 76.09 | 165.9385 | 75284.65095 | 20.91240304 | | | |
| H2 Consumed by Rxn | 355.3319 | 2 | 177.6659 | 80605.25351 | 22.3903482 | | | |
| H2O Produced by Rxn | 24480.48 | 18 | 1360.026 | 617030.4099 | 171.3973361 | | | |
| H2 Out | 2696.598 | 2 | 1348.299 | 611709.8073 | 169.9193909 | | | |
| H2O Out | 28244.57 | 18 | 1569.143 | 711904.3592 | 197.7512109 | | | |
| Acetol Out | 157.9415 | 74.08 | 2.13204 | 967.2852276 | 0.268690341 | | | |
| Ethylene Glycol Out | 297.7604 | 62.07 | 4.797171 | 2176.428511 | 0.604563475 | | | |
| Other Out | | | 4.797171 | 2176.428511 | 0.604563475 | | | |
| After scaling up to find 100% Conversion | | | | | | | | |
| | | | | | | | | |
| | lb/hr | MW | lbmol/hr | gram-mol/hr | gram-mol/s | | | |
| Glycerol In | lb/hr 15281.28 | MW 92.09 | lbmol/hr 165.9385 | gram-mol/hr 75284.65095 | gram-mol/s 20.91240304 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) | lb/hr 15281.28 2323.139 | MW 92.09 2 | lbmol/hr 165.9385 1161.57 | gram-mol/hr 75284.65095 526992.5567 | gram-mol/s 20.91240304 146.3868213 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In | lb/hr 15281.28 2323.139 2865.24 | MW 92.09 2 18 | lbmol/hr 165.9385 1161.57 159.18 | gram-mol/hr 75284.65095 526992.5567 72218.36986 | gram-mol/s 20.91240304 146.3868213 20.06065829 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out | lb/hr 15281.28 2323.139 2865.24 2827.037 | MW 92.09 2 18 92.09 | lbmol/hr 165.9385 1161.57 159.18 30.69863 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 | MW 92.09 2 18 92.09 76.09 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 | MW 92.09 2 18 92.09 76.09 2 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn H2O Produced by Rxn | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 17921.36 | MW 92.09 2 18 92.09 76.09 2 18 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 995.6312 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 451707.9057 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 125.4744183 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn H2O Produced by Rxn H2O Ut | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 17921.36 2052.66 | MW 92.09 2 18 92.09 76.09 2 18 2 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 995.6312 1026.33 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 451707.9057 465635.5661 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 125.4744183 129.3432128 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn H2O Produced by Rxn H2 Out H2 Out | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 17921.36 2052.66 20786.6 | MW 92.09 2 18 92.09 76.09 2 18 2 18 2 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 995.6312 1026.33 1154.811 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 451707.9057 465635.5661 523926.2756 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 125.4744183 129.3432128 145.5350765 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn H2O Produced by Rxn H2 Out H2O Out Acetol Out | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 17921.36 2052.66 20786.6 157.9415 | MW 92.09 2 18 92.09 76.09 2 18 2 18 2 18 74.08 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 995.6312 1026.33 1154.811 2.13204 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 451707.9057 465635.5661 523926.2756 967.2852276 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 125.4744183 129.3432128 145.5350765 0.268690341 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn H2O Produced by Rxn H2O Out H2O Out Acetol Out Ethylene Glycol Out | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 17921.36 2052.66 20786.6 157.9415 297.7604 | MW 92.09 2 18 92.09 76.09 2 18 2 18 74.08 62.07 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 995.6312 1026.33 1154.811 2.13204 4.797171 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 451707.9057 465635.5661 523926.2756 967.2852276 2176.428511 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 125.4744183 129.3432128 145.5350765 0.268690341 0.604563475 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn H2O Produced by Rxn H2O Ut H2O Out Acetol Out Ethylene Glycol Out Other Out | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 17921.36 2052.66 20786.6 157.9415 297.7604 | MW 92.09 2 18 92.09 76.09 2 18 2 18 74.08 62.07 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 995.6312 1026.33 1154.811 2.13204 4.797171 4.797171 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 451707.9057 465635.5661 523926.2756 967.2852276 2176.428511 2176.428511 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 125.4744183 129.3432128 145.5350765 0.268690341 0.604563475 0.604563475 | | | |
| Glycerol In H2 In (7:1 Mole ratio w/ Glycerol) H2O In Glycerol Out PG Out H2 Consumed by Rxn H2O Produced by Rxn H2 Out H2O Out Acetol Out Ethylene Glycol Out Other Out | lb/hr 15281.28 2323.139 2865.24 2827.037 12626.26 270.4798 17921.36 2052.66 20786.6 157.9415 297.7604 | MW 92.09 2 18 92.09 76.09 2 18 2 18 74.08 62.07 | lbmol/hr 165.9385 1161.57 159.18 30.69863 165.9385 135.2399 995.6312 1026.33 1154.811 2.13204 4.797171 | gram-mol/hr 75284.65095 526992.5567 72218.36986 13927.66043 75284.65095 61356.99053 451707.9057 465635.5661 523926.2756 967.2852276 2176.428511 2176.428511 | gram-mol/s 20.91240304 146.3868213 20.06065829 3.868794563 20.91240304 17.04360848 125.4744183 129.3432128 145.5350765 0.268690341 0.604563475 0.604563475 | | | |

| 0 | , | | | | | | | |
|-----------------------------|---------------|------|--------|------|-----|--|--|--|
| Selectivity % | | | | | | | | |
| Catalyst (3.0g Loading) | Glycerol C(PG | Ace | tol EG | Othe | r* | | | |
| 40% CuO; 40% ZnO; 20% Al2O3 | 81.5 | 93.4 | 1.2 | 2.7 | 2.7 | | | |

LHSV = 4.6 h-1

| Catalyst Mass | Flow of G | Flow of A | Flow of PG | Flow of W | Conversion of G |
|---------------|-------------|-------------|-------------|-------------|-----------------|
| (kg) | (mol/s) | (mol/s) | (mol/s) | (mal/s) | |
| 0 | 20.91240304 | 0 | 0 | 26.35443056 | |
| 10 | 20.80856741 | 0.103735628 | 0 | 26.45816618 | 0.004950483 |
| 20 | 20.70508926 | 0.193070255 | 0.014243529 | 26.56174434 | 0.009913437 |
| 30 | 20.60166158 | 0.25989431 | 0.040847154 | 26.66517202 | 0.014859194 |
| 40 | 20.49837839 | 0.335853607 | 0.078171041 | 25.7684552 | 0.019798043 |
| 50 | 20.3952346 | 0.392380933 | 0.124787511 | 26.871599 | 0.024730226 |
| 60 | 20.29222585 | 0.44072345 | 0.179453735 | 26.97460775 | 0.029655951 |
| 70 | 20.18934847 | 0.481966585 | 0.241087987 | 27.07748513 | 0.034575394 |
| 80 | 20.08659937 | 0.517054683 | 0.308748985 | 27.18023422 | 0.039488703 |
| 90 | 19.98397598 | 0.545809181 | 0.381617885 | 27.28285762 | 0.044396001 |
| 100 | 19.88147614 | 0.57194432 | 0.458982586 | 27.38535746 | 0.049297391 |
| 110 | 19.7790981 | 0.593080902 | 0.540224039 | 27.4877355 | 0.054192956 |
| 120 | 19.67584045 | 0.610758289 | 0.624804296 | 27.58999314 | 0.059082765 |
| 130 | 19.57470209 | 0.625444871 | 0.712256084 | 27.69213151 | 0.063966869 |
| 140 | 19.47268213 | 0.637547214 | 0.8021737 | 27.79415147 | 0.068845312 |
| 150 | 19.37077995 | 0.647418038 | 0.894205057 | 27.89605365 | 0.073718123 |
| 160 | 19.26899511 | 0.655363191 | 0.988044745 | 27.99783849 | 0.078585322 |
| 170 | 19.16732734 | 0.661647727 | 1.083427974 | 28.09950526 | 0.083446924 |
| 180 | 19.06577654 | 0.66650122 | 1.180125284 | 28.20105706 | 0.088302932 |
| 190 | 18.96434272 | 0.670122397 | 1.277937929 | 28.30249088 | 0.093153346 |
| 200 | 18.86302601 | 0.672683177 | 1.376693854 | 28,40380759 | 0.097998161 |
| 210 | 18.76182665 | 0.5743322 | 1.476244183 | 28.50500594 | 0.102837363 |
| 220 | 18.66074499 | 0.675197896 | 1.57646016 | 28.60608861 | 0.107670938 |
| 230 | 18.5597814 | 0.675391162 | 1.677230481 | 28.7070522 | 0.112498867 |
| 240 | 18.45893637 | 0.675007697 | 1.778458976 | 28.80789723 | 0.117321126 |
| 250 | 18.35821043 | 0.674130025 | 1.880062583 | 28.90862316 | 0.12213769 |
| 260 | 18.25760418 | 0.672829265 | 1.981969593 | 29.00922942 | 0.126948532 |
| 270 | 18.15711825 | 0.67116657 | 2.08411812 | 29.10971535 | 0.131753619 |
| 280 | 18.05675332 | 0.669194946 | 2.186454771 | 29.21008027 | 0.136552921 |
| 290 | 17.95551012 | 0.66695943 | 2.288933495 | 29.31032348 | 0.141346402 |
| 300 | 17.85538938 | 0.664499083 | 2.391514576 | 29.41044421 | 0.146134026 |
| 310 | 17.75639191 | 0.661847365 | 2.49416377 | 29.51044169 | 0.150915757 |
| 320 | 17.6555185 | 0.659032995 | 2.596851549 | 29.6103151 | 0.155691555 |
| 330 | 17.55677 | 0.655080601 | 2.699552445 | 29.7100536 | 0.16045138 |
| 340 | 17.45714726 | 0.653011291 | 2.802244491 | 29.80968634 | 0.165225191 |
| 350 | 17.35765117 | 0.649843141 | 2.90490873 | 29.90918243 | 0.169982946 |
| 360 | 17.25828263 | 0.645591623 | 3.007528789 | 30.00855097 | 0.174734602 |
| 370 | 17.15904256 | 0.643269966 | 3.110090519 | 30.10779104 | 0.179480114 |
| 380 | 17.05993189 | 0.639889478 | 3.212581675 | 30.20690171 | 0.184219439 |
| 390 | 16.96095158 | 0.635459815 | 3.314991648 | 30.30588202 | 0.18895253 |
| 400 | 16.86210259 | 0.632989219 | 3.417311228 | 30.404731 | 0.193679341 |
| 410 | 16.76338592 | 0.629484719 | 3.519532404 | 30.50344768 | 0.198399826 |
| 420 | 16.66480255 | 0.625952305 | 3.621648187 | 30.60203105 | 0.203113936 |

| 3130 | 0.409093483 | 0.015388397 | 20.48792116 | 46.85774011 | 0.980437758 |
|------|-------------|-------------|-------------|-------------|-------------|
| 3140 | 0.399440404 | 0.015025289 | 20.49793735 | 46.86739319 | 0.980899354 |
| 3150 | 0.389998313 | 0.014670116 | 20.50773461 | 46.87683528 | 0.981350861 |
| 3160 | 0.380763327 | 0.014322735 | 20.51731698 | 46.88607027 | 0.981792464 |
| 3170 | 0.371731603 | 0.013982999 | 20.52668844 | 46.89510199 | 0.982224348 |
| 3180 | 0.362899342 | 0.013650766 | 20.53585293 | 46.90393426 | 0.982646693 |
| 3190 | 0.354262785 | 0.013325894 | 20.54481436 | 46.91257081 | 0.983059681 |
| 3200 | 0.345818219 | 0.013008245 | 20.55357658 | 46.92101538 | 0.983453487 |
| 3210 | 0.337561973 | 0.012697679 | 20.56214339 | 46.92927162 | 0.983858289 |
| 3220 | 0.329490421 | 0.012394061 | 20.57051856 | 46.93734318 | 0.984244258 |
| 3230 | 0.321599982 | 0.012097255 | 20.5787058 | 46.94523362 | 0.984621567 |
| 3240 | 0.313887123 | 0.01180713 | 20.58670879 | 46.95294647 | 0.984990385 |
| 3250 | 0.306348354 | 0.011523552 | 20.59453114 | 46.96048524 | 0.985350878 |
| 3260 | 0.298980232 | 0.011246394 | 20.60217642 | 46.96785337 | 0.98570321 |
| 3270 | 0.291779363 | 0.010975528 | 20.60964815 | 46.97505423 | 0.986047545 |
| 3280 | 0.284742399 | 0.010710826 | 20.61694982 | 46.9820912 | 0.986384042 |
| 3290 | 0.277856038 | 0.010452166 | 20.62408484 | 46.98895756 | 0.986712859 |
| 3300 | 0.271147027 | 0.010199425 | 20.63105659 | 46.99568657 | 0.987034153 |
| 3310 | 0.264582161 | 0.009952482 | 20.6378684 | 47.00225144 | 0.987348075 |
| 3320 | 0.258168283 | 0.009711218 | 20.64452354 | 47.00865532 | 0.987654777 |
| 3330 | 0.251902281 | 0.009475517 | 20.65102524 | 47.01493132 | 0.987954408 |
| 3340 | 0.245781094 | 0.009245264 | 20.65737668 | 47.0210525 | 0.988247114 |
| 3350 | 0.239801708 | 0.009020344 | 20.66358099 | 47.02703189 | 0.988533039 |
| 3360 | 0.233961158 | 0.008800647 | 20.66964124 | 47.03287244 | 0.988812326 |
| 3370 | 0.228256523 | 0.008586062 | 20.67556046 | 47.03857707 | 0.989085113 |
| 3380 | 0.222684935 | 0.008376482 | 20.68134163 | 47.04414866 | 0.989351538 |
| 3390 | 0.217243568 | 0.008171801 | 20.68698767 | 47.04959003 | 0.989611736 |
| 3400 | 0.211929647 | 0.007971913 | 20.69250148 | 47.05490395 | 0.98986584 |
| 3410 | 0.206740443 | 0.007776717 | 20.69788588 | 47.06009315 | 0.99011398 |
| 3420 | 0.201673273 | 0.007586111 | 20.70314366 | 47.06516032 | 0.990356284 |
| 3430 | 0.196725502 | 0.007399995 | 20.70827754 | 47.0701081 | 0.990592879 |
| 3440 | 0.191894539 | 0.007218275 | 20.71329023 | 47.07493906 | 0.990823889 |
| 3450 | 0.187177843 | 0.007040853 | 20.71818435 | 47.07965576 | 0.991049434 |
| 3460 | 0.182572913 | 0.005867635 | 20.72295249 | 47.08425068 | 0.991269635 |
| 3470 | 0.178077297 | 0.005598528 | 20.72762722 | 47.0887563 | 0.991484609 |
| 3480 | 0.173688588 | 0.006533443 | 20.73218101 | 47.09314501 | 0.99169447 |
| 3490 | 0.169404421 | 0.005372291 | 20.73662633 | 47.09742918 | 0.991899333 |
| 3500 | 0.165222477 | 0.005214983 | 20.74095558 | 47.10161112 | 0.992099307 |
| 3510 | 0.161140479 | 0.006061435 | 20.74520113 | 47.10569312 | 0.992294502 |
| 3520 | 0.157156195 | 0.005911563 | 20.74933528 | 47.1096774 | 0.992485025 |
| 3530 | 0.153267434 | 0.005765284 | 20.75337032 | 47.11355616 | 0.992670979 |
| 3540 | 0.149472047 | 0.005622517 | 20.75730848 | 47.11736155 | 0.992852469 |
| 3550 | 0.145767929 | 0.005483184 | 20.76115193 | 47.12105567 | 0.993029595 |
| 3560 | 0.142153014 | 0.005347206 | 20.76490282 | 47.12468058 | 0.993202454 |
| 3570 | 0.138625276 | 0.005214507 | 20.76856326 | 47.12820832 | 0.993371146 |
| | | | | | |

Appendix

| 3580 | 0.135182732 | 0.005085013 | 20 7721353 | 47.13165087 | 0.993535763 |
|------|-------------|-------------|-------------|-------------|-------------|
| 3590 | 0.131823437 | 0.00495865 | 20,77562096 | 47.13501016 | 0.993696399 |
| 3600 | 0.128545483 | 0.004835347 | 20,77902221 | 47,13828811 | 0.993853146 |
| 3610 | 0.125342006 | 0.004715034 | 20.782341 | 47.14148659 | 0.994006093 |
| 3620 | 0.122226174 | 0.004597641 | 20,78557923 | 47.14460742 | 0.994155326 |
| 3630 | 0 119181195 | 0.004483101 | 20,78873874 | 47.1476524 | 0.994300933 |
| 3640 | 0.116210317 | 0.004371349 | 20.79182138 | 47 15062328 | 0.994442996 |
| 3650 | 0.113311819 | 0.00426232 | 20.7948289 | 47.15352178 | 0.994581597 |
| 3560 | 0.110484019 | 0.00415595 | 20 79775307 | 47 15634958 | 0.994716819 |
| 3670 | 0.10772527 | 0.004052177 | 20.80062559 | 47.15910833 | 0.994848738 |
| 3680 | 0.105033959 | 0.003950941 | 20.80341814 | 47.16179964 | 0.994977432 |
| 3690 | 0.102408505 | 0.003852183 | 20.80614235 | 47,16442509 | 0.995102978 |
| 3700 | 0.099847369 | 0.003755843 | 20.80879983 | 47.16698623 | 0.995225447 |
| 3710 | 0.097349033 | 0.003661866 | 20.81139214 | 47.16948456 | 0.995344914 |
| 3720 | 0.094912021 | 0.003570196 | 20.81392083 | 47,17192158 | 0.995461448 |
| 3730 | 0.092534884 | 0.003480778 | 20.81638738 | 47.17429871 | 0.99557512 |
| 3740 | 0.090216206 | 0.003393559 | 20.81879328 | 47.17661739 | 0.995685995 |
| 3750 | 0.087954603 | 0.003308487 | 20.82113995 | 47.17887899 | 0.995794142 |
| 3760 | 0.085748719 | 0.003225511 | 20.82342881 | 47.18108488 | 0.995899624 |
| 3770 | 0.083597229 | 0.00314458 | 20.82565123 | 47.18323637 | 0.996002505 |
| 3780 | 0.081498837 | 0.003065648 | 20.82783856 | 47.18533476 | 0.996102847 |
| 3790 | 0.079452275 | 0.002988665 | 20.8299621 | 47.18738132 | 0.99620071 |
| 3800 | 0.077456305 | 0.002913584 | 20.83203315 | 47.18937729 | 0.996296155 |
| 3810 | 0.075509716 | 0.002840352 | 20.83405296 | 47.19132388 | 0.996389238 |
| 3820 | 0.073611322 | 0.002768952 | 20.83602277 | 47.19322228 | 0.996480016 |
| 3830 | 0.071759965 | 0.002699312 | 20.83794376 | 47.19507363 | 0.996568545 |
| 3840 | 0.069954515 | 0.002631398 | 20.83981713 | 47.19687908 | 0.996654879 |
| 3850 | 0.068193865 | 0.00256517 | 20.84164401 | 47.19863973 | 0.996739071 |
| 3860 | 0.066476934 | 0.002500585 | 20.84342552 | 47.20035666 | 0.996821172 |
| 3870 | 0.064802665 | 0.002437607 | 20.84516277 | 47.20203093 | 0.996901233 |
| 3880 | 0.063170028 | 0.002376194 | 20.84685682 | 47.20366357 | 0.996979303 |
| 3890 | 0.061578011 | 0.002316309 | 20.84850872 | 47.20525559 | 0.997055431 |
| 3900 | 0.06002563 | 0.002257915 | 20.8501195 | 47.20680797 | 0.997129664 |
| 3910 | 0.058511922 | 0.002200975 | 20.85169014 | 47.20832168 | 0.997202047 |
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| 3970 | 0.050190632 | 0.001887963 | 20.86032445 | 47.21664297 | 0.997599959 |
| 3980 | 0.048922485 | 0.001840251 | 20.8616403 | 47.21791111 | 0.9976606 |
| 3990 | 0.047686071 | 0.001793752 | 20.86292322 | 47.21914753 | 0.997719723 |
| 4000 | 0.04648061 | 0.001748407 | 20.86417402 | 47.22035299 | 0.997777366 |
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| 4020 | 0.044159526 | 0.001661098 | 20.86658242 | 47.22267407 | 0.997888357 |

Appendix

| 4030 | 0.043042435 | 0.001619078 | 20.86774153 | 47.22379116 | 0.997941775 |
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| 4040 | 0.041953364 | 0.001578111 | 20.86887157 | 47.22488023 | 0.997993853 |
| 4050 | 0.04089162 | 0.001538173 | 20.86997325 | 47.22594198 | 0.998044624 |
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| 4140 | 0.032459535 | 0.001220993 | 20.87872251 | 47.23437406 | 0.998447833 |
| 4150 | 0.031636515 | 0.001190034 | 20.87957649 | 47.23519708 | 0.998487189 |
| 4160 | 0.030834232 | 0.001159856 | 20.88040895 | 47.23599937 | 0.998525553 |
| 4170 | 0.030052169 | 0.001130438 | 20.88122043 | 47.23678143 | 0.99855295 |
| 4180 | 0.029289825 | 0.001101752 | 20.88201146 | 47.23754377 | 0.998599404 |
| 4190 | 0.028546707 | 0.001073809 | 20.88278253 | 47.23828689 | 0.998634939 |
| 4200 | 0.027822335 | 0.001046551 | 20.88353415 | 47.23901126 | 0.998659577 |
| 4210 | 0.027116245 | 0.00102 | 20.8842558 | 47.23971735 | 0.998703341 |
| 4220 | 0.026427977 | 0.000994111 | 20.88498095 | 47.24040562 | 0.998736253 |
| 4230 | 0.025757088 | 0.000968875 | 20.88567708 | 47.24107651 | 0.998758334 |
| 4240 | 0.025103142 | 0.000944276 | 20.88635562 | 47.24173046 | 0.998799605 |
| 4250 | 0.024465717 | 0.000920299 | 20.88701703 | 47.24236788 | 0.998830086 |
| 4260 | 0.023844398 | 0.000896927 | 20.88765172 | 47.2429892 | 0.998859796 |
| 4270 | 0.023238784 | 0.000874147 | 20.88829011 | 47.24359481 | 0.998888756 |
| 4280 | 0.022648481 | 0.000851942 | 20.88890262 | 47.24418512 | 0.998916983 |
| 4290 | 0.022073105 | 0.000830299 | 20.88949964 | 47.24476049 | 0.998944497 |
| 4300 | 0.021512283 | 0.000809203 | 20.89008156 | 47.24532132 | 0.998971315 |
| 4310 | 0.020965648 | 0.000788641 | 20.89064875 | 47.24586795 | 0.998997454 |
| 4320 | 0.020432845 | 0.000768599 | 20.8912016 | 47.24640075 | 0.999022932 |
| 4330 | 0.01991353 | 0.000749064 | 20.89174045 | 47.24692007 | 0.999047765 |
| 4340 | 0.01940736 | 0.000730024 | 20.89226566 | 47.24742624 | 0.999071969 |
| 4350 | 0.018914005 | 0.000711455 | 20.89277757 | 47.24791959 | 0.99909556 |
| 4360 | 0.018433147 | 0.000693378 | 20.89327652 | 47.24840045 | 0.999118554 |
| 4370 | 0.017964468 | 0.000675749 | 20.89376283 | 47.24886913 | 0.999140966 |
| 4380 | 0.017507663 | 0.000658566 | 20.89423681 | 47.24932593 | 0.99915281 |
| 4390 | 0.017062434 | 0.000641818 | 20.89469879 | 47.24977116 | 0.9991841 |
| 4400 | 0.016628489 | 0.000625495 | 20.89514906 | 47.25020511 | 0.99920485 |
| 4410 | 0.016205544 | 0.000609585 | 20.89558791 | 47.25062805 | 0.999225075 |
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| 4430 | 0.015391552 | 0.000578966 | 20.89643252 | 47.25144205 | 0.999263999 |
| 4440 | 0.014999972 | 0.000564237 | 20.89683883 | 47.25183363 | 0.999282724 |
| 4450 | 0.014618325 | 0.000549881 | 20.89723484 | 47.25221527 | 0.999300973 |
| 4460 | 0.014246359 | 0.000535889 | 20.89762079 | 47.25258724 | 0.99931876 |
| 4470 | 0.013883832 | 0.000522252 | 20.89799596 | 47.25294977 | 0.999336096 |

Section 25

PROBLEM STATEMENT

Glycerol to Renewable Propylene Glycol (recommended by Bruce M. Vrana, DuPont)

Propylene glycol (PG), is used to make unsaturated polyester resins, cosmetics, aircraft deicer, environmentally-friendly (and pet-friendly) automotive antifreezes, etc. PG is conventionally made from propylene. Propylene is itself a byproduct of ethylene manufacture. Since demand for polypropylene is growing faster than ethylene, propylene is in short supply and prices are rising. Demand and prices for PG are expected to increase. And there will likely be a price premium and plenty of demand for PG made from renewable resources.

Glycerol is a byproduct of biodiesel manufacture, with relatively few industrial uses. As the production of biodiesel increases, particularly in Europe due to government regulations but also in the U.S. due to public demand for renewable fuels, the price of glycerol is expected to continue to decrease.

BASF has patented a catalyst to convert glycerol to propylene glycol in very high yield – up to 100% glycerol conversion and 98.5% selectivity. The patent does not disclose quantitatively what the byproducts are, but you can assume the 1.5% of glycerol that does not form propylene glycol goes to equal amounts of n-propanol and isopropanol.

CH₂OH-CH₂OH-CH₂OH + H₂ \rightarrow CH₂OH-CH₂OH-CH₃ + H₂O

Crude glycerol from biodiesel manufacture contains 15% water, 4% NaCl, 1% methanol. It also contains trace amounts of organic sulfur and chlorine compounds, which must be removed, as described in the patent.

Design a process to make 100MM lb/yr of propylene glycol from crude glycerol. Your plant is on the U.S. Gulf Coast. Crude glycerol delivered to you costs \$0.22/lb (for the crude stream at 80% concentration). Hydrogen is available on your plant site for \$0.50/lb.

Renewable propylene glycol can be sold for \$1.00/lb. Byproduct renewable n-propanol and isopropanol can be sold for \$0.80/lb. All products will need to meet normal specs for that product. All prices are forecasts by your marketing organization for long-term average prices, expressed in 2011 dollars on the Gulf Coast.

The plant design should be as environmentally friendly as possible. Recover and recycle process materials to the maximum economic extent. Also, energy consumption should be minimized, to the extent economically justified. The plant design must also be controllable and safe to operate. Remember that you will be there for the plant start-up and will have to live with whatever design decisions you have made.

Reference U.S. Patent 7.790,937, September 7, 2010, assigned to BASF Section 25

PATENT AND ASPEN SIMULATION



US007790937B2

(12) United States Patent

Henkelmann et al.

(54) PROCESS FOR THE PREPARATION OF 1,2-PROPANEDIOL

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 Wahl, Heidelberg (DE); Gerhard Theis,
 Maxdorf (DE); Stephen Maurer,
 Neustadt-Gimmeldingen (DE)
- (73) Assignee: **BASF SE**, Ludwigshafen (DE)
- (*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.
- (21) Appl. No.: 12/281,406
- (22) PCT Filed: Mar. 2, 2007
- (86) PCT No.: PCT/EP2007/051983
 § 371 (c)(1),
 - (2), (4) Date: Sep. 2, 2008
- (87) PCT Pub. No.: WO2007/099161

PCT Pub. Date: Sep. 7, 2007

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(30) Foreign Application Priority Data

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- (51) Int. Cl. *C07C 29/132* (2006.01)
- (52) U.S. Cl. 568/861
 (58) Field of Classification Search 568/861 See application file for complete search history.

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(45) **Date of Patent:** Sep. 7, 2010

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Primary Examiner—Sikarl A Witherspoon

(74) Attorney, Agent, or Firm—Connolly Bove Lodge & Hutz LLP

(57) **ABSTRACT**

The present invention relates to a process for the preparation of 1,2-propanediol, in which a glycerol-containing stream, in particular a stream obtained on an industrial scale in the production of biodiesel, is subjected to a hydrogenation.

16 Claims, No Drawings

PROCESS FOR THE PREPARATION OF 1,2-PROPANEDIOL

RELATED APPLICATIONS

This application is a national stage application (under 35 U.S.C. §371) of PCT/EP2007/051983, filed Mar. 2, 2007, which claims benefit of European Application No. 06004414.6, filed Mar. 3, 2006, and U.S. Provisional Application Ser. No. 60/778,371, filed Mar. 3, 2006.

The present invention relates to a process for the preparation of 1,2-propanediol, in which a glycerol-containing stream, in particular a stream obtained on an industrial scale in the preparation of biodiesel, is subjected to a hydrogenation.

Diminishing mineral oil reserves and increasing fuel prices are leading to a growing interest in the replacement of fuels produced on the basis of mineral oil by economical and environmentally friendly alternatives. Processes for the production of fuels from biogenic fat- or oil-containing starting 20 mixtures and used oils obtained, for example, in restaurants and animal fats have long been known, rapeseed oil currently predominantly being used in Central Europe as a starting material in the production of biogenic fuels. Biogenic oils and fats themselves are less suitable as engine fuel since they have 25 to be purified beforehand by generally complicated methods. These include the removal of lecithins, carbohydrates and proteins, the removal of the so-called oil sludge and the removal of the free fatty acids present in relatively large amounts, for example, in rapeseed oil. Vegetable oils treated 30 in this manner nevertheless differ from the technical properties of conventional diesel fuels in several respects. Thus, they have as a rule a higher density than diesel fuel, the cetane number of rapeseed oil is lower than that of diesel fuel and the viscosity is several times higher compared with that of diesel 35 fuel. This leads to an unacceptable deterioration in the fuel properties, such as to nonuniform running behavior of the engine, to substantially increased noise emission and, owing to the higher viscosity, to poorer atomization and combustion in the combustion chamber. In conventional engines, the use 40 of pure vegetable oils therefore leads to coking, associated with increased particle emission. It is known that these problems can be solved by converting the triglycerides (fatty acid esters of glycerol) present in the biogenic oil and fat starting mixtures into monoalkyl esters of fatty acids, in particular 45 methyl or ethyl esters. These esters, also referred to as "biodiesel", can as a rule be used in diesel engines without major retrofits, it often even being possible to reduce the emission of uncombusted hydrocarbons and soot particles in comparison with normal diesel fuel. The transesterification of 50 the triglycerides for biodiesel production also results in glycerol ($\approx 10\%$), which, for reasons of both cost-efficiency and sustainability, should be utilized. There is therefore a need for effective and economical processes which also permit utilization of the glycerol obtained in biodiesel production. These 55 processes should in particular also be suitable for the utilization of further glycerol streams available on an industrial scale.

U.S. Pat. No. 2,360,844 describes a process for the preparation of soaps, in which a crude glyceride is transesterified $_{60}$ with C₁-C₄-alkanols and the glycerol liberated is separated from the monoalkyl esters. The utilization of the glycerol obtained is not described.

U.S. Pat. No. 5,354,878 describes a process for the preparation of lower alkyl esters of higher fatty acids having a low 65 residual glycerol content by transesterification of fatty acid triglycerides and the use of these esters as diesel fuel.

DE 102 43 700 A1 describes a pressureless process for the preparation of alkyl esters of higher fatty acids, in particular biodiesel, from fatty acid triglyceride starting mixtures comprising free fatty acids by a combination of acidic esterification and basic transesterification. The glycerol obtained in the transesterification is partly used as an entraining agent in the esterification of the free fatty acids.

It is known that alcohols having a relatively high hydricity can be converted into alcohols having a lower hydricity by catalytic hydrogenation. Thus, German Patent 524 101 describes such a process in which, inter alia, glycerol is subjected to a gas-phase hydrogenation in the presence of a hydrogenation catalyst with hydrogen in considerable excess. Specifically, copper or cobalt catalysts activated with Cr are used for the hydrogenation of glycerol.

German patent 541 362 describes a process for the hydrogenation of polyoxy compounds such as, for example, glycerol, in the presence of catalysts at elevated temperatures above 150° C. and under superatmospheric pressure. Specifically, the hydrogenation of glycerol using a nickel catalyst at a temperature from 200 to 240° C. and a hydrogen pressure of 100 atm is described.

R. Connor and H. Adkins, in J. Am. Chem. Soc. 54, 1932, pages 4678-4690, describe the hydrogenolysis of oxygencontaining organic compounds, inter alia of 98% strength glycerol, to 1,2-propanediol in the presence of a copperchromium-barium oxide catalyst.

C. Montassier et al., in Bulletin de la Société Chimique de France 1989, No. 2, pages 148-155, describe investigations of the reaction mechanism of the catalytic hydrogenation of polyols in the presence of various metallic catalysts, such as, for example, of glycerol in the presence of Raney copper.

J. Chaminand et al., in Green Chem. 6, 2004, pages 359-361, describe the hydrogenation of aqueous glycerol solutions at 180° C. and 80 bar hydrogen pressure in the presence of supported metal catalysts based on Cu, Pd and Rh.

DE 43 02 464 A1 describes a process for the preparation of 1,2-propanediol by hydrogenation of glycerol in the presence of a heterogeneous catalyst at pressures of from 20 to 300 bar, in particular at from 100 to 250 bar, and temperatures of from 150° C. to 320° C., glycerol in vapor or liquid form being passed over a catalyst bed. Inter alia, copper chromite, copper zinc oxide, copper aluminum oxide and copper silicon dioxide are mentioned as catalysts. The use of glycerol-containing streams from biodiesel production and measures for the pre-treatment of such streams before their use for the hydrogenation are not described in this document.

EP 0 523 015 describes a process for the catalytic hydrogenation of glycerol for the preparation of 1,2-propanediol and 1,2-ethanediol in the presence of a Cu/Zn catalyst at a temperature of at least 200° C. In this process, the glycerol is used as an aqueous solution having a glycerol content of from 20 to 60% by weight, the maximum glycerol content in the working examples being 40% by weight.

WO 2005/095536 describes a low-pressure process for converting glycerol into propylene glycol, in which a glycerol-containing stream having a water content of not more than 50% by weight is subjected to a catalytic hydrogenation at a temperature in the range of from 150 to 250° C. and a pressure in the range of from 1 to 25 bar.

M. A. Dasari et al., in Appl. Chem. A: General 281, 2005, pages 225-231, describe a process for the low-pressure hydrogenation of glycerol to propylene glycol at a temperature of 200° C. and a hydrogen pressure of 200 psi (13.79 bar) in the presence of a nickel, palladium, platinum, copper or copper chromite catalyst. Different reaction parameters were tested, such as, inter alia, the water content of the glycerol

used. It was found that, although the conversion increased with decreasing water content, the highest selectivity was achieved in this low-pressure process at a water content of 20% by weight.

U.S. Pat. No. 5,616,817 describes a process for the prepa-5 ration of 1,2-propanediol by catalytic hydrogenation of glycerol at elevated temperature and superatmospheric pressure, in which glycerol having a water content of not more than 20% by weight is reacted in the presence of a catalyst which comprises from 40 to 70% by weight of cobalt, if appropriate, 10 manganese and/or molybdenum and a low copper content of from 10 to 20% by weight. The temperature is in the range of from about 180 to 270° C. and the pressure in a range of from 100 to 700 bar, preferably from 200 to 325 bar.

It is the object of the present invention to provide a process 15 for the preparation of 1,2-propanediol which permits the hydrogenation of glycerol-containing streams with high selectivity and/or low energy consumption, as required, for example, for separating off water. The process should be suitable in particular for the further processing of glycerol 20 streams obtained on an industrial scale, such as those obtained in the transesterification of fatty acid triglycerides for the preparation of alkyl esters of higher fatty acids.

The invention therefore relates to a process for the preparation of 1,2-propanediol, in which

- a) a glycerol-containing stream is provided and
- b) the glycerol-containing stream is subjected to a hydrogenation in the presence of a copper-containing, heterogeneous catalyst at a temperature of from 100 to 320° C. and a pressure of from 100 to 325 bar.

The hydrogenation product obtained in step b) can, if appropriate, be subjected to at least one working-up step (step c)).

In principle, all glycerol-containing streams, including those from processes carried out industrially and having the 35 purities resulting there, are suitable for use in the process according to the invention. These include in particular glycerol-containing streams from the processing of oil- and/or fat-containing starting materials, for example from soap production, fatty acid and fatty acid ester production, etc. The 40 glycerol-containing stream provided in step a) is preferably a glycerol-containing stream obtained in the preparation of alkyl esters of higher fatty acids by transesterification of fatty acid triglycerides, as obtained in particular in the production of "biodiesel". This embodiment of the process according to 45 the invention is described in more detail below.

The glycerol-containing stream used in step a) preferably has a water content of not more than 30% by weight, preferably of not more than 20% by weight. A water content corresponding to glyceryl monohydrate (water content 16.3% by 50 weight) or less is particularly preferred. In a special embodiment, a glycerol-containing stream which is substantially anhydrous is used. In the context of the present invention, "substantially anhydrous" is understood as meaning a water content of not more than 3% by weight, particularly prefer- 55 ably of not more than 1% by weight. The use of glycerolcontaining streams having a water content in the range of up to 30% by weight, in particular up to 20% by weight, permits the preparation of 1,2-propanediol in high yields and with high selectivity in the temperature and pressure range used for 60 the hydrogenation. The hydrogenation of glycerol-containing streams which are not substantially anhydrous and in particular of streams which have a higher water content than glyceryl monohydrate, is likewise possible in high yields and with high selectivities but, owing to the reduced space-time yield, 65 is less economical. Nevertheless, a water content in the range of from 3 to 30% by weight may be advantageous for the

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rheological properties during the hydrogenation. A special embodiment of the process according to the invention therefore relates to the use of glycerol-containing streams having a water content in the range of from 3 to 30% by weight, preferably from 5 to 20% by weight, for reducing the viscosity during the hydrogenation.

The glycerol-containing streams may have at least one further, preferably glycerol-miscible (and hence as a rule also water-miscible), organic solvent instead of or in addition to water. The glycerol-containing streams provided in step a) preferably have a total solvent content of not more than 20% by weight, particularly preferably not more than 15% by weight, in particular not more than 10% by weight and especially not more than 5% by weight. If solvent mixtures which comprise water and at least one glycerol- or water-miscible organic solvent are used, the proportion of the organic solvent is preferably not more than 50% by weight, particularly preferably not more than 20% by weight, based on the total weight of the solvent. Suitable glycerol-miscible organic solvents are C1-C4-alkanols, such as methanol, ethanol, n-propanol, isopropanol, n-butanol, tert-butanol, polyols and mono- and dialkyl ethers thereof, cyclic ethers, such as dioxane and tetrahydrofuran, etc. Other suitable solvents are aromatic hydrocarbons, such as benzene, toluene or the xylenes. Preferred organic solvents are C1-C4-alkanols, in particular methanol and/or ethanol, and mixtures thereof with water. However, the glycerol-containing streams used in step a) preferably have no organic solvents.

The glycerol-containing streams provided in step a) may be subjected to at least one working-up step. This includes, for example, at least one purification step for removing undesired components. This furthermore includes a reduction of the content of water and/or, if present, organic solvents.

Depending on the origin, the glycerol-containing streams may also comprise inorganic salts as undesired components. These can be removed from the crude glycerol by the working-up processes described below. Thermal working-up (for example with the use of a Sambay evaporator) is particularly suitable for this purpose.

Depending on the origin, the glycerol-containing streams may also comprise catalyst poisons, i.e. components which adversely affect the hydrogenation by deactivating the hydrogenation catalyst. These include, for example, nitrogen-containing compounds, such as amines, and sulfur-containing compounds, such as sulfuric acid, hydrogen-sulfide, thioalcohols, thioethers, e.g. dimethyl sulfide and dimethyl disulfide, carbon oxide sulfide, amino acids, e.g. amino acids comprising sulfur and additional nitrogen groups, fatty acids and salts thereof etc. The catalyst poisons furthermore include halogen compounds, traces of conventional extracting agents, e.g. acetonitrile or N-methylpyrrolidone, etc. and, if appropriate, organic phosphorus and arsenic compounds. A catalyst poison frequently present in glycerol-containing streams from oil and fat refining is sulfuric acid, which is used as a catalyst in the esterification or transesterification.

For example, thermal working-up, preferably distillation, adsorption, ion exchange, a membrane separation method, crystallization or extraction or a combination of two or more of these methods can be used for working up the glycerolcontaining streams in step a). Membrane separation methods with the use of membranes of defined pore sizes are especially suitable for reducing the water content and/or for salt removal. Crystallization is also understood as meaning the partial freezing of the glycerol-containing streams on cooled surfaces. Thus, it is possible to remove impurities which accumulate in the solid phase. In a first embodiment, the glycerol-containing stream in step a) is subjected to a distillation for reducing the water content and/or for removing components which adversely affect the catalytic hydrogenation. This can in principle be effected by conventional distillation methods known to the 5 person skilled in the art. Suitable apparatuses for the distillative working-up comprise distillation columns, such as tray columns, which may be equipped with caps, sieve plates, sieve trays, stacked packings, dumped packings, valves, side take-offs, etc., evaporators, such as thin-film evaporators, 10 falling-film evaporators, forced-circulation evaporators, Sambay evaporators, etc., and combinations thereof. The removal of sulfuric acid takes place even as a result of a simple distillation, in particular a short path distillation.

Suitable separation processes are described in the follow- 15 ing documents: Sattler, Klaus: Thermische Trennverfahren, 3^{rd} edition, Wiley VCH, 2001; Schlünder E. U., Thurner F.: Destillation, Absorption, Extraktion, Springer Verlag, 1995; Mersmann, Alfons: Thermische Verfahrenstechnik, Springer Verlag, 1980; Grassmann P., Widmer F.: Einführung in die 20 thermische Verfahrenstechnik, de Gruyter, 1997; Weiß S., Militzer K.-E., Gramlich K.: Thermische Verfahrenstechnik, Dt. Verlag für Grundstoffindustrie, Leipzig, Stuttgart, 1993. Reference is made here to these documents.

In a further embodiment, the glycerol-containing stream in 25 step a) is subjected to a catalytic desulfurization, if appropriate in the presence of hydrogen, for reducing the contents of sulfur-containing compounds, especially sulfur-containing aromatic compounds. Suitable desulfurization agents comprise a metal component, wherein the metals are preferably 30 selected from metals of groups 6, 7, 8, 9, 10, 11 and 12 of the periodic table of the elements. The metals are selected in particular from Mo, Ni, Cu, Ag, Zn and combinations thereof. Further suitable components of the desulfurization agents are doping agents. The metal component can be employed in 35 oxidized form, reduced form and in form of a mixture that comprises oxidized and reduced constituents. The active components of the desulfurization agents (metal component(s) and optionally doping agent(s)) can be applied on a supporting material. Suitable supports are in principal the 40 adsorbents and the catalyst supports mentioned in the following. Preferably, the supporting material is selected from active carbons, graphites, carbon black, Al₂O₃, SiO₂, TiO₂, ZrO₂, SiC, silicates, zeolithes, argillaceous earth (e.g. bentonite) and combinations thereof. The application of at least one 45 metal component and optionally further components to a supporting material can be carried out by methods known to a person skilled in the art, e.g. by (co)-precipitation or impregnation. The desulfurization agents may be present in form of a geometric body, e.g. in form of spheres, rings, cylinders, 50 cubes, cuboids or other geometric bodies. Unsupported desulfurization agents can be shaped by customary shaping processes, e.g. by extruding, tabletting, etc. The form of supported desulfurization agents is determined by the shape of the support. The desulfurization agents can be employed e.g. 55 in the form of pressed cylinders, tablets, lozenges, wagon wheels, rings, stars or extrudates, such as solid extrudates, polylobal extrudates (e.g. trilobal), hollow extrudates and honeycomb bodies. A preferred desulfurization agent which comprises copper and zinc in an atomic ratio of from 1:0.3 to 60 1:10, preferably from 1:0.5 to 1:3, in particular from 1:0.7 to 1:1.5, is preferably used for the catalytic desulfurization. A desulfurization agent which comprises from 35 to 45% by weight of copper oxide, from 35 to 45% by weight of zinc oxide and from 10 to 30% by weight of alumina is preferably 65 used. In a special embodiment, the desulfurization agent is a component capable of use as a hydrogenation catalyst in step

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b). In this respect, reference is made to the following disclosure of hydrogenation catalysts of the above mentioned composition and processes for their preparation.

In one configuration of this process variant, the glycerolcontaining streams are brought into contact in at least one desulfurization zone with the desulfurization agent and then hydrogenated in at least one reaction zone.

It is self-evident to the person skilled in the art that the specific configuration and arrangement of the desulfurization and reaction zone(s) can be effected in any known manner. It is possible to arrange the desulfurization and reaction zone(s) spatially separate from one another, i.e. to separate them structurally from one another by the configuration of the apparatus or to realize them in one or more common desulfurization/hydrogenation zone(s).

The copper-zinc desulfurization agent can be obtained, for example, by a conventional precipitation or coprecipitation method and used in oxidized as well as in reduced form.

In a particular embodiment, the copper-zinc desulfurization agent comprises at least copper, zinc and aluminum, the copper:zinc:aluminum atomic ratio being in the range of from 1:0.3:0.05 to 1:10:2, preferably from 1:0.6:0.3 to 1:3:1 and in particular from 1:0.7:0.5 to 1:1.5:0.9.

For conversion into the reduced form, it is possible to subject the desulfurization agent to a hydrogen reduction. This is carried out at from about 150 to 350° C., preferably from about 150 to 250° C., in the presence of hydrogen, the hydrogen being diluted by an inert gas, such as, for example, nitrogen, argon, or methane, in particular nitrogen, so that the hydrogen content is 10% by volume or less, preferably 6% by volume or less, in particular from 0.5 to 4% by volume. The copper-zinc desulfurization agent thus obtained ("reduced form") can be used in this form in the desulfurization.

In an embodiment, the desulfurization of the glycerolcontaining stream is carried out over the copper-zinc desulfurization agent in oxidized form without addition of hydrogen.

In a further embodiment, the desulfurization of the glycerol-containing stream is carried out over the copper-zinc desulfurization agent in oxidized form in the presence of hydrogen.

In a further embodiment, the desulfurization of the glycerol-containing stream is carried out over the copper-zinc desulfurization agent in reduced form without addition of hydrogen.

In a further embodiment, the desulfurization of the glycerol-containing stream is carried out over the copper-zinc desulfurization agent in reduced form in the presence of hydrogen.

Usually, the desulfurization is carried out in a temperature range of from 40 to 200° C., in particular at from 50 to 180° C., especially at from 60 to 160° C., preferably at from 70 to 120° C., at a pressure of from 1 to 40 bar, in particular at from 1 to 32 bar, preferably at from 1.5 to 5 bar, especially at from 2.0 to 4.5 bar. The desulfurization can be carried out in the presence of inert gases, such as, for example, nitrogen, argon or methane. As a rule, however, the desulfurization is carried out without addition of inert gases.

Usually—if desired—hydrogen having a purity of \geq 99.8% by volume, in particular of \geq 99.9% by volume, preferably of \geq 99.95% by volume, is used here. These purities apply analogously to the hydrogen which is used in the catalyst activations carried out if appropriate.

Usually, the weight ratio of glycerol-containing stream to hydrogen is in the range of from 40 000:1 to 1000:1, particularly in the range of from 38 000:1 to 5000:1, in particular in

the range of from $37\,000:1$ to $15\,000:1$, preferably in the range of from $36\,000:1$ to $25\,000:1$, especially in the range of from $35\,000:1$ to $30\,000:1$.

The glycerol-containing stream thus desulfurized generally has a content of sulfur-containing impurities, especially of aromatic sulfur compounds of not more than 70 ppb, preferably of not more than 50 ppb and the total sulfur content is ≤ 200 ppb, preferably ≤ 150 ppb, in particular ≤ 100 ppb altogether.

The desulfurization agents described above also make it 10 possible to reduce or to remove chlorine, arsenic and/or phosphorus or corresponding chlorine, arsenic- and/or phosphorus-containing compounds from the aromatic hydrocarbon or from the mixture of aromatic hydrocarbons.

In a further embodiment, the glycerol-containing stream in 15 step a) is brought into contact with at least one adsorbent for removing components which adversely affect the catalytic hydrogenation.

The adsorbents generally have a specific surface area, determined according to BET, in the range of from about 10 to 20 $2000 \text{ m}^2/\text{g}$, preferably in the range of from 10 to $1500 \text{ m}^2/\text{g}$, more preferably in the range of from 10 to $400 \text{ m}^2/\text{g}$, especially in the range of from 60 to $250 \text{ m}^2/\text{g}$.

Suitable adsorbents are, for example, active aluminas. They are prepared, for example, starting from aluminum 25 hydroxide, which is obtainable from aluminum salt solutions by conventional precipitation methods. Active aluminas suitable for the process according to the invention are also obtainable starting from aluminum hydroxide gels. For the preparation of such gels, for example, precipitated aluminum 30 hydroxide can be activated by conventional working-up steps, such as filtration, washing and drying, and then, if appropriate, milled or agglomerated. If desired, the resulting alumina can then also be subjected to a shaping method, such as extrusion, granulation, tabletting, etc. Suitable adsorbents 35 are preferably the SelexsorbTM types from Alcoa.

Suitable adsorbents are furthermore alumina-containing solids. These include, for example, the so-called clays, which likewise have aluminas as the main constituent.

Other suitable adsorbents are aluminum phosphates.

Other suitable adsorbents are silicas, which are obtainable, for example, by dehydration and activation of silica gels. A further process for the preparation of silica is the flame hydrolysis of silicon tetrachloride, it being possible to vary the desired surface properties of the resulting silica in wide 45 ranges by suitable variations of the reaction parameters, such as, for example, of the stoichiometric composition of the starting mixture and of the temperature.

Other suitable adsorbents are kieselguhrs, which likewise have silicas as the main constituent. These include, for 50 example, the diatomaceous earth obtained from silicic sediments.

Other suitable adsorbents are titanium dioxides and zirconium dioxides, as described, for example, in Römpp, Chemie-Lexikon, 9^{th} edition (paperback), vol. 6, page 4629 et 55 seq. and page 5156 et seq. and the literature cited there. Reference is made here to these in their entirety.

Other suitable adsorbents are phosphates, in particular condensed phosphates, such as, for example, fused or calcined phosphates, which have a large active surface area. 60 Suitable phosphates are described, for example, in Römpp, Chemie-Lexikon, 9^{th} edition (paperback) vol. 4, page 3376 et seq. and the literature cited there. Reference is made here to this in its entirety.

Other suitable adsorbents are carbon-containing adsor- 65 bents, preferably active carbon. Active carbon is understood here in general as meaning carbon having a porous structure

and large internal surface area. For the preparation of active carbon, vegetable, animal and/or mineral carbon-containing raw materials are heated, for example, with dehydrating agents, such as zinc chloride or phosphoric acid, or are carbonized by dry distillation and then oxidatively activated. For this purpose, for example, the carbonized material can be treated at elevated temperatures of from about 700 to 1000° C. with steam, carbon dioxide and/or mixtures thereof.

Use of ion exchangers and/or adsorber resins is also possible.

The adsorbents are preferably selected from titanium dioxides, zirconium dioxides, silicas, kieselguhr, aluminas, alumina-containing solids, aluminum phosphates, natural and synthetic aluminum silicates, phosphates, carbon-containing adsorbents and mixtures thereof.

The adsorbents generally have a specific surface area, determined according to BET, in the range of from about 10 to $2000 \text{ m}^2/\text{g}$, in particular in the range of from 10 to $1500 \text{ m}^2/\text{g}$ and especially in the range of from 20 to $600 \text{ m}^2/\text{g}$.

For the adsorptive removal of undesired components, in particular of components which adversely affect the catalytic hydrogenation, the glycerol-containing stream in step a) is brought into contact with at least one adsorbent in an adsorption zone.

In a special embodiment, an adsorbent which comprises at least one component also capable of use as a hydrogenation catalyst in step b) is used. The hydrogenation catalysts described in more detail below are referred to here in their entirety. Combinations of two or more than two adsorbents are also suitable for use as adsorbents. It is possible to use either exclusively components also capable of being hydrogenation catalysts, exclusively adsorbents not suitable as hydrogenation catalysts or combinations thereof.

In a preferred embodiment, the same component is used as adsorbent and as hydrogenation catalyst. If appropriate, one or more further, conventional adsorbents, as described above, differing from the hydrogenation catalyst, are additionally used here.

In a configuration of the process, glycerol-containing streams are brought into contact in at least one adsorption zone with the adsorbent and then hydrogenated in at least one reaction zone.

It is self-evident to the person skilled in the art that the specific configuration and arrangement of the adsorption and reaction zone(s) can be effected in any known manner. It is preferably to arrange the adsorption and reaction zone(s) spatially separate from one another, i.e. to separate them structurally from one another by the configuration of the apparatus.

If different adsorbents are used, for example, a first adsorption zone which comprises a first adsorbent can be provided in a first adsorption zone in a first reactor and separately, i.e. structurally separate therefrom, for example in a second reactor, a second adsorption zone which comprises a second adsorbent. Here, the first and/or the second adsorbent may comprise at least one component capable of use of a hydrogenation catalyst.

In a further embodiment, a conventional adsorbent is used together with an adsorbent capable of hydrogenation in a single adsorption zone, for example, in stratified form, mixed in the form of a random distribution or in the form of a gradient bed. The use in mixed form permits, if appropriate, better control of the temperature. In the case of a gradient bed, linear and non-linear gradients can be used. It may be advantageous here to implement the distribution within the bed in such a way that the glycerol-containing stream to be hydro-

genated is first brought into contact with the conventional adsorbent before it is brought into contact with the adsorbent capable of hydrogenation.

Advantageously, at least two adsorption zones will be arranged in such a way that the glycerol-containing stream to 5 be hydrogenated is brought into contact with a conventional adsorbent in the first adsorption zone and is brought into contact, in the second adsorption zone, with an adsorbent which comprises at least one component capable of use as a hydrogenation catalyst.

The glycerol-containing streams provided in step a) of the process according to the invention preferably originate from the production of biodiesel. In the context of the present invention, "biodiesel" is understood as meaning a mixture of fatty acid monoalkyl esters which can be obtained from bio-15 genic oil- and/or fat-containing starting mixtures and can be used as fuel in diesel engines.

In principle, all available biogenic oil- and/or fat-containing starting mixtures are suitable for providing the glycerolcontaining stream. Oils and fats are generally solid, semisolid 20 or liquid fatty acid triglycerides, in particular from vegetable and animal sources, which chemically substantially comprise glyceryl esters of higher fatty acids. Suitable higher fatty acids are saturated or mono- or polyunsaturated fatty acids having preferably 8 to 40, particularly preferably 12 to 30, 25 carbon atoms. These include, for example, n-nonanoic acid, n-decanoic acid, n-undecanoic acid, n-tridecanoic acid, myristic acid, pentadecanoic acid, palmitic acid, margaric acid, nonadecanoic acid, arachidic acid, behenic acid, lignoceric acid, cerotic acid, melissic acid, palmitoleic acid, oleic 30 acid, linoleic acid, linolenic acid, stearic acid, elaostearic acid, etc.

Vegetable fats and oils are substantially based on fatty acids having an even number of carbon atoms, whereas animal fats and oils may also comprise fatty acids having an odd 35 number of carbon atoms, in free form or bound as triglyceride esters. The unsaturated fatty acids occurring in vegetable fats and oils are present in the cis form, while animal fatty acids frequently have a trans configuration.

In principle, used or unused, unpurified or purified veg- 40 etable, animal or industrial oils or fats or mixtures thereof can be used for providing the glycerol-containing stream in step a). These may comprise proportions of further ingredients, for example free fatty acids. The proportion of free fatty acids is in general from 0% to 50%, e.g. from 0.1 to 20%, of the 45 starting mixture used for the transesterification of the fatty acid triglycerides. Free fatty acids can, if desired, be removed before or after the transesterification of the fatty acid triglycerides. Salts of these fatty acids (for example the alkali metal salts) can be converted into the free acid beforehand by acidi- 50 fication with a strong acid, e.g. HCl. The isolation of the free fatty acids is effected, for example, by centrifuging. Preferably, the free fatty acids present in the starting mixture are likewise converted into the alkyl esters. This can be effected before, during or after the transesterification of the fatty acid 55 triglycerides.

Used fats and oils suitable for providing the glycerol-containing stream in step a) are fat- and/or oil-containing components which, after their recovery from appropriate biogenic starting materials were first used for other purposes, for 60 example for technical purposes or purposes for food production, and may be chemically modified or unmodified as a result of this use or may have additional ingredients which in particular are associated with this use. These can, if desired, be at least partly removed by transesterification before the use 65 for providing the glycerol-containing stream. Unused fats and oils suitable for providing the glycerol-containing stream in

step a) are fat- or oil-containing components which still have not been used for any other purpose after their recovery from the appropriate vegetable or animal starting materials and which therefore have only ingredients which originate from the starting materials or are associated with the recovery from the starting materials. Ingredients other than fatty acid triglycerides (and, if appropriate, free fatty acids) can, if desired, also be at least partially removed from these starting materials by transesterification before the use for providing the glycerol-containing stream.

For the purification and/or enrichment, the unused or used fats or oils can be subjected to removal of undesired ingredients, such as lecithins, carbohydrates, proteins, oil sludge, water, etc.

Vegetable oils and fats are those which originate predominantly from vegetable starting materials, such as seeds, roots, leaves or other suitable plant parts. Animal fats or oils originate predominantly from animal starting materials, such as animal organs, tissues or other body parts or body fluids, such as milk. Industrial oils and fats are those which were obtained in particular from animal or vegetable starting materials and treated for technical purposes. The used or unused, unpurified or purified oils and/or fats used according to the invention are selected in particular from the group consisting of soapstock, brown grease, yellow grease, industrial tallow, industrial lard, frying oils, animal fat, edible tallow, crude vegetable oils, crude animal oils or fats or mixtures thereof.

"Soapstock" is understood as meaning a byproduct obtained in the processing of vegetable oils, in particular a byproduct of edible oil refineries which is based on soybean, colza or sunflower oil. Soapstock has a proportion of from about 50% to 80% of free fatty acids.

"Brown grease" is understood as meaning an animal fatcontaining waste product which has a proportion of from more than 15% to 40% of free fatty acids. "Yellow grease" comprises from about 5% to 15% of free fatty acids.

"Industrial tallow" and "industrial lard" are understood as meaning animal fats which are produced for industrial purposes and are obtained after the drying or wet melting process, for example from slaughter wastes. Industrial tallows are rated and handled according to their acid number, the content of free fatty acids being, for example, between 1 and 15 to 20% by weight and in some cases even higher, depending on origin.

The "animal fats" include in particular fat-containing waste products obtained in the utilization of poultry, cattle, pig, fish and marine mammal bodies, for example solar stearin, a solid residue which remains after lard oil has been forced out of pork lard.

The glycerol-containing stream in step a) is preferably provided from crude vegetable oils as starting material. It is possible to start from unpurified crude vegetable oils, i.e. from liquid or solid compositions which are obtained from vegetable starting materials, for example by pressing, these having undergone no other treatment than settling in generally customary periods and centrifuging or filtering, in which only mechanical forces, such as gravitational force, centrifugal force or pressure, are used for separating the oil from solid constituents. Such unpurified crude vegetable oils may also be vegetable oils obtained by extraction if the properties thereof do not differ, or differ only insignificantly, from the corresponding vegetable oils obtained by means of pressing. The proportion of free fatty acids in unpurified vegetable fats and oil differs and is, for example, from about 0 to 20%, such as, for example from 0.1 to 15%.

Before they are used for the transesterification, the vegetable oils can of course be subjected to one or more working-

up steps, as described in more detail below. Thus, purified vegetable oils, for example raffinates or semiraffinates, of the abovementioned vegetable oils may also be used as starting materials.

A vegetable oil or fat which is preferably selected from 5 rapeseed oil, palm oil, colza oil, soybean oil, sunflower oil, corn oil, cottonseed oil, palm kernel and coconut fat and mixtures thereof is preferably used for providing the glycerol-containing stream in step a). Particularly preferably used are rapeseed oil or a mixture containing rapeseed-oil.

Animal oil or fat which is preferably selected from milk fat, wool fat, beef tallow, pork lard, fish oils, blubber, etc. and mixtures thereof is also suitable for providing the glycerolcontaining stream in step a). Before they are used for the transesterification, these animal fats or oils, too can be sub- 15 jected to one or more working-up steps, as described in more detail below.

Preferably, the provision of the glycerol-containing stream in step a) comprises the following steps:

- a1) provision of a biogenic fat- and/or oil-containing starting 20 mixture,
- a2) transesterification of the fatty acid triglycerides present in the starting mixture with at least one C1-C9-monoalcohol and, if appropriate, esterification of the free fatty acids present in the starting mixture with formation of an esteri- ²⁵ fication mixture,
- a3) separation of the esterification mixture to obtain at least one fraction enriched with biodiesel and at least one fraction enriched with glycerol liberated in the esterification,
- a4) if appropriate, purification of the fraction enriched with glycerol.

Step a1)

In a preferred embodiment, the provision of the biogenic fat- and/or oil-containing starting mixture in step a1) com- 35 prises at least one purification step. For the purification, the fat- and/or oil-containing starting mixture can be subjected to at least one purification process usually used for fats and oils, such as clarification, filtration, treatment with bleaching earths or treatment with acids or alkali for separating off 40 troublesome impurities, such as proteins, phosphatides and slimes, and a combination of at least two of these purification steps.

Step a2)

At least one C1-C2-monoalcohol, in particular at least one C_1 - C_4 -monoalcohol is preferably used for the transesterification of the fatty acid triglycerides. The use of methanol or ethanol is preferred.

The transesterification of the fatty acid triglyceride can be effected by acidic or preferably basic catalysis. Suitable acids are, for example, mineral acids, such as HCl, H₂SO₄ or H₃PO₄.

At least one base is preferably used as the catalyst. Said base is preferably selected from alkali metal hydroxides, such 55 as NaOH and KOH, alkaline earth metal hydroxides, such as Ca(OH)₂, alkali and alkaline earth metal C₁-C₆-alkanolates, such as NaOCH₃, KOCH₃, Na(OCH₂CH₂) and Ca(OCH₂CH₂)₂ and mixtures thereof. NaOH, KOH or NaOCH₃ is particularly preferably used, very particularly ₆₀ preferably NaOCH₃.

The amount of base used is usually in the range of from 0.1 to 10% by weight, in particular from 0.2 to 5% by weight, based on the amount of fatty acid triglycerides used.

The base is preferably used in the form of an aqueous or 65 alcoholic, particularly preferably alcoholic, solution. The solvent already used for the alcoholysis of the triglycerides is

advantageously used as a solvent for the base. NaOCH₃ solution in methanol is preferably used for the transesterification. The transesterification is preferably effected at a tempera-

ture from about 20 to 150° C., in particular from 30 to 95° C.

The transesterification is effected in apparatuses customary for this purpose and known to the person skilled in the art. In a suitable embodiment, the transesterification is effected continuously. The transesterification is preferably effected in at least one column, the transesterification mixture obtained simultaneously being subjected to a separation. In general, a higher-boiling phase which with enriched with the basic catalyst, with unconverted monoalcohol and with the glycerol formed in the transesterification is obtained and a lowerboiling phase which is enriched with the transesterification product is obtained. If the transesterification product still contains triglycerides which have not undergone transesterification, these can also be separated off and subjected to a further transesterification in the first or a further transesterification stage.

The last transesterification mixture is then transferred to a drying unit, residual amounts of water again being removed. After the drying in the drying apparatus, the desired end product biodiesel is present in purified form and can be used directly as fuel.

If the fat- and/or oil-containing starting mixture used for providing the glycerol-containing stream in step a) comprises free fatty acids, these can preferably be subjected to an esterification for conversion into esters suitable for biodiesel.

The free fatty acids are preferably transesterified with the same C1-Co-monoalcohol which was used for the transesterification of the fatty acid triglycerides. The esterification of free fatty acids can be effected before, during or after the transesterification of the fatty acid triglycerides. In a preferred embodiment the esterification of free fatty acids is effected before the transesterification of the fatty acid triglycerides.

The esterification of the free fatty acids can be effected by basic or preferably acidic catalysis. Suitable acids are the abovementioned mineral acids, such as HCl, H₂SO₄ or H₃PO₄, p-toluene sulfonic acid, etc. The esterification is preferably effected at a temperature of from about 20 to 95° C., in particular from 40 to 80° C.

The esterification is effected in apparatuses customary for this purpose and known to the person skilled in the art. These include stirred vessels and/or columns which, if desired, are connected to form cascades. The esterification of the free fatty acids is preferably effected in at least one esterification apparatus designed as a column, the esterification mixture obtained simultaneously being subjected to a separation. In a suitable embodiment, the esterification is effected in the presence of an entraining agent for facilitating the separation.

Step a3)

During or after the transesterification and/or esterification, the esterification mixture is subjected to a separation to obtain at least one fraction enriched with C1-C9-monoalcohol esters and at least one fraction enriched with glycerol liberated in the transesterification. The separation is preferably effected by conventional distillation methods known to the person skilled in the art. Suitable distillation apparatuses are those mentioned above.

Step a4)

The fraction obtained after separation of the esterification mixture in step a3) and enriched with glycerol can, if appropriate, be subjected to at least one working-up step. This includes, for example, the removal of undesired components, such as salts, and of components which adversely affect the catalytic hydrogenation or the removal of water and, if present, organic solvent. Reference is made to the above statements on these working-up steps, in their entirety.

The catalysts used in the process according to the invention may be unsupported catalysts or supported catalysts. They 5 can be used in form of uniform-composition catalysts, impregnated catalysts, coated catalyst and precipitated catalysts.

In principle, a large number of copper-containing catalysts which may additionally comprise at least one further element 10 of main group I, II or III, IV, V, or sub group I, II, IV, V, VI, VII, or VIII and of the lanthanides (IUPAC: groups 1 to 15 and the lanthanides) are suitable, in particular Ca, Mg, Al, La, Ti, Zr, Cr, Mo, W, Mn, Ni, Co, Zn and combinations thereof.

A special embodiment of catalysts which are particularly 15 advantageous for use in the process according to the invention comprises skeletal or metal sponge catalysts, such as those referred to as "Raney catalysts". These include in particular Raney copper and copper-containing metal alloys in the form of a Ranev catalyst. Ranev catalysts whose metal component 20 comprises at least 95%, in particular at least 99%, of copper are preferred. Processes for the preparation of Raney catalysts are known to the person skilled in the art and are described, for example, in DE-A-43 35 360, DE-A-43 45 265, DE-A-44 46 907 and EP-A-842 699. Raney copper can be prepared in 25 a manner known per se by treating copper-aluminum alloys with alkali metal hydroxides. A Raney catalyst suitable for use in the process according to the invention is obtainable, for example, by preparation of a mixture of at least one coppercontaining catalyst alloy and at least one binder, the catalyst 30 alloy comprising copper and, if appropriate, at least one further catalytically active catalyst metal and a leachable alloy component, if appropriate with addition of moistening agents and/or additives, such as molding assistants, lubricants, plasticizers and/or pore formers, homogenization of this mixture 35 and molding to give the desired molding, calcination of the molding and activation of the catalyst precursor thus obtained, by partial or complete leaching out of the leachable alloy component and, if appropriate, final washing of the prepared catalyst. 40

A further special embodiment of catalysts which are particularly advantageously used in the process according to the invention comprises catalysts which comprise copper in oxidic form and, if appropriate, additionally in elemental form. The hydrogenation catalyst used in step b) then prefer-45 ably comprises at least 23% by weight, particularly preferably at least 35% by weight, of copper in oxidic and/or elemental form, based on the total weight of the catalyst.

A frequently used process for the preparation of such catalysts comprises the impregnation of support materials with 50 solutions of the catalyst components, which are then converted into the catalytically active state by thermal treatment, decomposition or reduction.

A further suitable process for the preparation of catalysts comprises the precipitation of a catalyst component or the 55 coprecipitation of two or more catalyst components. Thus, a copper compound, optionally at least one further metal compound and/or an additive are precipitated and subjected to subsequent drying, calcination and shaping to produce a shaped catalyst body. The precipitation can be performed in 60 the presence of a support material. Suitable starting materials for the precipitation are metal salts and metal complexes. As copper compound for the precipitation it is in principle known to use all Cu(I) and/or Cu(II) salts which are soluble in the solvents used for application to the support, for example 65 nitrates, carbonates, acetates, oxalates or ammonium complexes. Particular preference is given to using copper nitrate.

The catalytically active component of the catalyst may further comprise, apart from a copper compound, other elements as additive components, e.g. metals, nonmetals and their compounds. These preferably include a metal of groups 4 to 15 and the lanthanides. These preferably include metals as La, Ti, Zr, Cu, Mo, W, Mn, Re, Co, Ni, Cu, Ag, Au, Zn, Sn, Pb, As, Sb and Bi. Preferably, an aqueous medium is used for the precipitation.

Suitable aqueous media are substances or mixtures which are liquid under the process conditions and contain at least 10% by weight, preferably at least 30% by weight and in particular at least 50% by weight, of water. The part other than water is preferably selected from among inorganic or organic substances which are at least partially soluble in water or at least partially miscible with water. For example, the substances other than water are selected from among organic solvents, C1-C22-alkanols, in particular methanol, ethanol, n-propanol, isopropanol, n-butanol, sec-butanol, tert-butanol, pentanoles and hexanoles, C4-C8-cycloalkyl ethers, such as tetrahydrofurans, pyrans, dioxanes and trioxanes, C1-C12-dialkyl ethers, such as dimethyl ether, dibutyl ether and methyl butyl ether. The aqueous medium preferably contains less than 40%, in particular less than 30% and particularly preferably less than 20%, of organic solvent. In preferred embodiments of the process of the present invention, the aqueous medium is essentially free of organic solvents.

Precipitation can be induced by known methods, e.g. cooling a saturated solution, adding a precipitating agent, etc. Suitable precipitating agents are e.g. acids, bases, reducing agents, etc.

Precipitation can be induced by addition of an acid or a base to the aqueous medium containing the copper compound and optionally further compounds. Suitable acids are mineral acids, like HCl, H_2SO_4 and H_3PO_4 . The base is preferably selected from among metal oxides, metal hydroxides, in particular alkali metal hydroxides, such as sodium hydroxide and potassium hydroxide, metal carbonates, in particular alkali metal and alkaline earth metal carbonates, e.g. lithium carbonate, sodium carbonate, nitrogen bases, in particular ammonia, primary, secondary and tertiary amines.

Examples of suitable reducing agents are carboxylic acids, such as formic acid, citric acid, lactic acid, tartaric acid and in particular the salts of carboxylic acids, preferably the alkali metal, alkaline earth metal, ammonium and C1-C10-alkylammonium salts, phosphorus or hypophosphorus acid, the salts of phosphorus or hypophosphorus acid, in particular the alkali metal or alkaline earth metal salts, C1C10-alkanols, such as methanol, ethanol and isopropanol, sugars, such as aldoses and ketoses in the form of monosaccharides, disaccharides and oligosaccharides, in particular glucose, fructose and lactose, aldehydes, such as formaldehyde, boron-hydrogen compounds, such as boron hydrides, boranes, metal boranates and borane complexes, e.g. diborane, sodium borohydride and aminoboranes, in particular trimethylaminoborane, hydrazine and alkylhydrazines, such as methylhydrazine, hydrogendithionites and dithionites, in particular sodium and potassium hydrogendithionites, sodium, potassium and zinc dithionites, hydrogensulfites and sulfites, in particular sodium and potassium hydrogensulfites, sodium, potassium and calcium sulfites, hydroxylamine and urea, and also mixtures thereof.

For example, catalysts which comprise nickel and copper, in addition to other metals, as active constituents on a silica support are suitable for the hydrogenation. Such catalysts are described, for example, in DE-A 26 28 987. The active material of these catalysts comprises in particular from 40 to 80% by weight of nickel, from 10 to 50% by weight of copper and from 2% to 10% by weight of manganese.

EP-A-0 434 062 describes hydrogenation catalysts which are obtainable by reduction of a precursor comprising oxides of copper, of aluminum and at least of one further metal ⁵ selected from magnesium, zinc, titanium, zirconium, tin, nickel and cobalt.

The hydrogenation catalysts which are described in DE 102 18 849 and comprise from 0.1 to 10% by weight of chromium, calculated as Cr_2O_3 , from 0.1 to 10% by weight of ¹⁰ calcium, calculated as CaO_x and from 5 to 20% by weight of copper, calculated as CuO, deposited on a silica support material and based in each case on the total weight of the calcined catalyst, are also suitable.

DE-A-40 21 230 discloses copper/zirconium oxide cata-¹⁵ lysts, the ratio of copper atoms to zirconium atoms, expressed as a weight ratio, being from 1:9 to 9:1.

DE-A-4 028 295 describes copper/manganese hydrogenation catalysts.

EP-A-552463 describes hydrogenation catalysts in a first embodiment, the oxidic form substantially corresponding to the composition $Cu_aAl_bZr_cMn_aO_x$, the following relationships being applicable: a>0; b>0; c>/=0; d>0; a>b/2; b>a/4; a>c; a>d;

and x is the number of oxygen ions which is required for preserving the electroneutrality per formula unit. According to a further embodiment, the catalyst according to the invention comprises a smaller proportion of alumina. The catalyst according to this embodiment substantially corresponds to ³⁰ the composition $Cu_aAl_bZr_cMn_dO_x$, the following relationships being applicable: a>0; b=a/40 to a/4; c>/=0; d>0; a>c; a=0.5d to 0.95d and x is the number of oxygen ions which is required for preserving the electroneutrality per formula unit.

WO 2006/005505 discloses moulded catalyst bodies that ³⁵ are particularly advantageous for use in the process according to the invention. Those catalysts can be produced by a process in which

- (i) an oxidic material comprising copper oxide, aluminum oxide and at least one of the oxides of lanthanum, tungsten, ⁴⁰ molybdenum, titanium or zirconium, with preference being given to the oxides of lanthanum and/or tungsten, is made available,
- (ii) pulverulent metallic copper, copper flakes, pulverulent cement or a mixture thereof or a mixture thereof with graphite can be added to the oxidic material, and
- (iii) the mixture resulting from (ii) is shaped to form a catalyst pellet or a catalyst extrudate having a diameter d and/or a height h of <2.5 mm, catalyst spheres having a diameter d of <2.5 mm or catalyst honeycombs having a cell diameter r_z of <2.5 mm.

Among the oxides of lanthanum, tungsten, molybdenum, titanium or zirconium, lanthanum oxide is preferred. The composition of the oxidic material is generally such that the proportion of copper oxide is in the range from 40 to 90% by weight, the proportion of oxides of lanthanum, tungsten, molybdenum, titanium or zirconium is in the range from 0 to 50% by weight and the proportion of aluminum oxide is up to 50% by weight, in each case based on the total weight of the abovementioned oxidic constituents, with these three oxides together making up at least 80% by weight of the oxidic material after calcination and cement not being included as part of the oxidic material in the above sense.

In a preferred embodiment, the oxidic material comprises $_{65}$ (a) copper oxide in a proportion in the range $50 \le x \le 80\%$ by

weight, preferably $55 \le x \le 75\%$ by weight,

(b) aluminum oxide in a proportion in the range 15≦y≦35% by weight, preferably 20≦y≦30% by weight, and

(c) at least one of the oxides of lanthanum, tungsten, molybdenum, titanium or zirconium, preferably of lanthanum and/or tungsten, in a proportion in the range 2≦z≦20% by weight, preferably 3≦z≦15% by weight,

in each case based on the total weight of the oxidic material after calcination, where $80 \le x+y+z \le 100$, in particular $95 \le x+y+z \le 100$.

Preferred catalysts comprise the following metals in oxidic form, reduced form (elemental form) or a combination thereof. Metals that are stable in more than one oxidation state can be employed entirely in one of the oxidation states or a combination of different oxidation states:

Cu

Cu, Ti Cu, Zr

Cu, Mn

 C_{11} Ni M

Cu, Al, at least one further metal selected from La, W, Mo, Mn, Zn, Ti, Zr, Sn, Ni, Co

Cu, Zn, Zr

Cu, Cr, C

Especially preferred catalysts comprise the following metals:

Cu

Cu, Al

Cu, Al, La

Cu, Al, Zn

⁵ Cu, Zn, Zr

Cu, Al, Mn

Cu, Cr, C

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Virtually all support materials of the prior art, as advantageously used in the preparation of supported catalysts, for example, SiO₂ (quartz), porcelain, magnesium oxide, tin dioxide, silicon carbide, TiO₂ (rutile, anatas), ZrO₂, Al₂O₃ (alumina), aluminum silicate, steatite (magnesium silicate), zirconium silicate, cerium silicate or mixtures of these support materials, may be used as inert support material for the catalysts according to the invention. Preferred support materials are alumina and silica. Silica materials of different origin and preparation, for example pyrogenically produced silicas or silicas produced by a wet chemical method, such as silica gels, aero gels or precipitated silicas, can be used as silica support material for the catalyst preparation (for the preparation of various SiO₂ starting materials cf.: W. Büchner; R. Schliebs; G. Winter; K. H. Büchel: Industrielle Anorganische Chemie; 2nd edition, pages 532-533, VCH Verlagsgesellschaft, Weinheim 1986).

The catalysts may be present in the form of a geometric body, e.g. in form of spheres, rings, cylinders, cubes, cuboids or other geometric bodies. Unsupported catalysts can be shaped by customary processes, e.g. by extruding, tabletting, etc. The form of supported catalysts is usually determined by the shape of the support. In an alternative, the support can be subjected to a shaping process prior to or after application of the catalystically active compound(s) or a precursor thereof. The catalysts can be employed e.g. in the form of pressed cylinders, tablets, lozenges, wagon wheels, rings, stars or extrudates, such as solid extrudates, polylobal extrudates (e.g. trilobal), hollow extrudates and honeycomb bodies.

²⁰ Cu, Mil

Cu, Ni, Mn

²⁵ Cu, Cr, Ca

Cu, Al, Mn, optionally Zr

Cu, Ti

Cu, Ai, Mii

The catalyst particles generally have a mean value of the (largest) diameter of from 0.5 to 20 mm, preferably from 1 to 10 mm. These include, for example, catalysts in the form of tablets, for example having a diameter of from 1 to 7 mm, preferably 2 to 6 mm, and a height of from 3 to 5 mm, rings 5 having, for example an external diameter of from 4 to 7 mm, preferably 5 to 7 mm, a height of from 4 to 7 mm, preferably 2 to 5 mm, and a hole diameter of from 2 to 3 mm, or strands of different lengths having a diameter of, for example, from 1.0 to 5 mm. Such shapes can be obtained in a manner known 10 per se, by tabletting, extrusion molding or extrusion. For this purpose, conventional adjuvants, for example lubricants, such as graphite, polyethylene oxide, cellulose or fatty acids (such as stearic acid), and/or molding assistants and reinforcing agents, such as fibers of glass, asbestos or silicon carbide 15 can be added to the catalyst material.

A special embodiment of supported catalysts comprises coated catalysts. Coated catalysts are also preferably suitable for the process according to the invention. Coated catalysts comprise a catalytic material applied in the form of a coat to 20 a support. They may be present in the form of spheres, rings, cylinders, cubes, cuboids or other geometrical bodies. Regardless of the type and composition of the catalytically active material, coated catalyst particles can be provided in principle by bringing the support into contact with a liquid 25 binder and the catalytically active material, applying a layer of the material to the support and then, if appropriate, partially removing the binder. In order to provide the catalyst particles, the catalytically active material is applied already in its prepared catalytically active form, for example as calcined 30 mixed oxide. Suitable processes for the preparation of coated catalysts are described, for example, in DE-A-29 09 671 and in EP-A-714 700. According to the last-mentioned process, the support is first moistened with the liquid binder, a layer of active catalyst material is then bonded to the surface of the 35 moistened support body by bringing into contact with dry, finely divided, active catalyst material, and, if appropriate, the liquid binder is then partly removed. In a special embodiment, the steps of moistening of the support, bringing into contact with the catalyst material and removal of the liquid binder are 40 2:1 to 500:1, preferably from 3:1 to 100:1. repeated once or several times until the desired layer thickness of the coated catalyst is reached.

A further special embodiment of supported catalysts comprises catalysts prepared by impregnation methods. For this purpose, the catalytically active catalyst components or pre- 45 cursor compounds thereof can be applied to the support material. In general, aqueous salt solutions of the components, for example aqueous solutions of their halides, sulfates, nitrates, etc. are applied for impregnating the support material. The copper component can also be applied, for example, in the 50 form of an aqueous solution of its amine complex salts, for example as $[Cu(NH_3)_4]SO_4$ or as $[Cu(NH_3)_4](NO_3)_2$ solution, if appropriate in the presence of sodium carbonate, to the support material. Of course, copper-amine complexes other than those mentioned by way of example can also be used 55 with the same success for the catalyst preparation.

The impregnation of the support material with the precursor compounds of the catalytically active components can be effected in principle in one stage or in a plurality of stages. The impregnation can be carried out in conventional impreg- 60 nation apparatuses, for example impregnation drums. After drying and/or calcination, the prepared catalyst is then obtained. The drying of the impregnated catalyst moldings can be effected continuously or batchwise, for example in belt or tray furnaces. The drying can be effected at atmospheric 65 pressure or reduced pressure. Furthermore the drying can be effected in a gas stream, for example in an air stream or a

nitrogen stream. Depending on the pressure applied, the drying is generally carried out at temperatures of from 50 to 200° C., preferably from 80 to 150° C. The calcination of the catalyst, dried beforehand if appropriate is effected in general at temperatures of from 200 to 800° C., preferably from 500 to 700° C. The calcination, like the drying, can be carried out continuously or batchwise, for example in belt or tray furnaces. The calcination can be effected at atmospheric pressure or reduced pressure and/or in a gas stream, for example in an air stream or hydrogen stream. A pretreatment with hydrogen or gases comprising hydrogen, in general under conditions which correspond to the hydrogenation conditions, serves for preliminary reduction/activation of the hydrogenation catalyst. However, the catalyst can also be reduced in situ under the conditions specified in the case of the hydrogenation, preferably under pressure (for example at a hydrogen pressure of from about 100 to 325 bar).

In the hydrogenation, the glycerol and the resulting 1,2propanediol are preferably present in the liquid phase.

The catalysts may be arranged, for example, in a fixed bed or may be used as a suspension. The hydrogenation can accordingly be carried out, for example, by the trickle-bed procedure or the liquid-phase procedure. For the liquid-phase hydrogenation, the catalysts are preferably used in finely divided form, for example as powder, in suspension. In the hydrogenation in the trickle phase, the catalysts are used as moldings, as described above, for example in the form of pressed cylinders, tablets, lozenges, wagon wheels, rings, stars or extrudates, such as solid extrudates, polylobal extrudates, hollow extrudates and honeycomb bodies. Excess hydrogen is preferably circulated, it being possible for a small part to be discharged as waste gas for removing inert materials. It is possible to use one reactor or a plurality of reactors which can be connected in series or parallel to one another.

The temperature in the hydrogenation in step b) is preferably from 150 to 300° C., in particular from 175 to 250° C.

The reaction pressure in step b) is preferably from 140 bar to 250 bar.

The molar ratio of hydrogen to glycerol is preferably from

The catalyst space velocity in the continuous procedure is preferably from 0.1 to 1, more preferably from 0.2 to 0.6, and in particular from 0.3 to 0.6, kg of glycerol to be hydrogenated per kg (catalyst) per h.

The conversion, based on glycerol, is preferably at least 90%, in particular at least 95%. The selectivity, based on 1,2-propanediol is preferably at least 85%, particularly preferably at least 90%, in the process according to the invention. Often, even higher selectivities of up to 95% or more can be achieved.

The hydrogenation is expediently carried out continuously. The hydrogenation discharge substantially comprises 1,2propanediol. Further constituents are, inter alia, methanol, ethanol, n-propanol, isopropanol, 1,3-propanediol, glycerol, ethylene glycol and water. The hydrogenation discharge can then be worked up by conventional methods known to the person skilled in the art. For example, thermal working-up, preferably distillation, adsorption, ion exchange, a membrane separation method, crystallization or extraction or a combination of two or more of these methods can be used. Preferred is a working-up by distillation. This can in principle be effected by conventional distillation methods known to the person skilled in the art. Suitable apparatuses for the distillative working-up comprise distillation columns, such as tray columns, which may be equipped with caps, sieve plates, sieve trays, stacked packings, dumped packings, valves, side take-offs, etc. evaporators, such as thin-film evaporators, fall-

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ing-film evaporators, forced-circulation evaporators, Sambay evaporators, etc., and combinations thereof. Glycerol still present in the hydrogenation discharge can be recycled to the hydrogenation stage, if appropriate after being separated off by distillation.

The invention is explained in more detail with reference to the following, non-limiting examples.

EXAMPLES

Glycerol of the pharmaceutical glycerol and pure glycerol qualities from Biodiesel Schwarzheide GmbH was used as feedstock for the experiments for catalyst screening. Table 1 shows the analytical data of the glycerol used.

| ГA | BL | Æ | 1 |
|----|----|---|---|
| | | | |

| Glycerol quality | Water [%] | pH [pH] | chloride [ppm] | Glycerol content [%] | 20 |
|------------------|--------------|------------|-------------------|----------------------------|----|
| Pure | 0.1 | 7 | 2 | 97-98 | 25 |
| Pharmaceutical | 0.141 | 7 | 1.4 | 99.8 | |

The analysis of the feedstock glycerol and of the reaction discharge is effected by gas chromatography (data in GC $\%^{-30}$ by area).

The catalysts were activated for 10 h at a temperature of 200° C. and a hydrogen pressure of 50 bar before the reaction.

General Method for Carrying Out Catalyst Tests Batchwise. Pharmaceutical glycerol having a water content of 20% was used. The catalyst was initially taken in a 0.3 1 mini autoclave and the autoclave was closed and was tested for leaks with 200 bar N₂ at room temperature. Catalyst extrudates were used in the form of moldings, and the extrudates were comminuted beforehand for the preparation of catalyst suspensions.

Thereafter, the autoclave was depressurized and the activation of the catalyst carried out. For this purpose, 50 bar H₂ was forced in at room temperature, heating was then effected to an internal temperature of 200° C. and the temperature was maintained for about 10 h without stirring. After cooling to 30° C. and subsequent inertization with N₂ the autoclave was evacuated and the reaction solution was aspirated.

For the reaction of the glycerol, 50 bar H₂ was forced in at room temperature and the reaction mixture was heated to 215° C. with stirring (speed from 700 to 1000 rpm). The pressure resulting in the autoclave was supplemented with H₂ to the desired final pressure of 200 bar. Hydrogen consumed in the reaction was replenished. The run time of the experiments was 10 hours. After the end of the run time of the experiment, the autoclave was cooled to room temperature and depressurized. The analysis of the samples and discharges was effected by means of gas chromatography by integration of the areas of the peaks (% by area). The results are shown in table 3.

TABLE 3

| Apparatus: | HP 5890-2 with sampler | | Comparison of the catalyst in the fixed bed and suspension procedures | | | | | | |
|--------------------------------------------------------------------|--------------------------------------------------------------------------|-------------|--------------------------------------------------------------------------|--------|-----------------------------------|------------------------|------------------------|--------------------|----------|
| Range: Column: Sample volume: Carrier gas: Fluid rate: | 2 30 m DBWax; film thickness: 0.25 μm 1 μl Helium 100 ml/min | | Ex. No. | Cataly | Catalyst st form | Amount of catalyst [g] | Conver- sion [%] | Selectivity [%] | Yi [% |
| Injector temperature: Detector: Detector temperature: | 240° C. FID (Flame ionization detector) 250° C. | 40 | 1 2 | A B | Suspension Extrudates 5 × 3 | 20.0 10.0 | 97.6 91.5 | 95.3 91.2 | 93 83 |
| Temperature program: | 5 min at 40° C., 10° C./min to 240° C., 15 min at 240° C. | | 3 | С | Extrudates 5×3 | 18.3 | 100.0 | 87.8 | 87 |
| | Total run time 45 min | — 45 | 4 5 | D E | Suspension Extrudates 5 × 5 | 20.0 10.0 | 97.2 99.1 | 93.6 88.8 | 91 88 |

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Copper-containing catalysts of different compositions were tested (cf. table 2).

TABLE 2

| | Overview of the catalysts tested |
|----------|------------------------------------------------------------------------------------------------|
| Catalyst | Composition |
| А | 67% CuO; 5% La ₂ O ₃ ; Al ₂ O ₃ (ad 100%) + 15% Cu |
| В | 40% CuO; 40% ZnO; 20% Al ₂ O ₃ |
| C* | 40% CuO; 40% ZnO; 20% Al ₂ O ₃ |
| D | 61% CuO; 39% Al ₂ O ₃ |
| Е | 70% CuO; 24.5% ZnO; 5.5% Al ₂ O ₃ |
| F | 55% CuO/Al ₂ O ₃ |
| G | 16% CuO/64% Al ₂ O ₃ /20% ZnO |
| Н | 100% Cu (Raney-Cu**) |
| Ι | 40% Cu/TiO ₂ |
| J | 70% CuO/20% ZnO/10% ZrO ₂ |
| K | 60% CuO/30% Al ₂ O ₃ /10% MnO ₃ |
| L | 66-77% CuO/21-32% Cu-Chromit/2% Graphit |

*as for B, but higher calcination temperature (from 400 to 500° C.) **from Aldrich

| | bed and suspension procedures | | | | | | | | |
|---|-------------------------------|----------|---------------------|------------------------|------------------------|--------------------|--------------|--|--|
| | Ex. No. | Catalyst | Catalyst form | Amount of catalyst [g] | Conver- sion [%] | Selectivity [%] | Yield [%] | | |
| 0 | 1 | А | Suspension | 20.0 | 97.6 | 95.3 | 93.0 | | |
| | 2 | В | Extrudates 5 × 3 | 10.0 | 91.5 | 91.2 | 83.5 | | |
| | 3 | С | Extrudates 5 × 3 | 18.3 | 100.0 | 87.8 | 87.8 | | |
| | 4 | D | Suspension | 20.0 | 97.2 | 93.6 | 91.0 | | |
| 5 | 5 | Е | Extrudates 5 × 5 | 10.0 | 99.1 | 88.8 | 88.0 | | |
| | 6 | F | Suspension | 20.0 | 96.8 | 94.4 | 91.3 | | |
| | 7 | G | Suspension | 10.0 | 92.3 | 92.5 | 85.4 | | |
| | 8 | Η | Suspension | 10.0 | 99.6 | 96.1 | 95.8 | | |
| | 9 | J | tablets | 18.0 | 92.7 | 96.6 | 89.6 | | |
| 0 | 10 | K | tablets | 18.0 | 79.6 | 94.5 | 75.2 | | |
| v | 11 | L | tablets | 18.0 | 74.5 | 96.8 | 72.1 | | |

General Method for the Continuous Hydrogenation Using Fixed-Bed Catalysts

Pharmaceutical glycerol having a water content of 10% was used. The experiments were carried out in a continuously operated laboratory apparatus at from 200 to 240 bar. The experimental series 9 to 11 were operated for simulating the main reactor with liquid circulation in the liquid-phase procedure. In each case 70 ml of the catalysts were used.

The structure of the unit and the process description are described below:

The unit consists of a 75 ml tubular reactor R1 (internal 65 Ø=12 mm, L=800 mm) having three liquid-heated heating zones, which is operated by the liquid-phase procedure. If required a liquid circulation which is operated with flow control (Danfoss) via an HPLC pump can be connected. All parts of the unit are made from metal and designed for an operating pressure of up to 250 bar.

The glycerol solution (aqueous, 90% strength) is metered continuously, regulated by a balance, into the reactor R1 and is reacted under defined conditions (pressure, temperature, catalyst space velocity) with hydrogen to give the desired product. The hydrogen is supplied from 50 l steel cylinders which are compressed to the required pressure by means of a compressed air-operated compressor. The desired reaction pressure is established via pressure control (P2) in the waste gas stream, and the required amount of hydrogen is fed into the reactor R1 with flow control via a mass flow meter (Hi-Tec). The liquid reactor discharge is discharged with level control (container B2) via an HPLC pump and collected in the discharge container (B5). The gaseous reactor discharge is passed via a buffer vessel (B4) and depressurized by means of a pressure-controlled (P2) Recco valve.

In experimental series 15, the reactions were continued in 20 a modified unit (main reactor with liquid circulation in the trickle-bed procedure, downstream reactor without liquid circulation in the liquid-phase procedure). In all experiments, the catalyst was stable, no catalyst loss occurred as a result of so-called "leaching".

7. The process of claim 1, wherein said glycerol-containing stream is subjected to catalytic desulfurization prior to step b).

8. The process of claim **7**, wherein said catalytic desulfurization is performed in the presence of hydrogen.

9. The process of claim **7**, wherein said sulfur-containing compounds comprise sulfur-containing aromatic compounds.

10. The process of claim 1, wherein said glycerol-containing stream is contacted with at least one adsorbent for removing components which adversely affect catalytic hydrogenation.

11. The process according to claim **1**, wherein said adsorbent comprises at least one component capable of use as a hydrogenation catalyst.

12. The process according to claim **1**, wherein said glycerol-containing stream is provided by:

- a1) providing a biogenic fat-and/or oil-containing starting mixture comprising fatty acid triglycerides and free fatty acids;
- a2) transesterifying said fatty acid triglycerides with at least one C₁-C₉-monoalcohol and, optionally, esterifying said free fatty acids, to form an esterification mixture;

TABLE 4

| Results of the continuous experiments (best setting from each experimental series) | | | | | | | | | | | |
|------------------------------------------------------------------------------------|------------------|--------------------------|------------------------------|--------------------------|------------------------------|-------------------------------|-------------------------|----------------------|-----------------------|------------------------|------------------------|
| Expt. No. | Catalyst | Run time [h] | Temp. [° C.] | Pressure [bar] | Feed [g/h] | Cat. space vel. [kg/l · h] | LR g/h | pH Discharge | Glycerol conversion | Yield | Selectivity |
| 12 13 14 15 | I I A B | 230 153 287 253 | 217 220 200 170-190 | 200 240 200 200 | 31.1 31.1 23.3 44.4 | 0.4 0.4 0.3 0.4 | 155 0 300 1800 | 5.5 6 6 6-7 | 81 96 88 100 | 81 94 82 98.5 | 99 98 92 98.5 |

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LR = liquid recycling (circulation) from the reactor exit to the reactor entrance

We claim:

- 1. A process for preparing 1,2-propanediol, comprising
- a) providing a glycerol-containing stream; and
- b) hydrogenating said glycerol-containing stream in the presence of a heterogeneous catalyst which comprises copper at a temperature in the range of from 100° C. to 45 320° C. and at a pressure in the range of from 100 bar to 325 bar;
- wherein said glycerol-containing stream has a water content of 30% by weight or less.

2. The process of claim 1, wherein said glycerol-containing stream is obtained from the preparation of alkyl esters of higher fatty acids by transesterification of fatty acid triglycerides.

3. The process of claim **1**, wherein said glycerol-containing stream has a water content of 20% by weight or less. 55

4. The process of claim 1, wherein said glycerol-containing stream is substantially anhydrous.

5. The process of claim **1**, wherein said glycerol-containing stream is worked-up via at least one work-up process selected from the group consisting of thermal working-up, adsorption, ion exchange, membrane separation, crystallization, extraction, and combinations thereof.

6. The process of claim **1**, wherein the water content of said glycerol-containing stream is reduced and/or components 65 which adversely affect catalytic hydrogenation are removed from said glycerol-containing stream via distillation.

- a3) separating said esterification mixture so as to obtain at least one fraction enriched with C_1 - C_9 -monoalkyl esters and at least one fraction enriched with glycerol liberated in the transesterification of a2),
- a4) optionally, purifying said at least one fraction enriched with glycerol.

13. The process according to claim **1**, wherein said heterogenous catalyst comprises a Raney catalyst.

14. The process of claim 1, wherein said heterogenous catalyst comprises at least 23% by weight of copper, in oxidic and/or elemental form, based on the total weight of the catalyst.

15. The process of claim **1**, wherein said heterogenous catalyst comprises at least 35% by weight of copper, in oxidic and/or elemental form, based on the total weight of the catalyst.

16. The process of claim 1, wherein said heterogenous catalyst comprises metals or mixtures of metals selected from the group consisting of Cu; Cu and Ti; Cu and Zr; Cu and Mn; Cu and Al; Cu, Ni, and Mn; Cu, Al, and at least one further metal selected from the group consisting of La, W, Mo, Mn, Zn, Ti, Zr, Sn, Ni, and Co; Cu, Zn, and Zr; Cu, Cr, and Ca; Cu, Cr, and C; and Cu, Al, Mn, and optionally Zr; wherein each metal is present in oxidic form, elemental form, or a combination thereof.

* * * * *

ASPEN INPUT SUMMARY

;Input Summary created by Aspen Plus Rel. 24.0 at 18:36:42 Mon Apr 11, 2011 ;Directory C:\temp Filename c:\temp\~ap2e09.txt

DYNAMICS DYNAMICS RESULTS=ON

IN-UNITS ENG

DEF-STREAMS CONVEN ALL

SIM-OPTIONS OLD-DATABANK=YES

DATABANKS PURE22 / AQUEOUS / SOLIDS / INORGANIC / & NOASPENPCD

PROP-SOURCES PURE22 / AQUEOUS / SOLIDS / INORGANIC

COMPONENTS GLYCEROL C3H8O3 / ACETOL C3H6O2-D1 / WATER H2O / HYDROGEN H2 / PG C3H8O2-2 / N-PROP C3H8O-1 / ETHYL-01 C2H6O2 / ISO-PROP C3H8O-2 / SODIU-01 NACL /

METHA-01 CH4O

HENRY-COMPS HC-1 HYDROGEN

FLOWSHEET

BLOCK VALVE IN=S-132 OUT=S-133 BLOCK D-102 IN=S-133 OUT=S-137 S-136 S-134 BLOCK V-101 IN=S-128 OUT=S-130 S-129 BLOCK P-100 IN=S-114 OUT=S-115 BLOCK R-101 IN=S-116 S-120 OUT=S-121 BLOCK CP-101 IN=S-130 OUT=S-131 BLOCK M-101 IN=S-131 S-118 OUT=S-119 BLOCK T-101 IN=S-117 OUT=S-118 BLOCK HX-103 IN=S-126 S-119 OUT=S-127 S-120 BLOCK C-102 IN=S-127 OUT=S-128 BLOCK C-103 IN=S-134 OUT=S-135 BLOCK M-102 IN=S-101 S-102 S-112 OUT=S-103 BLOCK HX-101 IN=S-109 S-106 OUT=S-110 S-107 BLOCK D-101 IN=S-108 OUT=S-109 S-114 BLOCK H-101 IN=S-107 OUT=S-108 BLOCK FSPLIT IN=S-110 OUT=S-113 S-111 BLOCK F-101 IN=S-121 OUT=S-123 S-122 BLOCK HX-102 IN=S-123 S-115 OUT=S-124 S-116

BLOCK F-102 IN=S-124 OUT=S-126 S-125 BLOCK M-103 IN=S-122 S-129 S-125 OUT=S-132 BLOCK S-101 IN=S-103 OUT=S-104 S-105 BLOCK P-101 IN=S-105 OUT=S-106 BLOCK P-102 IN=S-111 OUT=S-112 PROPERTIES PENG-ROB TRUE-COMPS=YES **PROPERTIES NRTL-RK / UNIQ-RK PROP-DATA HENRY-1 IN-UNITS ENG PROP-LIST HENRY** BPVAL HYDROGEN WATER 198.2062671 - 12588.31790 - 26.31190000 & 8.35727785E-3 33.53000373 150.5300028 0.0 BPVAL HYDROGEN N-PROP -71.55669085 3641.940059 13.38100000 & -.0153788890 -75.99999539 77.00000338 0.0 **PROP-DATA NRTL-1 IN-UNITS ENG PROP-LIST NRTL** BPVAL GLYCEROL WATER -.7026000000 283.4269177 .3000000000 & 0.0 0.0 0.0 77.00000338 553.9999996 BPVAL WATER GLYCEROL -1.093700000 407.9753967 .3000000000 & 0.0 0.0 0.0 77.00000338 553.9999996 BPVAL GLYCEROL PG -.6973000000 -65.90807947 .3000000000 0.0 & 0.0 0.0 191.4800025 357.6200011 BPVAL PG GLYCEROL .0663000000 454.1135364 .3000000000 0.0 & 0.0 0.0 191.4800025 357.6200011 BPVAL WATER PG 0.0 -543.0875357 .300000000 0.0 0.0 0.0 & 59.00000353 212.0000023 BPVAL PG WATER 0.0 849.8359732 .3000000000 0.0 0.0 0.0 & 59.00000353 212.0000023 BPVAL WATER N-PROP 5.508500000 -1583.168747 .3000000000 0.0 & 0.0 0.0 77.00000338 212.0000023 BPVAL N-PROP WATER -1.809800000 1074.524751 .3000000000 0.0 & 0.0 0.0 77.00000338 212.0000023 BPVAL PG N-PROP 2.467600000 - 2322.928601 .3000000000 0.0 & 0.0 0.0 213.4400023 347.7200012 BPVAL N-PROP PG -5.774100000 5454.272836 .3000000000 0.0 & 0.0 0.0 213.4400023 347.7200012 BPVAL GLYCEROL ETHYL-01 0.0 -621.3814150 .300000000 0.0 & 0.0 0.0 205.7000024 307.0400015 BPVAL ETHYL-01 GLYCEROL 0.0 532.5047957 .3000000000 0.0 & 0.0 0.0 205.7000024 307.0400015 BPVAL WATER ETHYL-01 .3184000000 59.89895952 .3000000000 & 0.0 0.0 0.0 86.72000331 386.0600009 BPVAL ETHYL-01 WATER .0531000000 -314.0342975 .3000000000 & 0.0 0.0 0.0 86.72000331 386.0600009 BPVAL PG ETHYL-01 -1.262600000 1782.934906 .3000000000 0.0 & 0.0 0.0 182.4800025 314.6900015 BPVAL ETHYL-01 PG -.4391000000 -242.0443781 .3000000000 0.0 & 0.0 0.0 182.4800025 314.6900015

PROP-DATA PRKBV-1

IN-UNITS ENG

- PROP-LIST PRKBV
- BPVAL WATER METHA-01 -.0778000000 0.0 0.0 -459.6699923 &
- 1340.329993
- BPVAL METHA-01 WATER -.0778000000 0.0 0.0 -459.6699923 & 1340.329993

PROP-DATA UNIQ-1

IN-UNITS ENG

PROP-LIST UNIQ

- BPVAL GLYCEROL WATER .3575000000 -127.4470190 0.0 0.0 & 77.00000338 553.9999996 0.0
- BPVAL WATER GLYCEROL .8232000000 -261.0345579 0.0 0.0 & 77.00000338 553.9999996 0.0
- BPVAL GLYCEROL PG .4210000000 -8.234459934 0.0 0.0 & 191.4800025 357.6200011 0.0
- BPVAL PG GLYCEROL -.2466000000 -112.8707991 0.0 0.0 & 191.4800025 357.6200011 0.0
- BPVAL GLYCEROL ETHYL-01 0.0 327.0122974 0.0 0.0 & 205.7000024 307.0400015 0.0
- BPVAL ETHYL-01 GLYCEROL 0.0 -319.0148974 0.0 0.0 & 205.7000024 307.0400015 0.0
- BPVAL WATER PG 0.0 -492.1070361 0.0 0.0 59.00000353 & 212.0000023 0.0
- BPVAL PG WATER 0.0 451.2850164 0.0 0.0 59.00000353 & 212.0000023 0.0
- BPVAL WATER N-PROP -2.388600000 1109.333151 0.0 0.0 & 77.00000338 212.0000023 0.0
- BPVAL N-PROP WATER 1.826100000 -1202.000210 0.0 0.0 & 77.00000338 212.0000023 0.0
- BPVAL WATER ETHYL-01 -.4787000000 140.8377589 0.0 0.0 & 86.72000331 386.0600009 0.0
- BPVAL ETHYL-01 WATER .4661000000 49.24979961 0.0 0.0 & 86.72000331 386.0600009 0.0
- BPVAL PG N-PROP -.7915000000 943.5014925 0.0 0.0 & 213.4400023 347.7200012 0.0
- BPVAL N-PROP PG 1.948700000 -2146.453363 0.0 0.0 & 213.4400023 347.7200012 0.0
- BPVAL PG ETHYL-01 .4147000000 -859.2713931 0.0 0.0 & 182.4800025 314.6900015 0.0
- BPVAL ETHYL-01 PG .2677000000 173.6895586 0.0 0.0 & 182.4800025 314.6900015 0.0

STREAM S-101

SUBSTREAM MIXED TEMP=70. PRES=30. MASS-FLOW=19392.4873 MASS-FLOW GLYCEROL 15513.99 / WATER 2908.873 / SODIU-01 & 775.6994 / METHA-01 193.9249

STREAM S-102

SUBSTREAM MIXED TEMP=70. PRES=30. MOLE-FLOW=1. MASS-FRAC WATER 1.

STREAM S-117

SUBSTREAM MIXED TEMP=70. PRES=1000. MOLE-FLOW=1.

MOLE-FRAC HYDROGEN 1.

BLOCK M-101 MIXER

- BLOCK M-102 MIXER PARAM PRES=25.
- BLOCK M-103 MIXER PARAM PRES=-5.
- BLOCK FSPLIT FSPLIT FRAC S-113 0.34
- BLOCK S-101 SEP PARAM PRES=-5. FRAC STREAM=S-104 SUBSTREAM=MIXED COMPS=SODIU-01 FRACS=1. FLASH-SPECS S-104 NPHASE=1 FREE-WATER=NO PHASE=S FLASH-SPECS S-105 PRES=0. DELT=15.
- BLOCK C-102 HEATER PARAM TEMP=225. PRES=-5.
- BLOCK C-103 HEATER PARAM TEMP=90. PRES=-5.
- BLOCK H-101 HEATER PARAM TEMP=242. PRES=-5.
- BLOCK F-101 FLASH2 PARAM PRES=560. DUTY=0.
- BLOCK F-102 FLASH2 PARAM PRES=0. DUTY=0.
- BLOCK V-101 FLASH2 PARAM TEMP=225. PRES=0.

BLOCK HX-101 HEATX PARAM T-HOT=80. CALC-METHOD=TASCPLUS-RIG HETRAN-PARAM INPUT-FILE= & 'EDR_HX101.EDR' FEEDS HOT=S-109 COLD=S-106 PRODUCTS HOT=S-110 COLD=S-107 PROPERTIES PENG-ROB FREE-WATER=STEAM-TA SOLU-WATER=3 & TRUE-COMPS=YES / PENG-ROB FREE-WATER=STEAM-TA & SOLU-WATER=3 TRUE-COMPS=YES HOT-SIDE DP-OPTION=CONSTANT COLD-SIDE DP-OPTION=CONSTANT

BLOCK HX-102 HEATX PARAM T-COLD=410. CALC-METHOD=TASCPLUS-RIG HETRAN-PARAM INPUT-FILE= & 'EDR_HX102.EDR' FEEDS HOT=S-123 COLD=S-115
PRODUCTS HOT=S-124 COLD=S-116 PROPERTIES PENG-ROB FREE-WATER=STEAM-TA SOLU-WATER=3 & TRUE-COMPS=YES / PENG-ROB FREE-WATER=STEAM-TA & SOLU-WATER=3 TRUE-COMPS=YES HOT-SIDE DP-OPTION=CONSTANT COLD-SIDE DP-OPTION=CONSTANT

BLOCK HX-103 HEATX PARAM DECR-HOT=20. CALC-METHOD=TASCPLUS-RIG HETRAN-PARAM INPUT-FILE= & 'EDR_HX103.EDR' FEEDS HOT=S-126 COLD=S-119 PRODUCTS HOT=S-127 COLD=S-120 PROPERTIES PENG-ROB FREE-WATER=STEAM-TA SOLU-WATER=3 & TRUE-COMPS=YES / PENG-ROB FREE-WATER=STEAM-TA & SOLU-WATER=3 TRUE-COMPS=YES HOT-SIDE DP-OPTION=CONSTANT COLD-SIDE DP-OPTION=CONSTANT

BLOCK D-101 RADFRAC PARAM NSTAGE=6 COL-CONFIG CONDENSER=TOTAL FEEDS S-108 4 PRODUCTS S-109 1 L / S-114 6 L P-SPEC 1 20. COL-SPECS MOLE-B=375.5 MOLE-RR=0.2

BLOCK D-102 RADFRAC PARAM NSTAGE=16 COL-CONFIG CONDENSER=PARTIAL-V-L FEEDS S-133 10 PRODUCTS S-136 1 L / S-134 16 L / S-137 1 V P-SPEC 1 20. COL-SPECS MOLE-B=164.3 MOLE-RR=0.13 T1=130.

BLOCK R-101 RSTOIC
PARAM PRES=-15. DUTY=0. SERIES=YES
STOIC 1 MIXED GLYCEROL -1. / HYDROGEN -1. / PG 1. / & WATER 1.
STOIC 2 MIXED GLYCEROL -2. / HYDROGEN -4. / ISO-PROP 1. / & N-PROP 1. / WATER 4.
CONV 1 MIXED GLYCEROL 0.985
CONV 2 MIXED GLYCEROL 1.

BLOCK P-100 PUMP PARAM PRES=587.

BLOCK P-101 PUMP PARAM PRES=42.

BLOCK P-102 PUMP PARAM PRES=30.

BLOCK CP-101 COMPR

PARAM TYPE=ISENTROPIC PRES=600. SEFF=0.8

BLOCK T-101 COMPR PARAM TYPE=ISENTROPIC PRES=600. SEFF=0.8 MODEL-TYPE=TURBINE

BLOCK VALVE VALVE PARAM P-OUT=30.

DESIGN-SPEC DS-1 DEFINE MIXH2 MOLE-FLOW STREAM=S-119 SUBSTREAM=MIXED & COMPONENT=HYDROGEN SPEC "MIXH2" TO "665.5535" TOL-SPEC ".1" VARY STREAM-VAR STREAM=S-117 SUBSTREAM=MIXED & VARIABLE=MOLE-FLOW LIMITS "0" "400"

DESIGN-SPEC DS-2 DEFINE WATFRAC MASS-FRAC STREAM=S-103 SUBSTREAM=MIXED & COMPONENT=WATER SPEC "WATFRAC" TO ".5" TOL-SPEC ".01" VARY STREAM-VAR STREAM=S-102 SUBSTREAM=MIXED & VARIABLE=MOLE-FLOW LIMITS "0" "800"

EO-CONV-OPTI

SENSITIVITY S-1 DEFINE METHREC MOLE-FLOW STREAM=S-103 SUBSTREAM=MIXED & COMPONENT=METHA-01 TABULATE 1 "METHREC" VARY BLOCK-VAR BLOCK=FSPLIT SENTENCE=FRAC VARIABLE=FRAC & ID1=S-113 RANGE LOWER="0" UPPER=".5" NPOINT="20"

CONV-OPTIONS WEGSTEIN MAXIT=70

STREAM-REPOR MOLEFLOW MOLEFRAC MASSFRAC

DISABLE

SENSITIVITY S-1

;

ASPEN SIMULATION REPORT

ASPEN PLUS PLAT: WIN32 VER: 24.0

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RUN CONTROL SECTION

RUN CONTROL INFORMATION

THIS COPY OF ASPEN PLUS LICENSED TO UNIVERSITY OF PENNSYLVAN

TYPE OF RUN: NEW

INPUT FILE NAME: _1152wfu.inm

OUTPUT PROBLEM DATA FILE NAME: _1152wfu LOCATED IN:

PDF SIZE USED FOR INPUT TRANSLATION: NUMBER OF FILE RECORDS (PSIZE) = 0 NUMBER OF IN-CORE RECORDS = 256 PSIZE NEEDED FOR SIMULATION = 256

CALLING PROGRAM NAME: apmain LOCATED IN: C:\PROGRA~1\ASPENT~1\ASPENP~2.2\Engine\xeq

SIMULATION REQUESTED FOR ENTIRE FLOWSHEET ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 2

FLOWSHEET SECTION

FLOWSHEET CONNECTIVITY BY STREAMS

| STREAM | A SOU | RCE I | DEST | STREAM | SOURCE | DEST |
|--------|--------|--------|--------|----------|----------|------|
| S-117 | ' | Т-101 | S-102 | M | -102 | |
| S-101 |] | M-102 | S-133 | VALVE | D-102 | |
| S-137 | D-102 | | S-136 | D-102 | | |
| S-134 | D-102 | C-103 | S-130 | V-101 | CP-101 | |
| S-129 | V-101 | M-103 | S-11: | 5 P-100 | HX-102 | |
| S-121 | R-101 | F-101 | S-131 | CP-101 | M-101 | |
| S-119 | M-101 | HX-10 |)3 S-1 | 18 T-101 | M-101 | |
| S-127 | HX-103 | C-102 | 2 S-12 | 0 HX-10 | 03 R-101 | |
| S-128 | C-102 | V-101 | S-135 | C-103 | | |
| S-103 | M-102 | S-101 | S-110 |) HX-101 | FSPLIT | |
| S-107 | HX-101 | H-101 | S-10 | 09 D-101 | HX-101 | |
| S-114 | D-101 | P-100 | S-108 | H-101 | D-101 | |
| S-113 | FSPLIT | | S-111 | FSPLIT | P-102 | |
| S-123 | F-101 | HX-102 | 2 S-12 | 2 F-101 | M-103 | |
| S-124 | HX-102 | F-102 | S-11 | 6 HX-10 | 2 R-101 | |
| S-126 | F-102 | HX-103 | 3 S-12 | 5 F-102 | M-103 | |
| S-132 | M-103 | VALV | E S-1 | 04 S-10 | l | |
| S-105 | S-101 | P-101 | S-106 | P-101 | HX-101 | |

FLOWSHEET CONNECTIVITY BY BLOCKS

| BLOCK | INLETS | OUTLETS | |
|----------|-------------------|-------------------|----------|
| VALVE | S-132 | S-133 | |
| D-102 | S-133 | S-137 S-136 S-134 | |
| V-101 | S-128 | S-130 S-129 | |
| P-100 | S-114 | S-115 | |
| R-101 | S-116 S-120 | S-121 | |
| CP-101 | S-130 | S-131 | |
| M-101 | S-131 S-118 | S-119 | |
| T-101 | S-117 | S-118 | |
| HX-103 | S-126 S-119 | S-127 S-120 | |
| C-102 | S-127 | S-128 | |
| C-103 | S-134 | S-135 | |
| M-102 | S-101 S-102 S-112 | S-103 | |
| HX-101 | S-109 S-106 | S-110 S-107 | |
| D-101 | S-108 | S-109 S-114 | |
| H-101 | S-107 | S-108 | |
| FSPLIT | S-110 | S-113 S-111 | |
| F-101 | S-121 | S-123 S-122 | |
| HX-102 | S-123 S-115 | S-124 S-116 | |
| F-102 | S-124 | S-126 S-125 | |
| M-103 | S-122 S-129 S-125 | S-132 | |
| S-101 | S-103 | S-104 S-105 | |
| P-101 | S-105 | S-106 | |
| P-102 | S-111 | S-112 | |
| ASPEN PL | US PLAT: WIN32 | VER: 24.0 | 04/11/20 |
| | | | |

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FLOWSHEET SECTION

CONVERGENCE STATUS SUMMARY

DESIGN-SPEC SUMMARY

DESIGN CONV SPEC ERROR TOLERANCE ERR/TOL VARIABLE STAT BLOCK ----- ----- -----DS-1 0.34537E-01 0.10000 0.34537 171.84 # \$OLVER03 DS-2 0.43373E-02 0.10000E-01 0.43373 298.88 # \$OLVER04

TEAR STREAM SUMMARY

STREAM MAXIMUM MAXIMUM VARIABLE CONV ID ERROR TOLERANCE ERR/TOL ID STAT BLOCK ----- ----- -----

S-108 0.18110E-08 0.22262E-06 0.81349E-02 METHA-01MOLEFLOW # \$OLVER01

| S-106 0.10204E-04 0.11994E-04 0.85077 WATER MOLEFLOW # \$OLV S-121 0.12667E-06 0.62868E-07 2.0148 ISO-PROPMOLEFLOW * \$OLV S-119 0.11910E-06 0.46937E-07 2.5376 ISO-PROPMOLEFLOW * \$OLV | /ER01 /ER02 /ER02 |
|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------|
| # = CONVERGED * = NOT CONVERGED LB = AT LOWER BOUNDS UB = AT UPPER BOUNDS | |
| DESIGN-SPEC: DS-1 | |
| SAMPLED VARIABLES: MIXH2 : HYDROGENMOLEFLOW IN STREAM S-119 SUBSTREAM MIX | ED |
| SPECIFICATION: MAKE MIXH2 APPROACH 665.554 WITHIN 0.100000 | |
| MANIPULATED VARIABLES: VARY : TOTAL MOLEFLOW IN STREAM S-117 SUBSTREAM MIXED LOWER LIMIT = 0.0 LBMOL/HR UPPER LIMIT = 400.000 LBMOL/HR FINAL VALUE = 171.838 LBMOL/HR | |
| VALUES OF ACCESSED FORTRAN VARIABLES: VARIABLE VALUE AT START FINAL VALUE UNITS OF LOOP | |
| MIXH2 665.588 665.588 LBMOL/HR | |
| DESIGN-SPEC: DS-2 | |
| SAMPLED VARIABLES: WATFRAC : WATER MASSFRAC IN STREAM S-103 SUBSTREAM MIXEI ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 4 |) |
| FLOWSHEET SECTION | |
| DESIGN-SPEC: DS-2 (CONTINUED) | |
| SPECIFICATION: MAKE WATFRAC APPROACH 0.50000 WITHIN 0.0100000 | |
| MANIPULATED VARIABLES: VARY : TOTAL MOLEFLOW IN STREAM S-102 SUBSTREAM MIXED LOWER LIMIT = 0.0 LBMOL/HR UPPER LIMIT = 800.000 LBMOL/HR FINAL VALUE = 298.877 LBMOL/HR | |
| VALUES OF ACCESSED FORTRAN VARIABLES: VARIABLE VALUE AT START FINAL VALUE UNITS | |

| OF L | OOP | | | |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------|--------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| WATFRAC | 0.504337 | 0.5043 | 337 | |
| CONVERGENCE | BLOCK: \$ | OLVER01 | | |
| Tear Stream : S Tolerance used: Trace molefrac: | S-108 S-1 0.100D-03 0.100D-05 | 06 0.100D-03 0.100D-05 | 5 | |
| MAXIT= 70 W QMAX = 0.0 METHOD: WE TOTAL NUMB | /AIT 1 ITE QMIN = - GSTEIN S ER OF ITEF | RATIONS 5.0 STATUS: (RATIONS: | BEFORE ACC CONVERGED 59 | CELERATING |
| *** | FINAL VA | LUES *** | | |
| VAR# TEAR STR VALUE PREV | EAM VAR VALUE | STREAM ERR/TOL | SUBSTREA (| COMPONEN ATTRIBUT ELEMENT UNIT |
| 1 TOTAL MOLE 2 TOTAL MOLE 3 MOLE-FLOW | EFLOW S- EFLOW S- S-108 | 108 MIX 106 MIX MIXED | ED ED GLYCEROL | LBMOL/HR 1138.0639 1138.0639 4.8677-05 LBMOL/HR 1138.1451 1138.0639 0.7134 LBMOL/HR 168.4574 168.4574 3.6901- |
| 4 MOLE-FLOW 5 MOLE-FLOW | S-108 S-108 | MIXED MIXED | ACETOL WATER | LBMOL/HR 0.0 0.0 0.0 LBMOL/HR 951.9378 951.9378 -9.2861- |
| 6 MOLE-FLOW 7 MOLE-FLOW 8 MOLE-FLOW 9 MOLE-FLOW 10 MOLE-FLOW 11 MOLE-FLOW 12 MOLE-FLOW | S-108 S-108 S-108 S-108 S-108 S-108 S-108 S-108 S-108 | MIXED MIXED MIXED MIXED MIXED MIXED MIXED | HYDROGEN PG N-PROP ETHYL-01 ISO-PROP SODIU-01 METHA-01 | LBMOL/HR 0.0 0.0 0.0 LBMOL/HR 17.6687 17.6687 8.1349- |
| 13 PRESSURE 14 MASS ENTH 15 MOLE-FLOW | S-108 1 ALPY S-10 / S-106 | MIXED)8 MIXE MIXED | D GLYCEROL | PSIA 36.8160 36.8160 -9.8423-05 BTU/LB -4866.9331 -4866.9332 6.7809-05 LBMOL/HR 168.4574 168.4574 |
| 4.3773-06 16 MOLE-FLOW 17 MOLE-FLOW 18 MOLE-FLOW 19 MOLE-FLOW 20 MOLE-FLOW 21 MOLE-FLOW ASPEN PLUS PI | X X S-106 X X | MIXED MIXED MIXED MIXED MIXED MIXED VER: 2 | ACETOL WATER HYDROGEN PG N-PROP ETHYL-01 4.0 | LBMOL/HR 0.0 0.0 0.0 LBMOL/HR 952.0188 951.9378 0.8508 LBMOL/HR 0.0 0.0 0.0 LBMOL/HR 0.0 0.0 0.0 LBMOL/HR 0.0 0.0 0.0 04/11/2011 PAGE 5 |
| | FLOWSHEE | ET SECTIC | DN | |

| CONVERGENCE B | LOCK: \$OLVER01 | (CONTINUED |)) | | | | | |
|--------------------|-----------------|------------|------|----------|---------|-------|-----|--------|
| 22 MOLE-FLOW | S-106 MIXED | ISO-PROP | | LBMOL/HR | 0.0 | 0.0 | 0.0 | |
| 23 MOLE-FLOW | S-106 MIXED | SODIU-01 | | LBMOL/HR | 0.0 | 0.0 | 0.0 | |
| 24 MOLE-FLOW | S-106 MIXED | METHA-01 | | LBMOL/HR | 17.6689 | 17.66 | 87 | 0.1111 |
| 25 PRESSURE | S-106 MIXED | | PSIA | 42.0000 | 42.0000 | 0.0 | | |

*** ITERATION HISTORY *** ASPEN PLUS PLAT: WIN32 VER: 24.0

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FLOWSHEET SECTION

CONVERGENCE BLOCK: \$OLVER01 (CONTINUED)

TEAR STREAMS AND TEAR VARIABLES:

ITERATION MAX-ERR/TOL STREAM ID VARIABLE SUBSTREA COMPONEN ATTRIBUT ELEMENT

| 1 | 0.1000E+0 |)7 S-106 | PRESSURE | MIXED | |
|----|-----------|----------|-----------|-------|----------|
| 2 | 0.1000E+0 | 05 S-108 | PRESSURE | MIXED | |
| 3 | 6573. | S-106 | MOLE-FLOW | MIXED | METHA-01 |
| 4 | 6573. | S-108 | MOLE-FLOW | MIXED | METHA-01 |
| 5 | 2606. | S-106 | MOLE-FLOW | MIXED | METHA-01 |
| 6 | 2606. | S-108 | MOLE-FLOW | MIXED | METHA-01 |
| 7 | 1358. | S-106 | MOLE-FLOW | MIXED | METHA-01 |
| 8 | 1358. | S-108 | MOLE-FLOW | MIXED | METHA-01 |
| 9 | 785.3 | S-106 | MOLE-FLOW | MIXED | METHA-01 |
| 10 | 785.3 | S-108 | MOLE-FLOW | MIXED | METHA-01 |
| 11 | 478.2 | S-106 | MOLE-FLOW | MIXED | METHA-01 |
| 12 | 478.2 | S-108 | MOLE-FLOW | MIXED | METHA-01 |
| 13 | 448.4 | S-106 | MOLE-FLOW | MIXED | WATER |
| 14 | 448.4 | S-108 | MOLE-FLOW | MIXED | WATER |
| 15 | 283.2 | S-106 | MOLE-FLOW | MIXED | WATER |
| 16 | 283.2 | S-108 | MOLE-FLOW | MIXED | WATER |
| 17 | -236.2 | S-106 | MOLE-FLOW | MIXED | WATER |
| 18 | -236.2 | S-108 | MOLE-FLOW | MIXED | WATER |
| 19 | -159.6 | S-106 | MOLE-FLOW | MIXED | WATER |
| 20 | -159.6 | S-108 | MOLE-FLOW | MIXED | WATER |
| 21 | -107.0 | S-106 | MOLE-FLOW | MIXED | WATER |
| 22 | -107.0 | S-108 | MOLE-FLOW | MIXED | WATER |
| 23 | -71.40 | S-106 | MOLE-FLOW | MIXED | WATER |
| 24 | -71.40 | S-108 | MOLE-FLOW | MIXED | WATER |
| 25 | -47.45 | S-106 | MOLE-FLOW | MIXED | WATER |
| 26 | -47.45 | S-108 | MOLE-FLOW | MIXED | WATER |
| 27 | -31.46 | S-106 | MOLE-FLOW | MIXED | WATER |
| 28 | -31.46 | S-108 | MOLE-FLOW | MIXED | WATER |
| 29 | -20.82 | S-106 | MOLE-FLOW | MIXED | WATER |
| 30 | -20.82 | S-108 | MOLE-FLOW | MIXED | WATER |
| 31 | -13.77 | S-106 | MOLE-FLOW | MIXED | WATER |
| 32 | -13.77 | S-108 | MOLE-FLOW | MIXED | WATER |
| 33 | 200.2 | S-106 | MOLE-FLOW | MIXED | WATER |
| 34 | 200.2 | S-108 | MOLE-FLOW | MIXED | WATER |
| 35 | 129.5 | S-106 | MOLE-FLOW | MIXED | WATER |
| 36 | 129.5 | S-108 | MOLE-FLOW | MIXED | WATER |
| 37 | 84.36 | S-106 | MOLE-FLOW | MIXED | WATER |
| 38 | 84.36 | S-108 | MOLE-FLOW | MIXED | WATER |
| 39 | 55.20 | S-106 | MOLE-FLOW | MIXED | WATER |
| 40 | 55.20 | S-108 | MOLE-FLOW | MIXED | WATER |

| 41 | 36.23 | S-106 | MOLE-FLOW | MIXED | WATER | |
|-------|---------|------------|-----------|---------|----------|----|
| 42 | 36.23 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 43 | 23.82 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 44 | 23.82 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 45 | 15.68 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 46 | 15.68 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 47 | 10.33 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 48 | 10.33 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 49 | 6.811 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 50 | 6.811 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 51 | 4.491 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 52 | 4.491 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 53 | 2.962 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 54 | 2.962 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 55 | 1.954 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 56 | 1.954 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 57 | 1.289 | S-106 | MOLE-FLOW | MIXED | WATER | |
| 58 | 1.289 | S-108 | MOLE-FLOW | MIXED | WATER | |
| 59 | 0.8508 | S-106 | MOLE-FLOW | MIXED | WATER | |
| ASPEN | PLUS PI | LAT: WIN32 | VER: 24.0 | 04/11/2 | 011 PAGE | 27 |
| | | | | | | |

FLOWSHEET SECTION

CONVERGENCE BLOCK: \$OLVER02 _____ Tear Stream : S-121 S-119 Tolerance used: 0.100D-03 0.100D-03 Trace molefrac: 0.100D-05 0.100D-05 MAXIT= 70 WAIT 1 ITERATIONS BEFORE ACCELERATING $QMAX = 0.0 \quad QMIN = -5.0$ METHOD: WEGSTEIN STATUS: NOT CONVERGED TOTAL NUMBER OF ITERATIONS: 70 * * * BLOCK NOT CONVERGED * * *

* EXCEEDED MAXIMUM NUMBER OF ITERATIONS *

*** FINAL VALUES ***

VAR# TEAR STREAM VAR STREAM SUBSTREA COMPONEN ATTRIBUT ELEMENT UNIT VALUE PREV VALUE ERR/TOL

| 1 | TOTAL MOLEFLO | OW S- | 121 MIX | ED | LBMOL/ | 'HR | 1063.1236 | 1063.130 | -6.0587- |
|----|---------------|-------|---------|----------|--------|-------|-----------|----------|------------|
| 02 | | | | | | | | | |
| 2 | TOTAL MOLEFLO | OW S- | 119 MIX | ED | LBMOL/ | ΉR | 687.6446 | 687.6236 | 0.3051 |
| 3 | MOLE-FLOW | S-121 | MIXED | GLYCEROL | LBM | OL/HR | 0.0 | 0.0 | 0.0 |
| 4 | MOLE-FLOW | S-121 | MIXED | ACETOL | LBMOI | L/HR | 0.0 | 0.0 | 0.0 |
| 5 | MOLE-FLOW | S-121 | MIXED | WATER | LBMOL | _/HR | 395.0384 | 395.038 | 5 -3.6307- |
| | | | | | | | | | |

03 6 MOLE-FLOW S-121 MIXED HYDROGEN LBMOL/HR 494.5852 494.5904 -0.1047

 7
 MOLE-FLOW
 S-121
 MIXED
 PG
 LBMOL/HR
 166.1698
 166.1698
 -2.0683-04

 8
 MOLE-FLOW
 S-121
 MIXED
 N-PROP
 LBMOL/HR
 2.2627
 2.2628
 -0.4908

 9
 MOLE-FLOW
 S-121
 MIXED
 ETHYL-01
 LBMOL/HR
 0.0
 0.0

 10
 MOLE-FLOW
 S-121
 MIXED
 ISO-PROP
 LBMOL/HR
 4.9886
 4.9896
 -2.0148

 LBMOL/HR 4.9886 4.9896 -2.0148
 11 MOLE-FLOW
 S-121
 MIXED
 SODIU-01
 LBMOL/HR
 0.0
 0.0
 0.0

 12 MOLE-FLOW
 S-121
 MIXED
 METHA-01
 LBMOL/HR
 7.8826-02
 7.8827-02
 1.7315-02

 1.7513-02

 13 PRESSURE
 S-121 MIXED
 PSIA
 566.0365
 566.0365
 3.6707-04

 14 MASS ENTHALPY
 S-121 MIXED
 BTU/LB
 -3269.7542
 -3269.7482
 -1.8293-02

 15 MOLE-FLOW
 S-119 MIXED
 GLYCEROL
 LBMOL/HR
 0.0
 0.0

 16 MOLE-FLOW
 S-119 MIXED
 ACETOL
 LBMOL/HR
 0.0
 0.0

 17 MOLE-FLOW
 S-119 MIXED
 WATER
 LBMOL/HR
 17.0797
 17.0791
 0.3245

 18 MOLE-FLOW
 S-119 MIXED
 HYDROGEN
 LBMOL/HR
 665.5880
 665.5687

 0.2907
 19 MOLE-FLOW
 S-119 MIXED
 PG
 LBMOL/HR
 0.2401
 0.2401
 0.6432

 20 MOLE-FLOW
 S-119 MIXED
 N-PROP
 LBMOL/HR
 0.9994
 0.9993
 1.1812
 *
 21 MOLE-FLOW
 S-119
 MIXED
 ETHYL-01
 LBMOL/HR
 0.0
 0.0
 0.0

 22 MOLE-FLOW
 S-119
 MIXED
 ISO-PROP
 LBMOL/HR
 3.7261
 3.7252
 2.5376
 *
 23 MOLE-FLOW
 S-119
 MIXED
 SODIU-01
 LBMOL/HR
 0.0
 0.0
 0.0

 24 MOLE-FLOW
 S-119
 MIXED
 METHA-01
 LBMOL/HR
 1.1240-02
 1.1240-02
 0.4400 25 PRESSURE S-119 MIXED PSIA 600.0000 600.0000 0.0 26 MASS ENTHALPY S-119 MIXED BTU/LB -933.0325 -933.0067 -0.2767

*** ITERATION HISTORY *** ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 8

FLOWSHEET SECTION

CONVERGENCE BLOCK: \$OLVER02 (CONTINUED)

TEAR STREAMS AND TEAR VARIABLES:

ITERATION MAX-ERR/TOL STREAM ID VARIABLE SUBSTREA COMPONEN ATTRIBUT ELEMENT

| 1 | 0.1000E- | +07 S-121 | PRESSURE | MIXED | |
|----|----------|-----------|------------|--------|--------------|
| 2 | 0.1065E- | +05 S-121 | TOTAL MOLE | FLOW N | 1IXED |
| 3 | -0.1585E | +05 S-119 | MASS ENTHA | LPY MI | XED |
| 4 | 9985. | S-121 | MOLE-FLOW | MIXED | HYDROGEN |
| 5 | 0.5834E- | +05 S-119 | MOLE-FLOW | MIXE | D N-PROP |
| 6 | 7194. | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 7 | 6614. | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 8 | 3313. | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 9 | 3234. | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 10 | 1945. | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 11 | 1936. | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 12 | 1263. | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| | | | | | |

| 13 | 1257. | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
|----------------|-----------------------------|----------------|-----------|-------|----------------|
| 14 | 858.2 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 15 | 846.1 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 16 | 594.0 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 17 | 586.1 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 18 | 419.1 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 19 | 411.9 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 20 | 298.2 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 21 | 299.5 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 22 | 218.7 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 23 | 221.3 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 24 | 162.6 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 25 | 152.5 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 26 | 112.6 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 27 | 114.7 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 28 | 84.91 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 29 | 92.68 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 30 | 68.76 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 31 | 64.77 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 32 | 48.13 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 33 | -42.06 | S-119 | MOLE-FLOW | MIXED | WATER |
| 34 | 30.77 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 35 | 34.70 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 36 | 25.84 | S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 37 | 25.01 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 38 | 18.90 | S-121 | MOLE FLOW | MIXED | ISO-PROP |
| 39 | 18.19 | S-119 | MOLE FLOW | MIXED | ISO-PROP |
| 40 | 13.56 | S-121 | MOLE FLOW | MIXED | ISO-PROP |
| 40 41 | 13.30 | S-121 S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| $\frac{1}{12}$ | 9 9/9 | S-121 | MOLE TEOW | MIXED | ISO_PROP |
| 72 //3 | 0.575 | S_{-110} | MOLE-FLOW | MIXED | |
| ΛΛ | 7.373 7 1 AA | S-117 S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 44 15 | 6 780 | S-121 S-110 | MOLE-FLOW | MIXED | ISO-PROP |
| 45 46 | 5.060 | S-119 S-121 | MOLE-FLOW | MIXED | ISO-I KOI |
| 40 | 1 600 | S-121 S-110 | MOLE-FLOW | MIXED | ISO-I KOI |
| 47 19 | 4.009 | S-119 S 121 | MOLE-FLOW | MIXED | ISO-I KOI |
| 40 | 2.441 2.049 | S-121 S-110 | MOLE-FLOW | MIXED | ISO-FROF |
| 49 50 | 3.0 4 0 46.01 | S-119 S-110 | MOLE-FLOW | MIXED | WATED |
| 51 | 40.01 | S-119 S-110 | MOLE-FLOW | MIXED | WATED |
| 52 | -44.90 | S-119 S 121 | MOLE-FLOW | MIXED | |
| 52 53 | -9.99J 1.627 | S-121 S-110 | MOLE-FLOW | MIXED | N-FROF |
| 55 | -4.037 | S-119 S 110 | MOLE FLOW | MIXED | N-FKOF |
| 54 55 | 2 400 | S-119 S-110 | MOLE-FLOW | MIXED | FU METUA 01 |
| 55 56 | -3.499 | S-119 S-110 | MOLE-FLOW | MIXED | METHA-UI |
| 50 57 | 2.373 | S-119 S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 51 50 | 1.//4 | S-121 S-110 | MOLE-FLOW | MIXED | ISO-PROP |
| 30 50 | 1.939 | S-119 S-121 | MOLE-FLOW | MIXED | ISO-PROP |
| 59 60 | 1.44ð 46.00 | S-121 S-110 | MOLE-FLUW | MINED | 190-LKOL |
| 60 | 40.09 | 5-119 | MOLE-FLOW | MIXED | WATER |
| 01 | -43.64 | 5-119 | MOLE-FLOW | MIXED | WAIEK |
| 02 62 | -9.446 | S-121 | MOLE-FLOW | MIXED | N-PKUP |
| 63 | -4.204 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 04 | /.608 | 5-119 | MOLE-FLOW | MIXED | N-PKUP |
| 65 | -0.505 | S-119 | MOLE-FLOW | MIXED | N-PKOP |
| 60 | 4.037 | 8-119 | MOLE-FLOW | MIXED | 120-1401 |

| 6 | 7 - | 3.626 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
|------|------|-------|------------|-----------|---------|-------------|
| 6 | 8 | 3.239 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 6 | 9 - | 2.698 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| 7 | 0 | 2.538 | S-119 | MOLE-FLOW | MIXED | ISO-PROP |
| ASPE | N PL | US PI | LAT: WIN32 | VER: 24.0 | 04/11/2 | 2011 PAGE 9 |

FLOWSHEET SECTION

CONVERGENCE BLOCK: \$OLVER03

SPECS: DS-1 MAXIT= 30 STEP-SIZE= 1.0000 % OF RANGE MAX-STEP= 100. % OF RANGE XTOL= 1.000000E-08 THE NEW ALGORITHM WAS USED WITH BRACKETING=NO METHOD: SECANT STATUS: CONVERGED TOTAL NUMBER OF ITERATIONS: 152 NUMBER OF ITERATIONS ON LAST OUTER LOOP: 1

*** FINAL VALUES ***

| VAR# MA | NIPUL/TEAR-V | AR VARIABLE DESCRI | PTION | UNIT | VALUE | PREV |
|---------|--------------|--------------------|----------|----------|----------|--------|
| VALUE | ERR/TOL | | | | | |
| | | | | | | |
| 1 TOTAI | L MOLEFLOW | S-117.MIXED | LBMOL/HR | 171.8376 | 171.8376 | 0.3454 |

*** ITERATION HISTORY ***

DESIGN-SPEC ID: DS-1 ITERATED: TOTAL MOLEFLOW IN STREAM S-117 SUBSTREAM MIXED

ITERATION VARIABLE ERROR ERR/TOL

1 171.8 0.3454E-01 0.3454

CONVERGENCE BLOCK: \$OLVER04

SPECS: DS-2 MAXIT= 30 STEP-SIZE= 1.0000 % OF RANGE MAX-STEP= 100. % OF RANGE XTOL= 1.000000E-08 THE NEW ALGORITHM WAS USED WITH BRACKETING=NO METHOD: SECANT STATUS: CONVERGED TOTAL NUMBER OF ITERATIONS: 127 NUMBER OF ITERATIONS ON LAST OUTER LOOP: 1

*** FINAL VALUES ***

| VAR# MANIPUL/TEAR-VAR VARIABLE DESCRIPTION VALUE ERR/TOL | | UNIT | VALUE | PREV |
|----------------------------------------------------------|----------|----------|----------|--------|
| 1 TOTAL MOLEFLOW S-102.MIXED | LBMOL/HR | 298.8769 | 298.8769 | 0.4337 |

*** ITERATION HISTORY *** ASPEN PLUS PLAT: WIN32 VER: 24.0

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FLOWSHEET SECTION

CONVERGENCE BLOCK: \$OLVER04 (CONTINUED)

DESIGN-SPEC ID: DS-2 ITERATED: TOTAL MOLEFLOW IN STREAM S-102 SUBSTREAM MIXED

ITERATION VARIABLE ERROR ERR/TOL

1 298.9 0.4337E-02 0.4337

COMPUTATIONAL SEQUENCE

SEQUENCE USED WAS: \$OLVER01 D-101 HX-101 FSPLIT P-102 \$OLVER04 M-102 \$CRETURN \$OLVER04 \$S-101 P-101 H-101 \$CRETURN \$OLVER01 \$OLVER02 F-101 *HX-102 F-102 *HX-103 R-101 C-102 V-101 CP-101 \$OLVER03 T-101 M-101 \$CRETURN \$OLVER03 \$CRETURN \$OLVER03 \$CRETURN *\$OLVER03 \$CRETURN *\$OL

OVERALL FLOWSHEET BALANCE

| *** | MASS AND F | ENERGY BA | ALANCE * | ** | |
|------------|--------------|-----------|--------------|-------------|--------------|
| | IN OUT | GENER | ATION RE | ELATIVE DI | FF. |
| CONVENTION | IAL COMPON | ENTS | | | |
| (LBMOL/I | HR) | | | | |
| GLYCEROL | 168.457 | 0.286352 | 2E-03 -168.4 | 457 0.468 | 770E-09 |
| ACETOL | 0.00000 | 0.00000 | 0.00000 | 0.00000 | |
| WATER | 460.344 | 631.246 | 170.983 | 0.128936E- | 03 |
| HYDROGEN | 171.838 | 0.83994 | 7 -170.98 | 0.82447 | '1E-04 |
| PG | 0.00000 16 | 5.930 16 | 5.930 0.7 | 23638E-07 | |
| N-PROP | 0.00000 | 1.26342 | 1.26342 | 0.552076E-0 | 05 |
| ETHYL-01 | 0.00000 | 0.00000 | 0.00000 | 0.00000 | |
| ISO-PROP | 0.00000 | 1.26348 | 1.26342 | -0.474970E | -04 |
| SODIU-01 | 13.2729 | 13.2729 | 0.00000 | 0.00000 | |
| METHA-01 | 6.05218 | 6.05197 | 0.00000 | 0.348543E | E-04 |
| TOTAL BALA | NCE | | | | |
| MOLE(LBMOI | L/HR) 819. | .963 819 | .868 0.00 | 0000 0.11 | 6746E-03 |
| MASS(LB/HR |) 25123.2 | 25121.7 | 7 | 0.596787E-0 | 04 |
| ENTHALPY(B | TU/HR) -0.1 | 07053E+09 | -0.116457E | +09 | 0.807452E-01 |
| ASPEN PLUS | PLAT: WIN32 | VER: 24.0 | C | 04/11/2011 | PAGE 11 |

COMPONENTS

ID TYPE FORMULA NAME OR ALIAS **REPORT NAME** C3H8O3 GLYCEROL C C3H8O3 GLYCEROL ACETOL C C3H6O2-D1 C3H6O2-D1 ACETOL WATER C H2O H2O WATER HYDROGEN C H2 H2 HYDROGEN PG C C3H8O2-2 C3H8O2-2 PG N-PROP C C3H8O-1 N-PROP C3H8O-1 ETHYL-01 C C2H6O2 C2H6O2 ETHYL-01 ISO-PROP C C3H8O-2 C3H8O-2 **ISO-PROP** SODIU-01 C NACL NACL SODIU-01 METHA-01 C CH4O CH4O METHA-01 LISTID SUPERCRITICAL COMPONENT LIST HC-1 HYDROGEN ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 12 U-O-S BLOCK SECTION BLOCK: C-102 MODEL: HEATER _____ INLET STREAM: S-127 OUTLET STREAM: S-128 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 888.498 888.498 0.00000 MASS(LB/HR) 12004.1 12004.1 0.00000 ENTHALPY(BTU/HR) -0.443889E+08 -0.518861E+08 0.144493 *** INPUT DATA *** TWO PHASE TP FLASH

IWO THASE IT TEASHSPECIFIED TEMPERATUREF225.000PRESSURE DROPPSI5.00000MAXIMUM NO. ITERATIONS30CONVERGENCE TOLERANCE0.000100000

*** RESULTS *** OUTLET TEMPERATURE F 225.00 OUTLET PRESSURE PSIA 553.28 HEAT DUTY BTU/HR -0.74972E+07 OUTLET VAPOR FRACTION 0.58054 PRESSURE-DROP CORRELATION PARAMETER 0.27858E+06 V-L PHASE EQUILIBRIUM :

 COMP
 F(I)
 X(I)
 Y(I)
 K(I)

 WATER
 0.36651
 0.82793
 0.33113E-01
 0.39995E-01

 HYDROGEN
 0.55587
 0.36804E-03
 0.95724
 2600.8

 PG
 0.70286E-01
 0.16692
 0.46548E-03
 0.27886E-02

 N-PROP
 0.21637E-02
 0.24765E-02
 0.19376E-02
 0.78236

 ISO-PROP
 0.50959E-02
 0.21508E-02
 0.72239E-02
 3.3586

 METHA-01
 0.77815E-04
 0.15535E-03
 0.21791E-04
 0.14027

 ASPEN PLUS
 PLAT: WIN32
 VER: 24.0
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U-O-S BLOCK SECTION

BLOCK: C-103 MODEL: HEATER

INLET STREAM: S-134 OUTLET STREAM: S-135 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 164.300 164.300 0.00000 MASS(LB/HR) 12379.8 12379.8 0.00000 ENTHALPY(BTU/HR) -0.324322E+08 -0.350141E+08 0.737392E-01

*** INPUT DATA ***TWOPHASE TP FLASHSPECIFIED TEMPERATUREF90.0000PRESSURE DROPPSI5.00000MAXIMUM NO. ITERATIONS30CONVERGENCE TOLERANCE0.000100000

*** RESULTS *** OUTLET TEMPERATURE F 90.000 OUTLET PRESSURE PSIA 15.000 HEAT DUTY BTU/HR -0.25819E+07 OUTLET VAPOR FRACTION 0.0000 PRESSURE-DROP CORRELATION PARAMETER 0.13144E+08

V-L PHASE EQUILIBRIUM :

COMPF(I)X(I)Y(I)K(I)WATER0.12859E-010.12859E-010.927860.12296PG0.987140.987140.72136E-010.12451E-03N-PROP0.22134E-070.22134E-070.17198E-050.13239ISO-PROP0.55639E-110.55639E-110.21457E-080.65712METHA-010.99242E-080.99242E-080.14898E-050.25580ASPEN PLUSPLAT: WIN32VER: 24.004/11/2011PAGE 14

BLOCK: CP-101 MODEL: COMPR

INLET STREAM:S-130OUTLET STREAM:S-131PROPERTY OPTION SET:PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 515.807 515.807 0.00000 MASS(LB/HR) 1605.65 1605.65 0.141608E-15 ENTHALPY(BTU/HR) -0.181851E+07 -0.174555E+07 -0.401195E-01

*** INPUT DATA ***

ISENTROPIC CENTRIFUGAL COMPRESSOROUTLET PRESSURE PSIA600.000ISENTROPIC EFFICIENCY0.80000MECHANICAL EFFICIENCY1.00000

*** RESULTS ***

| | INDICATE | ED HORSEPC | WER F | REQUIREM | ENT 1 | HP | 2 | 28.6734 |
|---|-----------|--------------------|--------------|------------|-------|----------|-------|---------|
| | BRAKE | HORSEPOW | ER RE | QUIREMEN | NT HP | | 28. | 6734 |
| | NET WOR | K REQUIRED |) | HP | | 28.6734 | | |
| | POWER L | OSSES | H | ΗP | 0.0 | | | |
| | ISENTRO | PIC HORSEPO | WER F | REQUIREM | ENT 1 | HP | - | 22.9387 |
| | CALCULA | TED OUTLE | Т ТЕМІ | P F | | 244.41 | 7 | |
| | ISENTRO | PIC TEMPERA | ATURE | F | | 240.520 | | |
| | EFFICIEN | CY (POLYTR | /ISENT | R) USED | | 0.80 | 0000 | |
| | OUTLET V | VAPOR FRAC | TION | | | 1.00000 | | |
| | HEAD DE | VELOPED, | FT-LI | BF/LB | | 28,286.7 | | |
| | MECHAN | ICAL EFFICIE | ENCY U | JSED | | 1.000 | 000 | |
| | INLET HE | AT CAPACIT | Y RAT | IO | | 1.3882 | 5 | |
| | INLET VC | DLUMETRIC F | FLOW F | RATE , CUF | T/HR | 6 | 5,949 | 9.72 |
| | OUTLET V | VOLUMETRI | CFLOW | V RATE, CU | JFT/H | R | 6,5 | 97.33 |
| | INLET CO | OMPRESSIBII | LITY FA | ACTOR | | 1.01 | 458 | |
| | OUTLET (| COMPRESSIB | ILITY I | FACTOR | | 1.(|)156 | 6 |
| | AV. ISEN | Г. VOL. EXPC | NENT | | | 1.40856 | | |
| | AV. ISEN | Г. TEMP EXP | ONENT | 1 | | 1.38215 | | |
| | AV. ACTU | JAL VOL. EX | PONEN | Т | | 1.5579 | 1 | |
| | AV. ACTU | JAL TEMP EX | KPONE | T | | 1.5266 | 53 | |
| A | ASPEN PLU | S PLAT: WI | N32 V | VER: 24.0 | | 04/11/2 | 011 | PAGE 15 |

U-O-S BLOCK SECTION

BLOCK: CP-101 MODEL: COMPR (CONTINUED)

BLOCK: D-101 MODEL: RADFRAC

INLETS - S-108 STAGE 4 OUTLETS - S-109 STAGE 1

S-114 STAGE 6 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT **RELATIVE DIFF.** TOTAL BALANCE MOLE(LBMOL/HR) 1138.06 1138.06 0.00000 MASS(LB/HR) 33229.6 33229.6 -0.217931E-11 ENTHALPY(BTU/HR) -0.161726E+09 -0.162466E+09 0.455082E-02 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 16

U-O-S BLOCK SECTION

BLOCK: D-101 MODEL: RADFRAC (CONTINUED)

**** INPUT PARAMETERS ****

NUMBER OF STAGES 6 ALGORITHM OPTION **STANDARD** ABSORBER OPTION NO INITIALIZATION OPTION **STANDARD** HYDRAULIC PARAMETER CALCULATIONS NO INSIDE LOOP CONVERGENCE METHOD BROYDEN DESIGN SPECIFICATION METHOD NESTED MAXIMUM NO. OF OUTSIDE LOOP ITERATIONS 25 MAXIMUM NO. OF INSIDE LOOP ITERATIONS 10 MAXIMUM NUMBER OF FLASH ITERATIONS 30 FLASH TOLERANCE 0.000100000 OUTSIDE LOOP CONVERGENCE TOLERANCE 0.000100000

**** COL-SPECS ****

MOLAR VAPOR DIST / TOTAL DIST0.0MOLAR REFLUX RATIO0.20000MOLAR BOTTOMS RATELBMOL/HR375.500

**** PROFILES ****

P-SPEC STAGE 1 PRES, PSIA 20.0000

**** RESULTS ****

*** COMPONENT SPLIT FRACTIONS ***

OUTLET STREAMS

S-109 S-114 COMPONENT: GLYCEROL .49995E-05 1.0000 WATER .78257 .21743 METHA-01 .99617 .38252E-02 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/20

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U-O-S BLOCK SECTION

BLOCK: D-101 MODEL: RADFRAC (CONTINUED)

*** SUMMARY OF KEY RESULTS ***

TOP STAGE TEMPERATURE F 221.380 BOTTOM STAGE TEMPERATURE F 247.890 TOP STAGE LIQUID FLOW LBMOL/HR 152.513 BOTTOM STAGE LIQUID FLOW LBMOL/HR 375.500 TOP STAGE VAPOR FLOW LBMOL/HR 0.0 LBMOL/HR 896.933 BOILUP VAPOR FLOW MOLAR REFLUX RATIO 0.20000 MOLAR BOILUP RATIO 2.38864 CONDENSER DUTY (W/O SUBCOOL) BTU/HR -0.164793+08REBOILER DUTY BTU/HR 0.157399+08

**** MAXIMUM FINAL RELATIVE ERRORS ****

DEW POINT0.37845E-06STAGE= 5BUBBLE POINT0.14269E-05STAGE= 5COMPONENT MASS BALANCE0.92840E-06STAGE= 2COMPONENT MASS0.93821E-06STAGE= 1

**** PROFILES ****

NOTE REPORTED VALUES FOR STAGE LIQUID AND VAPOR RATES ARE THE FLOWS FROM THE STAGE INCLUDING ANY SIDE PRODUCT.

| | | | ENTHALPY | | | | |
|----|--------|-----------|------------|-------------|-----------|-----------|----------|
| ST | AGE TE | EMPERATUR | E PRESSUI | RE | BTU/LBM | IOL HI | EAT DUTY |
| | F | PSIA | LIQUID | VAPOR | BTU/H | R | |
| | | | | | | | |
| 1 | 221.38 | 20.000 | -0.12036E+ | -06 -99448 | 81647 | 79+08 | |
| 2 | 229.97 | 20.000 | -0.12067E+ | 06 -0.1023 | 35E+06 | | |
| 3 | 230.24 | 20.000 | -0.12096E+ | -06 -0.1024 | 41E+06 | | |
| 4 | 233.48 | 20.000 | -0.14016E+ | -06 -0.1024 | 45E+06 | | |
| 5 | 234.16 | 20.000 | -0.14034E+ | 06 -0.1027 | 70E+06 | | |
| 6 | 247.89 | 20.000 | -0.18824E+ | 06 -0.102 | 73E+06 .1 | 5740+08 | |
| | | | | | | | |
| ST | AGE | FLOW RATE | E F | EED RAT | E] | PRODUCT I | RATE |
| | LBN | /IOL/HR | LBMO | L/HR | LBM | IOL/HR | |
| | LIQUII | O VAPOR | LIQUID | VAPOR | MIXED | LIQUID | VAPOR |
| 1 | 915.1 | 0.000 | | 762.56 | 539 | | |

| 4 | 155.1 | 71J.1 | | | |
|----|--------|--------|-----------|-----------|----|
| 3 | 151.7 | 915.6 | 17.2 | 337 | |
| 4 | 1278. | 897.0 | 1120.8301 | | |
| 5 | 1272. | 902.3 | | | |
| 6 | 375.5 | 896.9 | | 375.500 |)0 |
| AS | PEN PL | US PLA | T: WIN32 | VER: 24.0 | 04 |
| AS | PEN PL | US PLA | T: WIN32 | VER: 24.0 | C |

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U-O-S BLOCK SECTION

BLOCK: D-101 MODEL: RADFRAC (CONTINUED)

**** MASS FLOW PROFILES ****

2 152 1

015 1

STAGE FLOW RATE FEED RATE PRODUCT RATE LB/HR LB/HR LB/HR LIQUID VAPOR LIQUID VAPOR MIXED LIQUID VAPOR 1 0.1678E+05 0.000 .13985 + 052 2764. 0.1678E+05 3 2760. 0.1675E+05 329.1384 4 0.3558E+05 0.1642E+05 .32900+05 5 0.3548E+05 0.1634E+05 6 0.1924E+05 0.1623E+05 .19245 + 05

**** **** MOLE-X-PROFILE STAGE GLYCEROL WATER METHA-01 1 0.11044E-05 0.97692 0.23082E-01 2 0.11364E-03 0.99763 0.22605E-02 3 0.21073E-02 0.99595 0.19438E-02 0.86465 4 0.13209 0.32605E-02 5 0.13306 0.86615 0.79037E-03 6 0.44862 0.55120 0.17999E-03

**** **** MOLE-Y-PROFILE STAGE **GLYCEROL** WATER METHA-01 0.73210E-08 0.81160 0.18840 1 2 0.11044E-05 0.97692 0.23082E-01 3 0.19916E-04 0.98038 0.19601E-01 4 0.35090E-03 0.98115 0.18501E-01 5 0.36087E-03 0.99510 0.45424E-02

6 0.94577E-03 0.99801

**** **** K-VALUES STAGE **GLYCEROL** WATER METHA-01 1 0.66287E-02 0.83078 8.1622 2 0.97191E-02 0.97924 10.211 3 0.94510E-02 0.98437 10.084 4 0.26566E-02 1.1347 5.6743 5 0.27122E-02 1.1489 5.7473 6 0.21082E-02 1.8106 5.8109

**** MASS-X-PROFILE **** STAGE GLYCEROL WATER METHA-01 1 0.55463E-05 0.95967 0.40328E-01 2 0.57963E-03 0.99541 0.40115E-02

0.10459E-02

| 3 | 0.10664E | E-01 0.98591 | 0.34225E-02 | | |
|------|----------|--------------|-------------|------------|---------|
| 4 | 0.43685 | 0.55939 | 0.37518E-02 | | |
| 5 | 0.43947 | 0.55962 | 0.90826E-03 | | |
| 6 | 0.80614 | 0.19375 | 0.11253E-03 | | |
| ASPE | EN PLUS | PLAT: WIN3 | 2 VER: 24.0 | 04/11/2011 | PAGE 19 |

BLOCK: D-101 MODEL: RADFRAC (CONTINUED)

 MASS-Y-PROFILE

 STAGE
 GLYCEROL
 WATER
 METHA-01

 1
 0.32638E-07
 0.70778
 0.29222

 2
 0.55463E-05
 0.95967
 0.40328E-01

 3
 0.10027E-03
 0.96556
 0.34336E-01

 4
 0.17658E-02
 0.96584
 0.32392E-01

 5
 0.18356E-02
 0.99013
 0.80389E-02

 6
 0.48122E-02
 0.99334
 0.18515E-02

BLOCK: D-102 MODEL: RADFRAC

INLETS - S-133 STAGE 10 OUTLETS - S-137 STAGE 1 S-136 STAGE 1 S-134 STAGE 16 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 547.323 547.323 -0.207714E-15 MASS(LB/HR) 19591.2 19591.2 -0.928471E-15 ENTHALPY(BTU/HR) -0.781100E+08 -0.795924E+08 0.186258E-01 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 20

U-O-S BLOCK SECTION

BLOCK: D-102 MODEL: RADFRAC (CONTINUED)

**** INPUT PARAMETERS ****

NUMBER OF STAGES16ALGORITHM OPTIONSTANDARDINITIALIZATION OPTIONSTANDARDHYDRAULIC PARAMETER CALCULATIONSNOINSIDE LOOP CONVERGENCE METHODNEWTONDESIGN SPECIFICATION METHODNESTEDMAXIMUM NO. OF OUTSIDE LOOP ITERATIONS25

MAXIMUM NO. OF INSIDE LOOP ITERATIONS 10 MAXIMUM NUMBER OF FLASH ITERATIONS 30 FLASH TOLERANCE 0.000100000 OUTSIDE LOOP CONVERGENCE TOLERANCE 0.000100000

**** COL-SPECS ****

| MOLAR REFLUX RATIO | | 0.13000 |
|-----------------------|----------|---------|
| MOLAR BOTTOMS RATE | LBMOL/HR | 164.300 |
| CONDENSER TEMPERATURE | F | 130.000 |

**** PROFILES ****

P-SPEC STAGE 1 PRES, PSIA 20.0000

> ******* **** RESULTS **** *************

*** COMPONENT SPLIT FRACTIONS ***

OUTLET STREAMS

S-137 S-136 S-134 COMPONENT: .85286E-03 .99356 WATER .55898E-02 HYDROGEN .99992 .83472E-04 0.0000 .53556E-05 .22549E-01 .97745 PG N-PROP .72514 .27486 .28784E-05 ASPEN PLUS PLAT: WIN32 VER: 24.0

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U-O-S BLOCK SECTION

BLOCK: D-102 MODEL: RADFRAC (CONTINUED)

*** COMPONENT SPLIT FRACTIONS ***

OUTLET STREAMS

S-137 S-136 S-134 COMPONENT: ISO-PROP .96809 .31911E-01 .72352E-09 METHA-01 .12954E-01 .98702 .24125E-04

*** SUMMARY OF KEY RESULTS ***

TOP STAGE TEMPERATURE F 130.000 BOTTOM STAGE TEMPERATURE F 364.027 TOP STAGE LIQUID FLOW LBMOL/HR 49.7930 BOTTOM STAGE LIQUID FLOW LBMOL/HR 164.300 TOP STAGE VAPOR FLOW LBMOL/HR 3.30330 BOILUP VAPOR FLOW LBMOL/HR 252.798

MOLAR REFLUX RATIO0.13000MOLAR BOILUP RATIO1.53864CONDENSER DUTY (W/O SUBCOOL)BTU/HRREBOILER DUTYBTU/HR7,106,170.

**** MAXIMUM FINAL RELATIVE ERRORS ****

DEW POINT0.33817E-05STAGE= 15BUBBLE POINT0.52790E-05STAGE= 1COMPONENT MASS BALANCE0.72629E-08STAGE= 5ENERGY BALANCE0.44363E-06STAGE= 14

ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 22

U-O-S BLOCK SECTION

BLOCK: D-102 MODEL: RADFRAC (CONTINUED)

**** PROFILES ****

NOTE REPORTED VALUES FOR STAGE LIQUID AND VAPOR RATES ARE THE FLOWS FROM THE STAGE INCLUDING ANY SIDE PRODUCT.

| | | | ENTHALPY | ľ | | | |
|-----|---------|----------|------------|------------|----------|-----------|-----------|
| ST. | AGE TEN | MPERATUR | RE PRESSU | RE | BTU/LB | MOL | HEAT DUTY |
| | F | PSIA | LIQUID | VAPOR | BTU/ | HR | |
| | | | | | | | |
| 1 | 130.00 | 20.000 | -0.12347E- | +06 -8309 | 185 | 886+07 | |
| 2 | 231.02 | 20.000 | -0.12239E- | +06 -0.103 | 32E+06 | | |
| 3 | 231.13 | 20.000 | -0.12289E- | +06 -0.103 | 43E+06 | | |
| 4 | 231.20 | 20.000 | -0.12323E- | +06 -0.103 | 49E+06 | | |
| 8 | 231.41 | 20.000 | -0.12462E- | +06 -0.103 | 63E+06 | | |
| 9 | 231.52 | 20.000 | -0.12591E- | +06 -0.103 | 69E+06 | | |
| 10 | 233.20 | 20.000 | -0.14789E | 406 -0.10 | 399E+06 | | |
| 11 | 233.32 | 20.000 | -0.14788E | 406 -0.10 | 398E+06 | | |
| 12 | 233.34 | 20.000 | -0.14790E | +06 -0.10 | 398E+06 | | |
| 15 | 316.56 | 20.000 | -0.19288E | +06 -0.12 | 300E+06 | | |
| 16 | 364.03 | 20.000 | -0.19740E | +06 -0.16 | 184E+06 | .71062+07 | |
| | | | | | | | |
| ST. | AGE F | LOW RATI | E F | FEED RAT | Έ | PRODUC | ΓRATE |
| | LBM | OL/HR | LBMO | OL/HR | LB | MOL/HR | |
| | LIQUID | VAPOR | LIQUID | VAPOR | MIXEI |) LIQUII | O VAPOR |
| 1 | 429.5 | 3.303 | | 379.7 | 197 3.30 |)33 | |
| 2 | 55.17 | 432.8 | | | | | |
| 3 | 55.06 | 438.2 | | | | | |
| 4 | 54.99 | 438.1 | | | | | |
| 8 | 54.64 | 437.8 | | | | | |
| 9 | 54.02 | 437.7 | 91.8493 | | | | |
| 10 | 511.4 | 345.2 4 | 55.4736 | | | | |
| 11 | 511.5 | 347.1 | | | | | |
| 12 | 511.4 | 347.2 | | | | | |
| 15 | 417.1 | 285.0 | | | | | |
| 16 | 164.3 | 252.8 | | 164.3 | 000 | | |

| LOW RATE | F | EED RATH | E P | RODUCT F | RATE |
|------------|----------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| R | LB/HR LB/HR | | | | |
| VAPOR | LIQUID | VAPOR | MIXED | LIQUID | VAPOR |
| 136.2 | | 7075.31 | 26 136.16 | 01 | |
| 8139. | | | | | |
| 8274. | | | | | |
| 8292. | | | | | |
| 8336. | | | | | |
| 8353. | 1803.477 | 6 | | | |
| +05 6587 | .17788+05 | | | | |
| +05 6596. | | | | | |
| JS PLAT: W | IN32 VE | R: 24.0 | 04/ | 11/2011 PA | GE 23 |
| | LOW RATE R VAPOR 136.2 8139. 8274. 8292. 8336. 8353. +05 6587. +05 6596. VS PLAT: W | LOW RATE F R LB/HR VAPOR LIQUID 136.2 8139. 8274. 8292. 8336. 8353. 1803.477 +05 658717788+05 +05 6596. US PLAT: WIN32 VE | LOW RATE FEED RATH R LB/HR L VAPOR LIQUID VAPOR 136.2 7075.31 8139. 8274. 8292. 8336. 8353. 1803.4776 +05 658717788+05 +05 6596. US PLAT: WIN32 VER: 24.0 | LOW RATE FEED RATE P R LB/HR LB/HR VAPOR LIQUID VAPOR MIXED 136.2 7075.3126 136.160 8139. 8274. 8292. 8336. 8353. 1803.4776 +05 658717788+05 +05 6596. US PLAT: WIN32 VER: 24.0 04/ | LOW RATE FEED RATE PRODUCT F R LB/HR LB/HR VAPOR LIQUID VAPOR MIXED LIQUID 136.2 7075.3126 136.1601 8139. 8274. 8292. 8336. 8353. 1803.4776 +05 658717788+05 +05 6596. US PLAT: WIN32 VER: 24.0 04/11/2011 PA |

BLOCK: D-102 MODEL: RADFRAC (CONTINUED)

**** MASS FLOW PROFILES ****

STAGEFLOW RATEFEED RATEPRODUCT RATELB/HRLB/HRLB/HRLIQUIDVAPORLIQUIDVAPOR12 0.1898E+056593.150.2930E+0515 0.2930E+059912..12380+05

**** MOLE-X-PROFILE ****

| STA | GE W | ATER | HYDR | ROGEN | PG | N-PRO | P ISO-PROP |
|-----|---------|--------|------------|----------|------|--------------|-------------|
| 1 | 0.9889 | 5 0.1 | 8464E-06 | 0.98535E | -02 | 0.91452E-03 | 0.10618E-03 |
| 2 | 0.9784 | 6 0.6 | 58408E-08 | 0.21505E | -01 | 0.98687E-05 | 0.16374E-05 |
| 3 | 0.9721 | 5 0.7 | 3785E-08 | 0.27815E | -01 | 0.11224E-04 | 0.19598E-05 |
| 4 | 0.9677 | 9 0.7 | '8268E-08 | 0.32174E | -01 | 0.12617E-04 | 0.22329E-05 |
| 8 | 0.9503 | 9 0.9 | 7553E-08 | 0.49569E | -01 | 0.19399E-04 | 0.36102E-05 |
| 9 | 0.9344′ | 7 0.1 | 1705E-07 | 0.65473E | -01 | 0.27469E-04 | 0.53297E-05 |
| 10 | 0.6712 | . 0. | 14078E-10 | 0.32845 | 0 | .22769E-03 0 | .27967E-04 |
| 11 | 0.6715 | 61 0.0 | 66732E-15 | 0.32843 | 0 | .46357E-04 0 | .16307E-05 |
| 12 | 0.6712 | .0. 8 | 31658E-19 | 0.32871 | 0 | .94426E-05 0 | .95170E-07 |
| 15 | 0.1006 | 60 0. | .0000 0. | .89940 | 0.15 | 563E-06 0.72 | 542E-10 |
| 16 | 0.1285 | 9E-01 | 0.95267E-2 | 0.9871 | 14 | 0.22134E-07 | 0.55639E-11 |

**** MOLE-X-PROFILE ****

STAGE METHA-01

- 1 0.17568E-03
- 2 0.20983E-04
- 3 0.19715E-04
- 4 0.20420E-04
- 8 0.23301E-04
- 0 0.25501E-04
- 9 0.25804E-04
- 10 0.33827E-04
- 11 0.14080E-04
- 12 0.58571E-05
- 15 0.11739E-06

| | **** MOLE-Y-PROFILE **** | | | | | | | | |
|------|--------------------------|--------------|------------|-----------|------------|--------------|--|--|--|
| STA | GE WA | TER HY | DROGEN | PG | N-PROI | P ISO-PROP | | | |
| 1 | 0.97583E | E-01 0.25425 | 0.26902 | 2E-03 0.2 | 27734 0.1 | 37029 | | | |
| 2 | 0.98215 | 0.19407E- | 02 0.97804 | 4E-02 0.3 | 30243E-02 | 0.29314E-02 | | | |
| 3 | 0.98091 | 0.19168E- | 02 0.1124 | 8E-01 0.2 | 28845E-02 | 0.28836E-02 | | | |
| 4 | 0.98012 | 0.19173E- | 02 0.1203 | 9E-01 0.2 | 28853E-02 | 0.28843E-02 | | | |
| 8 | 0.97817 | 0.19186E- | 02 0.1398 | 0E-01 0.2 | 28880E-02 | 0.28864E-02 | | | |
| 9 | 0.97741 | 0.19192E- | 02 0.1473 | 9E-01 0.2 | 28892E-02 | 0.28873E-02 | | | |
| 10 | 0.98091 | 0.43733E | -06 0.1662 | 4E-01 0. | 16420E-02 | 0.70432E-03 | | | |
| 11 | 0.98288 | 0.20742E | -10 0.1669 | 3E-01 0. | 33545E-03 | 0.41205E-04 | | | |
| 12 | 0.98320 | 0.98310E | -15 0.1671 | 3E-01 0. | 68283E-04 | 0.24024E-05 | | | |
| 15 | 0.71131 | 0.0000 | 0.28869 | 0.10907 | 7E-05 0.10 | 955E-08 | | | |
| 16 | 0.15763 | 0.43055E | -18 0.8423 | 0.24 | 239E-06 0 | .11607E-09 | | | |
| ASPE | N PLUS | PLAT: WIN | 32 VER: 2 | 24.0 | 04/11/ | 2011 PAGE 24 | | | |

BLOCK: D-102 MODEL: RADFRAC (CONTINUED)

**** MOLE-Y-PROFILE ****

STAGE METHA-01

- 1 0.26504E-03
- 2 0.17636E-03
- 3 0.15688E-03
- 4 0.15675E-03
- 8 0.15717E-03
- 9 0.15733E-03
- 10 0.11946E-03
- 11 0.49834E-04
- 12 0.20738E-04
- 15 0.13554E-05
- 16 0.18723E-06

| | | **** | K-VALU | ES | **: | ** | | | | |
|-----|--------|--------|-----------|--------|--------|--------|--------|--------|-------|-----|
| STA | AGE V | WATER | . HYD | DROG | EN | PG | N-P | ROP | ISO-P | ROP |
| 1 | 0.9867 | 73E-01 | 0.13770E | +07 (| 0.2730 | 2E-01 | 303.26 | 348 | 7.3 | |
| 2 | 1.003 | 8 0.2 | 28369E+06 | 5 0.45 | 5479 | 306. | 45 | 1790.3 | | |
| 3 | 1.009 | 0 0.2 | 25979E+06 | 5 0.40 | 0438 | 257. | 00 | 1471.3 | | |
| 4 | 1.012 | .7 0.2 | 24496E+06 | 5 0.3 | 7417 | 228. | 69 | 1291.7 | | |
| 8 | 1.029 | 2 0.1 | 19668E+06 | 5 0.28 | 8204 | 148. | .88 ~ | 799.55 | | |
| 9 | 1.045 | 9 0.1 | 16396E+06 | 5 0.22 | 2512 | 105. | 18 | 541.75 | | |
| 10 | 1.461 | 13 3 | 1064. (| 0.5061 | 2E-01 | 7.21 | 16 2 | 25.184 | | |
| 11 | 1.463 | 37 3 | 1082. (| 0.5082 | 28E-01 | 7.23 | 62 2 | 25.267 | | |
| 12 | 1.464 | 47 3 | 1053. (| 0.5084 | 4E-01 | 7.23 | 14 2 | 25.244 | | |
| 15 | 7.070 | 04 6 | 467.1 (| 0.3209 | 98 | 7.0087 | 15. | 102 | | |
| 16 | 12.25 | 59 4 | 519.5 (| 0.8533 | 34 | 10.951 | 20. | 862 | | |
| | | | | | | | | | | |

**** K-VALUES ****

- STAGE METHA-01
 - 1 1.5087
 - 2 8.4048
 - 3 7.9573

- 4 7.6762 8 6.7452 9 6.0971 10 3.5313 11 3.5394 12 3.5407 15 11.547
- 16 18.866

**** MASS-X-PROFILE ****

| STA | GE WA | TER HYDR | OGEN P | G N-PF | ROP IS | O-PROP |
|------|----------|-------------|------------|--------------|------------|--------|
| 1 | 0.95616 | 0.19976E-07 | 0.40241E-0 | 1 0.29496E-0 | 0.34246 | E-03 |
| 2 | 0.91499 | 0.71582E-09 | 0.84943E-0 | 1 0.30785E-0 | 0.51077 | 'E-05 |
| 3 | 0.89211 | 0.75766E-09 | 0.10782 | 0.34358E-04 | 0.59994E- | -05 |
| 4 | 0.87680 | 0.79346E-09 | 0.12312 | 0.38130E-04 | 0.67482E- | -05 |
| 8 | 0.81938 | 0.94113E-09 | 0.18051 | 0.55792E-04 | 0.10383E- | -04 |
| 9 | 0.77154 | 0.10814E-08 | 0.22833 | 0.75655E-04 | 0.14679E- | -04 |
| 10 | 0.32593 | 0.76490E-12 | 0.67363 | 0.36879E-03 | 0.45299E | -04 |
| 11 | 0.32614 | 0.36267E-16 | 0.67377 | 0.75106E-04 | 0.26421E | -05 |
| 12 | 0.32590 | 0.17199E-20 | 0.67408 | 0.15293E-04 | 0.15413E | -06 |
| 15 | 0.25799E | E-01 0.0000 | 0.97420 | 0.13313E-06 | 0.62055E- | 10 |
| ASPE | N PLUS | PLAT: WIN32 | VER: 24.0 | 04/ | 11/2011 PA | AGE 25 |

U-O-S BLOCK SECTION

BLOCK: D-102 MODEL: RADFRAC (CONTINUED)

**** MASS-X-PROFILE ****

 STAGE
 WATER
 HYDROGEN
 PG
 N-PROP
 ISO-PROP

 16
 0.30745E-02
 0.25488E-23
 0.99693
 0.17653E-07
 0.44376E-11

**** MASS-X-PROFILE ****

STAGE METHA-01

- 1 0.30211E-03
- 2 0.34900E-04
- 3 0.32178E-04
- 4 0.32905E-04
- 8 0.35730E-04
- 9 0.37894E-04
- 10 0.29213E-04
- 11 0.12163E-04
- 12 0.50576E-05
- 15 0.53540E-07
- 16 0.42203E-08

**** MASS-Y-PROFILE ****

| STA | GE | WAT | ER | HYDR | OGEN | PG | N-PROI | P ISO-PROP |
|-----|------|--------|--------|--------|---------|--------|-------------|-------------|
| 1 | 0.42 | 649E-0 | 0.12 | 435E-0 | 1 0.496 | 64E-03 | 0.40435 | 0.53986 |
| 2 | 0.94 | 088 | 0.2080 | 3E-03 | 0.39576 | E-01 (| 0.96646E-02 | 0.93679E-02 |
| 3 | 0.93 | 584 | 0.2046 | 4E-03 | 0.45329 | E-01 (| 0.91801E-02 | 0.91773E-02 |
| 4 | 0.93 | 282 | 0.2041 | 9E-03 | 0.48397 | E-01 (| 0.91605E-02 | 0.91573E-02 |
| 8 | 0.92 | 544 | 0.2031 | 1E-03 | 0.55866 | E-01 (| 0.91145E-02 | 0.91096E-02 |
| 9 | 0.92 | 258 | 0.2027 | 1E-03 | 0.58766 | E-01 (| 0.90971E-02 | 0.90914E-02 |
| 10 | 0.92 | 2612 | 0.4620 |)3E-07 | 0.66295 | 5E-01 | 0.51714E-02 | 0.22183E-02 |

| 11 0.93187 0.22005E-11 0.66853E-01 0.10609E-02 0.13032E-03 |
|------------------------------------------------------------|
| 12 0.93277 0.10437E-15 0.66973E-01 0.21610E-03 0.76031E-05 |
| 15 0.36842 0.0000 0.63158 0.18846E-05 0.18928E-08 |
| 16 0.42423E-01 0.12966E-19 0.95758 0.21761E-06 0.10421E-09 |
| |
| **** MASS-Y-PROFILE **** |
| STAGE METHA-01 |
| 1 0.20603E-03 |
| 2 0.30050E-03 |
| 3 0.26620E-03 |
| 4 0.26534E-03 |
| 8 0.26447E-03 |
| 9 0.26413E-03 |
| 10 0.20060E-03 |
| 11 0.84035E-04 |
| 12 0.34993E-04 |
| 15 0.12487E-05 |
| 16 0.89620E-07 |
| ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 26 |
| |
| U-O-S BLOCK SECTION |

BLOCK: F-101 MODEL: FLASH2

INLET STREAM: S-121 OUTLET VAPOR STREAM: S-123 OUTLET LIQUID STREAM: S-122 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 1063.13 1063.13 0.00000 MASS(LB/HR) 21196.9 21196.9 0.343256E-15 ENTHALPY(BTU/HR) -0.693085E+08 -0.693085E+08 -0.166199E-06

*** INPUT DATA *** TWO PHASE PQ FLASH SPECIFIED PRESSURE PSIA SPECIFIED HEAT DUTY BTU/HR MAXIMUM NO. ITERATIONS CONVERGENCE TOLERANCE

560.000 0.0 30 0.000100000

*** RESULTS ***OUTLET TEMPERATURE F455.16OUTLET PRESSURE PSIA560.00VAPOR FRACTION0.92669

V-L PHASE EQUILIBRIUM :

| COMP | F(I) | X(I) | Y(I) | K(I) | |
|-------|---------|---------|------|------|--------|
| WATER | 0.37158 | 0.34956 | 0.37 | 332 | 1.0680 |

| HYDROGEN 0.46522 0.4442 | 9E-02 0.50167 112.92 |
|---------------------------------|-----------------------------------|
| PG 0.15630 0.64158 (| 0.11791 0.18378 |
| N-PROP 0.21285E-02 0.18346 | E-02 0.21517E-02 1.1728 |
| ISO-PROP 0.46933E-02 0.2530 | 8E-02 0.48644E-02 1.9221 |
| METHA-01 0.74146E-04 0.491 | 10E-04 0.76126E-04 1.5501 |
| ASPEN PLUS PLAT: WIN32 VER: 2 | 4.0 04/11/2011 PAGE 27 |
| | |
| U-O-S BLOCK SECTIO | ON |
| BLOCK: F-102 MODEL: FLASH2 | |
| INLET STREAM: S-124 | |
| OUTLET VAPOR STREAM S-126 | |
| OUTLET LIQUID STREAM: S-125 | |
| PROPERTY OPTION SET: PENG-RO | B STANDARD PR EQUATION OF STATE |
| | |
| *** MASS AND ENERGY | BALANCE *** |
| IN OUT | RELATIVE DIFF |
| TOTAL BALANCE | |
| $MOLE(LBMOL/HR) \qquad 985.192$ | 985 192 -0.115396E-15 |
| MASS(LB/HR) 16879.8 | 16879.8 -0.215523E-15 |
| ENTHALPY(BTU/HR) -0.585529 | 9E+08 -0.585529E+08 -0.872072E-07 |
| | |
| *** INPUT DATA *** | |
| TWO PHASE PO FLASH | |
| PRESSURE DROP PSI | 0.0 |
| SPECIFIED HEAT DUTY BTU/HR | 0.0 |
| MAXIMUM NO. ITERATIONS | 30 |
| CONVERGENCE TOLERANCE | 0.000100000 |
| | |
| *** RESULTS *** | |
| OUTLET TEMPERATURE F | 429.05 |
| OUTLET PRESSURE PSIA | 559.14 |
| VAPOR FRACTION | 0.90185 |
| | |
| | |
| | |
| V-L PHASE EQUILIBRIUM : | |
| | |
| COMP F(I) X(I) Y | (I) K(I) |
| WATER 0.37332 0.43594 | 0.36651 0.84074 |
| HYDROGEN 0.50167 0.3687 | 0E-02 0.55587 150.77 |
| | 202055 01 0 12552 |

| PG | 0.11/91 0.555 | 54 0.70286 | E-01 0.12652 | |
|----------|---------------|-------------|--------------|--------|
| N-PROP | 0.21517E-02 | 0.20419E-02 | 0.21637E-02 | 1.0596 |
| ISO-PROP | 0.48644E-02 | 0.27370E-02 | 0.50959E-02 | 1.8619 |
| METHA-01 | 0.76126E-04 | 0.60607E-04 | 0.77815E-04 | 1.2839 |

BLOCK: FSPLIT MODEL: FSPLIT

INLET STREAM: S-110 OUTLET STREAMS: S-113 S-111 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 28

BLOCK: FSPLIT MODEL: FSPLIT (CONTINUED) *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 762.564 762.564 0.00000 MASS(LB/HR) 13984.8 13984.8 -0.130069E-15 ENTHALPY(BTU/HR) -0.939180E+08 -0.939180E+08 0.158661E-15 *** INPUT DATA *** FRACTION OF FLOW STRM=S-113 FRAC= 0.34000 *** RESULTS *** STREAM= S-113 SPLIT= 0.34000 KEY= 0 STREAM-ORDER= 1 0.66000 0 S-111 2 BLOCK: H-101 MODEL: HEATER -----INLET STREAM: S-107 OUTLET STREAM: S-108 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 1138.06 1138.06 0.486765E-08 MASS(LB/HR) 33229.6 33229.6 0.908441E-08 ENTHALPY(BTU/HR) -0.164137E+09 -0.161726E+09 -0.146863E-01 *** INPUT DATA *** TWO PHASE TP FLASH SPECIFIED TEMPERATURE F 242.000 PRESSURE DROP PSI 5.00000 MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 29 **U-O-S BLOCK SECTION** BLOCK: H-101 MODEL: HEATER (CONTINUED) *** RESULTS *** OUTLET TEMPERATURE F 242.00 OUTLET PRESSURE PSIA 36.816 HEAT DUTY BTU/HR 0.24106E+07 0.0000 OUTLET VAPOR FRACTION PRESSURE-DROP CORRELATION PARAMETER 0.20788E+07

V-L PHASE EQUILIBRIUM :

COMP F(I) X(I) Y(I)K(I)0.37835E-03 0.17172E-02 GLYCEROL 0.14802 0.14802 0.83645 WATER 0.83645 0.92100 0.73970 METHA-01 0.15525E-01 0.15525E-01 0.78625E-01 3.4022 BLOCK: HX-101 MODEL: HEATX _____ THIS BLOCK RUNS WITH ASPEN EDR 25.0 WITH ADVANCED METHOD FOR SHELL&TUBE HOT SIDE: -----INLET STREAM: S-109 OUTLET STREAM: S-110 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE COLD SIDE: _____ INLET STREAM: S-106 OUTLET STREAM: S-107 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 1900.71 1900.63 0.427128E-MASS(LB/HR) 47215.8 47214.4 0.310346E-04 0.427128E-04 ENTHALPY(BTU/HR) -0.258065E+09 -0.258055E+09 -0.387716E-04 *** INPUT DATA *** FLASH SPECS FOR HOT SIDE: TWO PHASE FLASH MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 30 **U-O-S BLOCK SECTION** BLOCK: HX-101 MODEL: HEATX (CONTINUED) FLASH SPECS FOR COLD SIDE: TWO PHASE FLASH MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 SHELL&TUBE INPUT FILE NAME EDR HX101.EDR SHELL&TUBE PROGRAM MODE DESIGN HEAT CURVE GENERATION

HEAT CURVE GENERATION HOT HEAT CURVE GENERATED BY COLD HEAT CURVE GENERATED BY

ASPEN PLUS ASPEN PLUS

| STREAMS: | | |
|---------------------------------------------------------------------|-------------|------------------------------------------------------------|
| S-109> T= 2.2138D+02 P= 2.0000D+01 V= 0.0000D+00 | HOT (SHELL) | > S-110 T= 8.0000D+01 P= 1.9741D+01 V= 0.0000D+00 |
| S-107 < T= 1.5183D+02 P= 4.1816D+01 V= 0.0000D+00 | COLD (TUBE) | < S-106 T= 6.9907D+01 P= 4.2000D+01 V= 0.0000D+00 |
| UNIT RESULTS: | | |

| CALCULATED HEAT | DUTY 1 | BTU/HR | 2136339.45 | 528 |
|---------------------|-------------|------------|------------|---------|
| CALCULATED (REQU | IRED) AREA | SQFT | 2043.42 | 209 |
| ACTUAL EXCHANGE | R AREA | SQFT | 2052.035 | 0 |
| % OVER (UNDER) DE | SIGN (DIRTY |) | 0.4216 | |
| % OVER (UNDER) DE | SIGN (CLEA) | (V | 0.4216 | |
| AVERAGE COEFFICIE | ENT (DIRTY) | BTU/HR-S | QFT-R | 33.9741 |
| AVERAGE COEFFICIE | ENT (CLEAN) | BTU/HR-S | SQFT-R | 33.9741 |
| UA (DIRTY) | BTU/HR-R | 69423 | 3.3987 | |
| LMTD (CORRECTED) | F | 30. | 7726 | |
| LMTD CORRECTION | FACTOR | | 0.9989 | |
| NUMBER OF SHELLS | IN SERIES | | 1 | |
| NUMBER OF SHELLS | IN PARALLE | EL | 1 | |
| ASPEN PLUS PLAT: WI | IN32 VER: | 24.0 | 04/11/2011 | PAGE 31 |

BLOCK: HX-101 MODEL: HEATX (CONTINUED)

| SHELLSIDE RESULTS: | | | |
|-------------------------------|-------------|------------|----------|
| AVERAGE WALL TEMPERAT | URE F | 122.82 | 254 |
| CLEAN PRESSURE DROP | PSI | 0.2593 | |
| WINDOW PRESSURE DROP | PSI | 0.0007 | |
| PRESSURE DROP IN ENDS | PSI | 0.0007 | |
| CROSSFLOW PRESSURE DRO | P PSI | 0.0029 | 1 |
| BULK FILM COEFFICIENT | BTU/HR-S | QFT-R 13 | 1.2348 |
| WALL FILM COEFFICIENT | BTU/HR-S | SQFT-R 13 | 1.2348 |
| THERMAL RESISTANCE | HR-SQFT- | R/BTU 0. | 0076 |
| MAXIMUM FOULING RESISTA | ANCE HR- | SQFT-R/BTU | 0.000062 |
| FOULING RESISTANCE | HR-SQFT-R | R/BTU 0.0 | 000 |
| CROSSFLOW VELOCITY | FT/SEC | 0.0885 | |
| WINDOW VELOCITY | FT/SEC | 0.1210 | |
| MIDPOINT VELOCITY | FT/SEC | 0.1047 | |
| SHELL ENTRANCE RHOV^2 | LB/FT-S | QSEC 95. | .0789 |
| SHELL EXIT RHOV^2 | LB/FT-SQSEC | C 87.2858 | |
| BUNDLE ENTRANCE RHOV^2 | LB/FT- | SQSEC (| 0.4808 |
| BUNDLE EXIT RHOV^2 | LB/FT-SQSI | EC 0.441 | 4 |
| FOULING % OF OVERALL RE | SISTANCE | 0.0 | 000 |

| FILM % OF OVERALL RE | ESISTANCE | 25.888 | 30 |
|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|------------------------------------------------------------------------|----------------------------------------------|
| FRICTIONAL PRESSURE | DROP PSI | 0.2593 | |
| TUBESIDE RESULTS: AVERAGE WALL TEMPE CLEAN PRESSURE DROF BULK FILM COEFFICIEN WALL FILM COEFFICIEN THERMAL RESISTANCE MAXIMUM FOULING RE FOULING RESISTANCE | RATURE F PSI T BTU/HR-SQ T BTU/HR-SQ HR-SQFT-R SISTANCE HR-S HR-SQFT-R/ | 115.0 0.1840 QFT-R 4 QFT-R 4 /BTU 0 QFT-R/BTU BTU 0(| 198 5.3992 6.3992 .0216 0.000062 |
| INPUT VELOCITY | FT/SEC | 0.1329 | |
| OUTLET VELOCITY | FT/SEC | 0.1384 | |
| MIDPOINT VELOCITY | FT/SEC | 0.1384 | |
| FOULING % OF OVERAL | L RESISTANCE | 0.0 | 0000 |
| FILM % OF OVERALL RE | ESISTANCE | 73.221 | .4 |
| FRICTIONAL PRESSURE | DROP PSI | 0.1840 | |
| BLOCK: HX-102 MODEL: | HEATX | | |
| THIS BLOCK RUNS WITH HOT SIDE: | ASPEN EDR 25.0 W | ITH ADVANCE | ED METHOD FOR SHELL&TUBE |
| INLET STREAM: S-12 OUTLET STREAM: S- PROPERTY OPTION SET: COLD SIDE: | 23 124 PENG-ROB STAND | ARD PR EQUA | ATION OF STATE |
| INLET STREAM: S-11 OUTLET STREAM: S- PROPERTY OPTION SET: ASPEN PLUS PLAT: WIN3 | .5 116 PENG-ROB STAND 2 VER: 24.0 | ARD PR EQUA 04/11/2011 | ATION OF STATE PAGE 32 |
| U-O-S BLO | CK SECTION | | |
| BLOCK: HX-102 MODEL: | HEATX (CONTINUE | D) | |
| ************************************** | ************************************** | ***** | ***** |
| * A POTENTIAL TUBE * | VIBRATIEON PROB | LEM IS INDICA | ATED * |
| ****** | ******** | ****** | **** |
| *** MASS AND IN | ENERGY BALANCE OUT RELATIV | 5 *** 'E DIFF. | |
| TOTAL BALANCE | | · | |
| MOLE(LBMOL/HR) | 1360.69 1360.6 | 9 0.00000 | |
| MASS(LB/HR) 30 | 6124.6 36124.6 | 0.00000 | |
| ENTHALPY(BTU/HR) | -0.127173E+09 -0. | 127173E+09 0 | .234345E-15 |

*** INPUT DATA ***

FLASH SPECS FOR HOT SIDE: TWO PHASE FLASH MAXIMUM NO. ITERATIONS CONVERGENCE TOLERANCE

FLASH SPECS FOR COLD SIDE: TWO PHASE FLASH MAXIMUM NO. ITERATIONS CONVERGENCE TOLERANCE

SHELL&TUBE INPUT FILE NAME SHELL&TUBE PROGRAM MODE

HEAT CURVE GENERATION HOT HEAT CURVE GENERATED BY COLD HEAT CURVE GENERATED BY

*** OVERALL RESULTS ***

STREAMS:

| S-123> | HOT (SHELL) | > S-124 |
|---------------|-------------|------------------|
| T= 4.5516D+02 | | T= 4.2905D+02 |
| P= 5.6000D+02 | | P= 5.5914D+02 |
| V= 1.0000D+00 | | V= 9.0185D-01 |
| | | |
| S-116 < | COLD (TUBE) | < S-115 |
| T= 4.1000D+02 | | T= 2.5213D+02 |
| P= 5.8104D+02 | | P= 5.8700D+02 |
| V= 0.0000D+00 | | V = 0.0000D + 00 |
| | | |

ASPEN PLUS PLAT: WIN32 VER: 24.0

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U-O-S BLOCK SECTION

BLOCK: HX-102 MODEL: HEATX (CONTINUED)

| UNIT RESULTS: | | | | |
|--------------------|--------------|---------|----------|----------|
| CALCULATED HEAT | DUTY B | TU/HR | 1990193. | 7789 |
| CALCULATED (REQU | JIRED) AREA | SQFT | 66.0 | 988 |
| ACTUAL EXCHANGE | R AREA | SQFT | 72.845 | 56 |
| % OVER (UNDER) DE | SIGN (DIRTY) |) | 10.2072 | |
| % OVER (UNDER) DE | SIGN (CLEAN | () | 10.2072 | |
| AVERAGE COEFFICIE | ENT (DIRTY) | BTU/HR- | SQFT-R | 344.7762 |
| AVERAGE COEFFICIE | ENT (CLEAN) | BTU/HR | -SQFT-R | 344.7762 |
| UA (DIRTY) | BTU/HR-R | 2278 | 39.2816 | |
| LMTD (CORRECTED) | F | 87 | 7.3303 | |
| LMTD CORRECTION | FACTOR | | 0.9052 | |
| NUMBER OF SHELLS | IN SERIES | | 1 | |
| NUMBER OF SHELLS | IN PARALLE | L | 1 | |
| | | | | |
| SHELLSIDE RESULTS: | | | | |
| AVERAGE WALL TEN | MPERATURE | F | 441 | .7477 |

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30

0.000100000

30 0.000100000

> EDR_HX102.EDR DESIGN

ASPEN PLUS ASPEN PLUS

| CLEAN PRESSURE DROP | PSI | 0.8597 | |
|---------------------------|-------------|----------------|----------|
| WINDOW PRESSURE DROP | PSI | 0.2903 | |
| PRESSURE DROP IN ENDS | PSI | 0.2903 | |
| CROSSFLOW PRESSURE DR | OP PSI | 0.2178 | |
| BULK FILM COEFFICIENT | BTU/HR-SO | QFT-R 754.86 | 509 |
| WALL FILM COEFFICIENT | BTU/HR-S | QFT-R 754.80 | 509 |
| THERMAL RESISTANCE | HR-SQFT-F | R/BTU 0.001 | 3 |
| MAXIMUM FOULING RESIS | TANCE HR-S | SQFT-R/BTU | 0.000148 |
| FOULING RESISTANCE | HR-SQFT-R/ | BTU 0.0000 |) |
| CROSSFLOW VELOCITY | FT/SEC | 17.0614 | |
| WINDOW VELOCITY | FT/SEC | 30.5421 | |
| MIDPOINT VELOCITY | FT/SEC | 23.8017 | |
| SHELL ENTRANCE RHOV^2 | LB/FT-SQ | SEC 332.01 | 35 |
| SHELL EXIT RHOV^2 | LB/FT-SQSEC | 296.9255 | |
| BUNDLE ENTRANCE RHOV | ^2 LB/FT-S | QSEC 246.7 | 540 |
| BUNDLE EXIT RHOV^2 | LB/FT-SQSE | C 220.2565 | |
| FOULING % OF OVERALL R | ESISTANCE | 0.0000 | |
| FILM % OF OVERALL RESIS | STANCE | 45.6741 | |
| FRICTIONAL PRESSURE DR | OP PSI | 0.8866 | |
| | | | |
| TUBESIDE RESULTS: | | | |
| AVERAGE WALL TEMPERA | TURE F | 397.6940 | |
| CLEAN PRESSURE DROP | PSI | 5.9635 | |
| BULK FILM COEFFICIENT | BTU/HR-SC | QFT-R 770.42 | .99 |
| WALL FILM COEFFICIENT | BTU/HR-S | QFT-R 770.42 | 299 |
| THERMAL RESISTANCE | HR-SQFT-F | R/BTU 0.001 | 3 |
| MAXIMUM FOULING RESIS | TANCE HR-S | SQFT-R/BTU | 0.000148 |
| FOULING RESISTANCE | HR-SQFT-R/ | BTU 0.0000 | |
| INPUT VELOCITY | FT/SEC | 4.9940 | |
| OUTLET VELOCITY | FT/SEC | 5.4123 | |
| MIDPOINT VELOCITY | FT/SEC | 5.4123 | |
| FOULING % OF OVERALL R | ESISTANCE | 0.0000 | |
| FILM % OF OVERALL RESIS | STANCE | 44.7511 | |
| FRICTIONAL PRESSURE DR | OP PSI | 5.9310 | |
| ASPEN PLUS PLAT: WIN32 | VER: 24.0 | 04/11/2011 PAG | GE 34 |
| | | | |

BLOCK: HX-103 MODEL: HEATX

PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE
* A POTENTIAL TUBE VIBRATIEON PROBLEM IS INDICATED * *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 1576.12 1576.12 0.00000 MASS(LB/HR) 13956.0 13956.0 0.00000 ENTHALPY(BTU/HR) -0.450774E+08 -0.450774E+08 -0.165284E-15 *** INPUT DATA *** FLASH SPECS FOR HOT SIDE: TWO PHASE FLASH MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 FLASH SPECS FOR COLD SIDE: TWO PHASE FLASH MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 SHELL&TUBE INPUT FILE NAME EDR HX103.EDR SHELL&TUBE PROGRAM MODE DESIGN HEAT CURVE GENERATION HOT HEAT CURVE GENERATED BY ASPEN PLUS COLD HEAT CURVE GENERATED BY ASPEN PLUS ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 35 **U-O-S BLOCK SECTION** BLOCK: HX-103 MODEL: HEATX (CONTINUED) *** OVERALL RESULTS *** STREAMS: S-126 ----> HOT (SHELL) |----> S-127 T= 4.2905D+02 | P= 5.5914D+02 | T = 4.0905D + 02| P= 5.5828D+02 V= 1.0000D+00 | | V= 9.3825D-01 S-120 <-----| COLD (TUBE) |<---- S-119

UNIT RESULTS:

T = 4.1649D + 02

P= 5.9891D+02 |

V= 1.0000D+00 |

*

T = 2.0241D + 02

V= 9.9431D-01

P = 6.0000D + 02

| CALCULATED HEAT DUTY | BTU/HR | 1132628.0668 |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------|---------------------|
| CALCULATED (REQUIRED) AREA | A SQFT | 156.3568 |
| ACTUAL EXCHANGER AREA | SQFT | 164.4385 |
| % OVER (UNDER) DESIGN (DIRTY | () | 5.1688 |
| % OVER (UNDER) DESIGN (CLEA | Ń) | 5.1688 |
| AVERAGE COEFFICIENT (DIRTY) | BTU/HR-SOF | Г-В 145.2866 |
| AVERAGE COEFFICIENT (CLEAN |) $BTU/HR-SOF$ | $T_{-R} = 145.2866$ |
| $\frac{1}{1} \frac{1}{1} \frac{1}$ | 22716 55 | 18 |
| I MTD (COPPECTED) = E | /0.850 | 2 2 |
| LWID CORRECTED) | 49.039 | 2 |
| LMID CORRECTION FACTOR | (| 1 |
| NUMBER OF SHELLS IN DADALL | | 1 |
| NUMBER OF SHELLS IN PARALL | EL | 1 |
| | | |
| SHELLSIDE RESULTS: | | 101 1 105 |
| AVERAGE WALL TEMPERATURE | | 421.1405 |
| CLEAN PRESSURE DROP | PSI | 0.8607 |
| WINDOW PRESSURE DROP | PSI | 0.3807 |
| PRESSURE DROP IN ENDS | PSI | 0.3807 |
| CROSSFLOW PRESSURE DROP | PSI | 0.1580 |
| BULK FILM COEFFICIENT | BTU/HR-SQFT-I | R 569.6667 |
| WALL FILM COEFFICIENT | BTU/HR-SQFT- | R 569.6667 |
| THERMAL RESISTANCE | HR-SQFT-R/BTU | J 0.0018 |
| MAXIMUM FOULING RESISTANC | E HR-SQFT | -R/BTU 0.000178 |
| FOULING RESISTANCE H | R-SOFT-R/BTU | 0.0000 |
| CROSSFLOW VELOCITY | FT/SEC | 13.7680 |
| WINDOW VELOCITY FT | T/SEC | 27.6842 |
| MIDPOINT VELOCITY FT | T/SEC | 20.7261 |
| SHELL ENTRANCE RHOV^2 | LB/FT-SOSEC | 125 6469 |
| SHELL EXIT RHOV^2 | FT-SOSEC | 466 0312 |
| BUNDI E ENTRANCE RHOV^2 | I B/FT-SOSE(| 7 36 9687 |
| BUNDLE EXIT RHOV^2 | B/FT_SOSEC | 53 8235 |
| FOULING % OF OVERALL RESIST | TANCE | 0,0000 |
| FILM % OF OVERALL RESISTANCE | TE | 25 5038 |
| EDICTIONAL DESSUE DOOD | | 25.5058 |
| A CDEN DI LIC DI AT. WIN22 VED. | PSI 0 | 0.0755 |
| ASPEN PLUS PLAT: WIN52 VER: | 24.0 0 | 4/11/2011 PAGE 30 |
| U-O-S BLOCK SECT | ION | |
| BLOCK: HX-103 MODEL: HEATX (| CONTINUED) | |
| becer. In 105 mobile. Herrin (| contracted) | |
| TUBESIDE RESUITS: | | |
| AVERAGE WALL TEMPERATURE | F | 405 7931 |
| CLEAN PRESSURE DROP | | 1 0853 |
| PLILK EILM COFFEICIENT | ι 51 ρτι /μρ ςλετ Ι | 2 - 206 1210 |
| WALL EILM COEFFICIENT | DIU/IIK-SQI'I-I | 200.1210 |
| | $DIU/\Pi K-SQFI-$ | R 200.1210 |
| THERMAL RESISTANCE | HK-SQFI-K/BIU | D (DTU 0.00170 |
| MAXIMUM FOULING RESISTANC | E HR-SQFT | -R/BTU 0.0001/8 |
| FOULING RESISTANCE H | R-SQFT-R/BTU | 0.0000 |
| INPUT VELOCITY FT/S | EC 40 | .3146 |
| OUTLET VELOCITY FT/ | SEC 5 | 3.5906 |
| MIDPOINT VELOCITY FT | T/SEC | 53.5906 |
| FOULING % OF OVERALL RESIST | TANCE | 0.0000 |
| FILM % OF OVERALL RESISTANC | CE | 70.4861 |
| FRICTIONAL PRESSURE DROP | PSI | 1 0606 |

BLOCK: M-101 MODEL: MIXER -----INLET STREAMS:S-131OUTLET STREAM:S-119 S-118 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 687.645 687.624 0.305083E-04 MASS(LB/HR) 1952.06 1951.94 0.584380E-04 ENTHALPY(BTU/HR) -0.182133E+07 -0.182118E+07 -0.861022E-04 *** INPUT DATA *** TWO PHASE FLASH MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 OUTLET PRESSURE: MINIMUM OF INLET STREAM PRESSURES BLOCK: M-102 MODEL: MIXER -----INLET STREAMS: S-101 S-102 S-112 OUTLET STREAM: S-103 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 37 **U-O-S BLOCK SECTION** BLOCK: M-102 MODEL: MIXER (CONTINUED) *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 1151.42 1151.42 0.00000 MASS(LB/HR) 34006.8 34006.8 0.00000 ENTHALPY(BTU/HR) -0.169033E+09 -0.169033E+09 0.176311E-15 *** INPUT DATA *** TWO PHASE FLASH MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 OUTLET PRESSURE PSIA 25.0000 BLOCK: M-103 MODEL: MIXER -----INLET STREAMS: S-122 S-129 S-125 OUTLET STREAM: S-132 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 547.323 547.323 0.00000

MASS(LB/HR) 19591.2 19591.2 0.371389E-15 ENTHALPY(BTU/HR) -0.781100E+08 -0.781100E+08 -0.381543E-15

*** INPUT DATA *** TWO PHASE FLASH MAXIMUM NO. ITERATIONS CONVERGENCE TOLERANCE PRESSURE DROP PSI

30 0.000100000 5.00000

BLOCK: P-100 MODEL: PUMP

INLET STREAM: S-114 OUTLET STREAM: S-115 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 38

U-O-S BLOCK SECTION

BLOCK: P-100 MODEL: PUMP (CONTINUED)

*** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 375.500 375.500 0.00000 MASS(LB/HR) 19244.9 -0.189036E-15 ENTHALPY(BTU/HR) -0.706840E+08 -0.706100E+08 -0.104774E-02

*** INPUT DATA ***OUTLET PRESSURE PSIA587.000DRIVER EFFICIENCY1.00000

FLASH SPECIFICATIONS:LIQUID PHASE CALCULATIONNO FLASH PERFORMEDMAXIMUM NUMBER OF ITERATIONS30TOLERANCE0.000100000

*** RESULTS *** VOLUMETRIC FLOW RATE CUFT/HR 267.014 PRESSURE CHANGE PSI 567.000 NPSH AVAILABLE FT-LBF/LB 0.0 FLUID POWER HP 11.0107 BRAKE POWER HP 29.1061 ELECTRICITY KW 21.7044 PUMP EFFICIENCY USED 0.37829 NET WORK REQUIRED HP 29.1061 HEAD DEVELOPED FT-LBF/LB 1,132.83

BLOCK: P-101 MODEL: PUMP

INLET STREAM: S-105 OUTLET STREAM: S-106 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** OUT RELATIVE DIFF. IN TOTAL BALANCE MOLE(LBMOL/HR) 1138.15 1138.15 0.00000 MASS(LB/HR) 33231.1 0.00000 ENTHALPY(BTU/HR) -0.166287E+09 -0.166283E+09 -0.249300E-04 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 39 **U-O-S BLOCK SECTION** BLOCK: P-101 MODEL: PUMP (CONTINUED) *** INPUT DATA *** OUTLET PRESSURE PSIA 42.0000 DRIVER EFFICIENCY 1.00000 FLASH SPECIFICATIONS: LIQUID PHASE CALCULATION NO FLASH PERFORMED MAXIMUM NUMBER OF ITERATIONS 30 TOLERANCE 0.000100000 *** RESULTS *** VOLUMETRIC FLOW RATE CUFT/HR 471.696 PRESSURE CHANGE PSI 22.0000 NPSH AVAILABLE FT-LBF/LB 40.1994 FLUID POWER HP 0.75471 BRAKE POWER HP 1.62926 ELECTRICITY KW 1.21494 PUMP EFFICIENCY USED 0.46322 NET WORK REQUIRED HP 1.62926 HEAD DEVELOPED FT-LBF/LB 44.9679 BLOCK: P-102 MODEL: PUMP -----INLET STREAM: S-111 OUTLET STREAM: S-112 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 503.292 503.292 0.00000

MASS(LB/HR) 9229.94 9229.94 0.00000 ENTHALPY(BTU/HR) -0.619858E+08 -0.619849E+08 -0.156051E-04

*** INPUT DATA *** OUTLET PRESSURE PSIA 30.0000 DRIVER EFFICIENCY 1.00000

FLASH SPECIFICATIONS: LIQUID PHASE CALCULATION NO FLASH PERFORMED MAXIMUM NUMBER OF ITERATIONS

30

ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 40

U-O-S BLOCK SECTION

BLOCK: P-102 MODEL: PUMP (CONTINUED)

| *** RESULTS *** | |
|------------------------------|---------|
| VOLUMETRIC FLOW RATE CUFT/HR | 150.641 |
| PRESSURE CHANGE PSI | 10.2593 |
| NPSH AVAILABLE FT-LBF/LB | 44.9718 |
| FLUID POWER HP | 0.11240 |
| BRAKE POWER HP | 0.38016 |
| ELECTRICITY KW | 0.28349 |
| PUMP EFFICIENCY USED | 0.29566 |
| NET WORK REQUIRED HP | 0.38016 |
| HEAD DEVELOPED FT-LBF/LB | 24.1115 |
| | |

BLOCK: R-101 MODEL: RSTOIC

INLET STREAMS: S-116 S-120 OUTLET STREAM: S-121 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT GENERATION RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 1063.12 1063.13 0.00000 -0.605872E-05 MASS(LB/HR) 21196.8 21196.9 -0.379189E-05 ENTHALPY(BTU/HR) -0.693083E+08 -0.693085E+08 0.196263E-05

*** INPUT DATA *** STOICHIOMETRY MATRIX:

REACTION # 1: SUBSTREAM MIXED :

GLYCEROL -1.00 WATER 1.00 HYDROGEN -1.00 PG 1.00

REACTION # 2: SUBSTREAM MIXED : GLYCEROL -2.00 WATER 4.00 HYDROGEN -4.00 N-PROP 1.00 ISO-PROP 1.00

REACTION CONVERSION SPECS: NUMBER= 2 REACTION # 1: SUBSTREAM:MIXED KEY COMP:GLYCEROL CONV FRAC: 0.9850 REACTION # 2: SUBSTREAM:MIXED KEY COMP:GLYCEROL CONV FRAC: 1.000

TWO PHASE PQ FLASH

PRESSURE DROP PSI 15.0000 SPECIFIED HEAT DUTY BTU/HR 0.0 MAXIMUM NO. ITERATIONS 30 CONVERGENCE TOLERANCE 0.000100000 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 41 **U-O-S BLOCK SECTION** BLOCK: R-101 MODEL: RSTOIC (CONTINUED) SERIES REACTIONS GENERATE COMBUSTION REACTIONS FOR FEED SPECIES NO *** RESULTS *** OUTLET TEMPERATURE F 455.84 OUTLET PRESSURE PSIA 566.04 VAPOR FRACTION 0.92623 **REACTION EXTENTS:** REACTION REACTION NUMBER EXTENT LBMOL/HR 1 165.93 2 1.2634 V-L PHASE EQUILIBRIUM : COMP F(I) $X(I) \qquad Y(I)$ K(I)WATER 0.37158 0.35104 0.37322 1.0632 0.46522 0.45085E-02 0.50191 HYDROGEN 111.32 PG 0.15630 0.64002 0.11778 0.18402 N-PROP 0.21284E-02 0.18411E-02 0.21513E-02 1.1684 ISO-PROP 0.46924E-02 0.25428E-02 0.48636E-02 1.9127 METHA-01 0.74146E-04 0.49353E-04 0.76121E-04 1.5424 BLOCK: S-101 MODEL: SEP -----INLET STREAM: S-103 OUTLET STREAMS: S-104 S-105 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE *** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE 1151.42 MOLE(LBMOL/HR) 1151.42 0.394945E-15 MASS(LB/HR) 34006.8 34006.8 0.213956E-15 ENTHALPY(BTU/HR) -0.169033E+09 -0.168638E+09 -0.233928E-02 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 42 **U-O-S BLOCK SECTION**

BLOCK: S-101 MODEL: SEP (CONTINUED)

| *** INPUT DATA *** | |
|--------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------|
| INLET PRESSURE DROP PSI | 5.00000 |
| FLASH SPECS FOR STREAM S-104 ONE PHASE TP FLASH SPECIFIED P PRESSURE DROP PSI MAXIMUM NO. ITERATIONS CONVERGENCE TOLERANCE | HASE IS SOLID 0.0 30 0.000100000 |
| FLASH SPECS FOR STREAM S-105 TWO PHASE TP FLASH SPECIFIED TEMPERATURE CHANGE H PRESSURE DROP PSI MAXIMUM NO. ITERATIONS CONVERGENCE TOLERANCE | F 15.0000 0.0 30 0.000100000 |
| FRACTION OF FEED SUBSTREAM= MIXED STREAM= S-104 CPT= SODIU-01 FR | RACTION= 1.00000 |
| *** RESULTS *** | |
| HEAT DUTY BTU/HR | 0.39542E+06 |
| COMPONENT = GLYCEROL STREAM SUBSTREAM SPLIT FRAC S-105 MIXED 1.00000 | TION |
| COMPONENT = WATER STREAM SUBSTREAM SPLIT FRAC S-105 MIXED 1.00000 | TION |
| COMPONENT = SODIU-01 STREAM SUBSTREAM SPLIT FRAC S-104 MIXED 1.00000 | TION |
| COMPONENT = METHA-01 STREAM SUBSTREAM SPLIT FRAC S-105 MIXED 1.00000 ASPEN PLUS PLAT: WIN32 VER: 24.0 | TION 04/11/2011 PAGE 43 |
| U-O-S BLOCK SECTION | |
| BLOCK: T-101 MODEL: COMPR | |
| INLET STREAM: S-117 | |

INLET STREAM:S-117OUTLET STREAM:S-118PROPERTY OPTION SET:PENG-ROBSTANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE ***

IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 171.838 171.838 0.00000 MASS(LB/HR) 346.404 346.404 0.00000 ENTHALPY(BTU/HR) -5244.32 -75785.5 0.930800

*** INPUT DATA ***

| 600.000 |
|---------|
| 0.80000 |
| 1.00000 |
| |

*** RESULTS ***

| | INDICATED | HORSEPO | WER RI | EQUIREME | NT H | Р | -2 | 27.7237 |
|---|------------|-------------------|--------|------------|-------|----------|------|---------|
| | BRAKE H | IORSEPOW | ER REQ | UIREMENT | Г НР | -1 | 27. | 7237 |
| | NET WORK | REQUIRED | | HP | -2 | 27.7237 | | |
| | POWER LOS | SSES | H | Р | 0.0 | | | |
| | ISENTROPIO | C HORSEPO | WER R | EQUIREME | ENT H | Р | -3 | 34.6546 |
| | CALCULAT | ED OUTLEI | TEMP | F | | 11.7339 | | |
| | ISENTROPIO | C TEMPERA | TURE | F | | -3.14840 | | |
| | EFFICIENCY | Y (POLYTR/ | ISENTR |) USED | | 0.800 | 00 | |
| | OUTLET VA | POR FRAC | TION | | | 1.00000 | | |
| | HEAD DEVI | ELOPED, | FT-LB | F/LB | -1 | 98,081. | | |
| | MECHANIC | AL EFFICIE | NCY US | SED | | 1.0000 | 00 | |
| | INLET HEAT | T CAPACIT | Y RATI | С | | 1.41692 | | |
| | INLET VOL | UMETRIC F | LOW R | ATE , CUFT | T/HR | 1,0 |)10 | .06 |
| | OUTLET VC | DLUMETRIC | FLOW | RATE, CUI | FT/HR | . 1 | 1,4′ | 79.73 |
| | INLET COM | IPRESSIBIL | ITY FA | CTOR | | 1.034 | 11 | |
| | OUTLET CC | MPRESSIB | LITY F | ACTOR | | 1.02 | 13 | 3 |
| | AV. ISENT. | VOL. EXPO | NENT | | | 1.45958 | | |
| | AV. ISENT. | TEMP EXPO | DNENT | | | 1.41031 | | |
| | AV. ACTUA | L VOL. EXF | ONENT | | | 1.33775 | | |
| | AV. ACTUA | L TEMP EX | PONEN | Т | | 1.29557 | | |
| A | SPEN PLUS | PLAT: WIN | N32 V | ER: 24.0 | | 04/11/20 | 11 | PAGE 44 |

U-O-S BLOCK SECTION

BLOCK: T-101 MODEL: COMPR (CONTINUED)

BLOCK: V-101 MODEL: FLASH2

INLET STREAM: S-128 OUTLET VAPOR STREAM: S-130 OUTLET LIQUID STREAM: S-129 PROPERTY OPTION SET: PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 888.498 888.498 0.00000 MASS(LB/HR) 12004.1 12004.1 0.226374E-10 ENTHALPY(BTU/HR) -0.518861E+08 -0.518861E+08 0.143595E-15

| *** INPUT DATA *** | |
|-------------------------|-------------|
| TWO PHASE TP FLASH | |
| SPECIFIED TEMPERATURE F | 225.000 |
| PRESSURE DROP PSI | 0.0 |
| MAXIMUM NO. ITERATIONS | 30 |
| CONVERGENCE TOLERANCE | 0.000100000 |
| | |
| *** RESULTS *** | |

| | REDUCTIO | |
|--------------|-----------|---------|
| OUTLET TEMPI | ERATURE F | 225.00 |
| OUTLET PRESS | URE PSIA | 553.28 |
| HEAT DUTY | BTU/HR | 0.0000 |
| VAPOR FRACTI | ION | 0.58054 |
| | | |

ASPEN PLUS PLAT: WIN32 VER: 24.0

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U-O-S BLOCK SECTION

BLOCK: V-101 MODEL: FLASH2 (CONTINUED)

V-L PHASE EQUILIBRIUM :

| COMP | F(I) | X(I) | Y(I) | K(I) | | |
|----------|-------------|----------|----------------|----------|------------|------|
| WATER | 0.36651 | 0.827 | '93 0.3 | 3113E-01 | 0.39995E | -01 |
| HYDROGEN | N 0.555 | 87 0. | 36804E-03 | 0.95724 | 2600. | 8 |
| PG | 0.70286E-01 | 0.1669 | 0.46 | 548E-03 | 0.27886E-0 |)2 |
| N-PROP | 0.21637E | -02 0.2 | 4765E-02 | 0.19376E | E-02 0.782 | 236 |
| ISO-PROP | 0.50959 | E-02 0.2 | 21508E-02 | 0.72239 | E-02 3.35 | 586 |
| METHA-01 | 0.77815 | 5E-04 0 | .15535E-03 | 3 0.2179 | 1E-04 0.1- | 4027 |

BLOCK: VALVE MODEL: VALVE

INLET STREAM:S-132OUTLET STREAM:S-133PROPERTY OPTION SET:PENG-ROB STANDARD PR EQUATION OF STATE

*** MASS AND ENERGY BALANCE *** IN OUT RELATIVE DIFF. TOTAL BALANCE MOLE(LBMOL/HR) 547.323 547.323 0.00000 MASS(LB/HR) 19591.2 19591.2 0.00000 ENTHALPY(BTU/HR) -0.781100E+08 -0.781100E+08 0.00000

*** INPUT DATA ***

VALVE OUTLET PRESSURE PSIA VALVE FLOW COEF CALC. NO

FLASH SPECIFICATIONS: NPHASE 2 MAX NUMBER OF ITERATIONS CONVERGENCE TOLERANCE

30 0.000100000

30.0000

VALVE PRESSURE DROP PSI 518.280 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 46 STREAM SECTION S-101 S-102 S-103 S-104 S-105 _____ STREAM ID S-103 S-101 S-102 S-104 S-105 FROM : M-102 S-101 S-101 ----____ P-101 TO : M-102 M-102 S-101 ____ SUBSTREAM: MIXED PHASE: LIQUID LIQUID LIQUID SOLID LIQUID COMPONENTS: LBMOL/HR GLYCEROL 168.4569 0.0 168.4574 0.0 168.4574 ACETOL 0.0 0.0 0.0 0.0 0.0 161.4670 298.8769 952.0188 0.0 WATER 952.0188 **HYDROGEN** 0.0 0.0 0.0 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 N-PROP 0.0 0.0 0.0 0.0 0.0 ETHYL-01 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 **ISO-PROP** 13.2729 13.2729 SODIU-01 13.2729 0.0 0.0 METHA-01 6.0522 0.0 17.6689 0.0 17.6689 COMPONENTS: MOLE FRAC 0.4823 0.0 **GLYCEROL** 0.1463 0.0 0.1480 0.0 0.0 0.0 ACETOL 0.0 0.0 WATER 0.4623 1.0000 0.8268 0.0 0.8365 0.0 0.0 HYDROGEN 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 N-PROP 0.0 0.0 ETHYL-01 0.0 0.0 0.0 0.0 0.0 **ISO-PROP** 0.0 0.0 0.0 0.0 0.0 3.8004-02 0.0 1.1527-02 1.0000 0.0 SODIU-01 METHA-01 1.7329-02 0.0 1.5345-02 0.0 1.5524-02 COMPONENTS: MASS FRAC GLYCEROL 0.8000 0.0 0.4562 0.0 0.4669 ACETOL 0.0 0.0 0.0 0.0 0.0 WATER 0.1500 1.0000 0.5043 0.0 0.5161 HYDROGEN 0.0 0.0 0.0 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 N-PROP 0.0 0.0 0.0 0.0 0.0 ETHYL-01 0.0 0.0 0.0 0.0 0.0 **ISO-PROP** 0.0 0.0 0.0 0.0 0.0 SODIU-01 4.0000-02 0.0 2.2810-02 1.0000 0.0 1.0000-02 METHA-01 0.0 1.6648-02 0.0 1.7037-02 TOTAL FLOW: LBMOL/HR 349.2489 298.8769 1151.4180 13.2729 1138.1451 1.9392+04 5384.3506 3.4007+04 775.6994 3.3231+04 LB/HR CUFT/HR 242.1154 86.4478 470.7215 5.7432 471.6956 STATE VARIABLES:

TEMP F 70.0000 70.0000 54.8014 54.8182 69.8182 PRES PSIA 30.0000 30.0000 25.0000 20.0000 20.0000 VFRAC 0.0 0.0 0.0 0.0 0.0 **LFRAC** 1.0000 1.0000 1.0000 0.0 1.0000 SFRAC 0.0 0.0 0.0 1.0000 0.0 ENTHALPY: BTU/LBMOL -2.0053+05 -1.2384+05 -1.4680+05 -1.7708+05 -1.4610+05 BTU/LB -3611.4585 -6874.1822 -4970.5698 -3029.9052 -5003.9710 -7.0035+07 -3.7013+07 -1.6903+08 -2.3503+06 -1.6629+08 BTU/HR ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 47

STREAM SECTION

S-101 S-102 S-103 S-104 S-105 (CONTINUED)

STREAM ID S-101 S-102 S-103 S-104 S-105

ENTROPY:

| BTU/LBMOL-R | -86.4419 -40.3582 -54.1619 -22.1726 -53.7260 |
|--------------|----------------------------------------------|
| BTU/LB-R | -1.5568 -2.2402 -1.8338 -0.3794 -1.8401 |
| DENSITY: | |
| LBMOL/CUFT | 1.4425 3.4573 2.4461 2.3111 2.4129 |
| LB/CUFT | 80.0960 62.2844 72.2439 135.0641 70.4503 |
| AVG MW | 55.5263 18.0153 29.5347 58.4425 29.1976 |
| ASPEN PLUS P | PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 48 |

STREAM SECTION

S-106 S-107 S-108 S-109 S-110

| STREAM ID | S-106 | 5 S-107 | S-108 | S-109 | S-110 |
|-----------|--------|---------|-------|--------|--------|
| FROM : | P-101 | HX-101 | H-101 | D-101 | HX-101 |
| TO : | HX-101 | H-101 | D-101 | HX-101 | FSPLIT |

MAX CONV. ERROR: 8.5077-05 0.0 8.1349-07 0.0 0.0 SUBSTREAM: MIXED LIQUID LIQUID LIQUID LIQUID PHASE: LIQUID COMPONENTS: LBMOL/HR 168.4574 168.4574 168.4574 8.4221-04 8.4221-04 GLYCEROL ACETOL 0.0 0.0 0.0 0.0 0.0 WATER 952.0188 951.9378 951.9378 744.9620 744.9620 0.0 HYDROGEN 0.0 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 N-PROP ETHYL-01 0.0 0.0 0.0 0.0 0.0 **ISO-PROP** 0.0 0.0 0.0 0.0 0.0 SODIU-01 0.0 0.0 0.0 0.0 0.0 METHA-01 17.6689 17.6687 17.6687 17.6011 17.6011 COMPONENTS: MOLE FRAC GLYCEROL 0.1480 0.1480 0.1480 1.1044-06 1.1044-06 ACETOL 0.0 0.0 0.0 0.0 0.0 WATER 0.8365 0.8365 0.8365 0.9769 0.9769

HYDROGEN 0.0 0.0 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 0.0 N-PROP 0.0 0.0 0.0 0.0 0.0 ETHYL-01 0.0 0.0 0.0 0.0 0.0 **ISO-PROP** 0.0 0.0 0.0 0.0 0.0 SODIU-01 0.0 0.0 0.0 0.0 0.0 1.5524-02 1.5525-02 1.5525-02 2.3082-02 2.3082-02 METHA-01 COMPONENTS: MASS FRAC 0.4669 0.4669 0.4669 5.5463-06 5.5463-06 GLYCEROL ACETOL 0.0 0.0 0.0 0.0 0.0 WATER 0.5161 0.5161 0.5161 0.9597 0.9597 0.0 **HYDROGEN** 0.0 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 0.0 N-PROP 0.0 0.0 0.0 0.0 0.0 ETHYL-01 0.0 0.0 0.0 0.0 0.0 **ISO-PROP** 0.0 0.0 0.0 0.0 0.0 SODIU-01 0.0 0.0 0.0 0.0 0.0 METHA-01 1.7037-02 1.7037-02 1.7037-02 4.0328-02 4.0328-02 TOTAL FLOW: LBMOL/HR 1138.1451 1138.0639 1138.0639 762.5639 762.5639 3.3231+04 3.3230+04 3.3230+04 1.3985+04 1.3985+04 LB/HR 471.7154 491.1869 515.9014 248.6216 228.2434 CUFT/HR STATE VARIABLES: TEMP F 69.9067 151.8282 242.0000 221.3804 80.0000 PRES PSIA 42.0000 41.8160 36.8160 20.0000 19.7407 **VFRAC** 0.0 0.0 0.0 0.0 0.0 LFRAC 1.0000 1.0000 1.0000 1.0000 1.0000 **SFRAC** 0.0 0.0 0.0 0.0 0.0 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 49

STREAM SECTION

S-106 S-107 S-108 S-109 S-110 (CONTINUED)

STREAM ID S-106 S-107 S-108 S-109 S-110

ENTHALPY:

| BTU/LBMOL | -1.4610+05 -1.4422+05 -1.4211+05 -1.2036+05 -1.2316+05 |
|--------------|--------------------------------------------------------|
| BTU/LB | -5003.8462 -4939.4756 -4866.9332 -6562.9766 -6715.7386 |
| BTU/HR | -1.6628+08 -1.6414+08 -1.6173+08 -9.1782+07 -9.3918+07 |
| ENTROPY: | |
| BTU/LBMOL-F | R -53.7228 -50.4286 -47.1981 -35.6303 -40.2390 |
| BTU/LB-R | -1.8400 -1.7271 -1.6165 -1.9429 -2.1942 |
| DENSITY: | |
| LBMOL/CUFT | 2.4128 2.3170 2.2060 3.0672 3.3410 |
| LB/CUFT | 70.4473 67.6517 64.4108 56.2492 61.2712 |
| AVG MW | 29.1976 29.1984 29.1984 18.3391 18.3391 |
| ASPEN PLUS F | PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 50 |

STREAM SECTION

S-111 S-112 S-113 S-114 S-115

STREAM ID S-112 S-113 S-114 S-111 S-115 FROM : FSPLIT P-102 FSPLIT D-101 P-100 TO : P-102 M-102 P-100 HX-102 ----SUBSTREAM: MIXED PHASE: LIQUID LIQUID LIQUID LIQUID LIQUID COMPONENTS: LBMOL/HR GLYCEROL 5.5586-04 5.5586-04 2.8635-04 168.4566 168.4566 ACETOL 0.0 0.0 0.0 0.0 0.0 WATER 491.6749 491.6749 253.2871 206.9758 206.9758 HYDROGEN 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 PG N-PROP 0.0 0.0 0.0 0.0 0.0 ETHYL-01 0.0 0.0 0.0 0.0 0.0**ISO-PROP** 0.0 0.0 0.0 0.0 0.0 SODIU-01 0.0 0.0 0.0 0.0 0.0 11.6167 11.6167 5.9844 6.7587-02 6.7587-02 METHA-01 COMPONENTS: MOLE FRAC 1.1044-06 1.1044-06 1.1044-06 0.4486 GLYCEROL 0.4486 ACETOL 0.0 0.0 0.0 0.0 0.0 0.9769 0.9769 0.9769 0.5512 0.5512 WATER HYDROGEN 0.0 0.0 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 0.0 N-PROP 0.0 0.0 0.0 0.0 0.0 0.0 0.0 ETHYL-01 0.0 0.0 0.0 **ISO-PROP** 0.0 0.0 0.0 0.0 0.0 SODIU-01 0.0 0.0 0.0 0.0 0.0 METHA-01 2.3082-02 2.3082-02 2.3082-02 1.7999-04 1.7999-04 COMPONENTS: MASS FRAC 5.5463-06 5.5463-06 5.5463-06 GLYCEROL 0.8061 0.8061 ACETOL 0.0 0.0 0.0 0.0 0.0 0.9597 0.9597 WATER 0.9597 0.1938 0.1938 HYDROGEN 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 PG 0.0 0.0 0.0 0.0 N-PROP 0.0 0.0 0.0 0.0 0.0 0.0 ETHYL-01 0.0 **ISO-PROP** 0.0 0.0 0.0 0.0 SODIU-01 0.0 0.0 0.0 0.0 0.0 4.0328-02 4.0328-02 4.0328-02 1.1253-04 1.1253-04 METHA-01 TOTAL FLOW: 503.2922 503.2922 259.2717 375.5000 375.5000 LBMOL/HR 9229.9377 9229.9377 4754.8164 1.9245+04 1.9245+04 LB/HR CUFT/HR 150.6406 150.6465 77.6027 267.0142 267.5415 STATE VARIABLES: TEMP F 80.0000 80.0714 80.0000 247.8897 252.1323 19.7407 PRES PSIA 30.0000 19.7407 20.0000 587.0000 **VFRAC** 0.0 0.0 0.0 0.0 0.0 LFRAC 1.0000 1.0000 1.0000 1.0000 1.0000 SFRAC 0.0 0.00.0 0.0 0.0 ENTHALPY: BTU/LBMOL -1.2316+05 -1.2316+05 -1.2316+05 -1.8824+05 -1.8804+05BTU/LB -6715.7386 -6715.6338 -6715.7386 -3672.8791 -3669.0308 **BTU/HR** -6.1986+07 -6.1985+07 -3.1932+07 -7.0684+07 -7.0610+07 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 51

STREAM SECTION

S-111 S-112 S-113 S-114 S-115 (CONTINUED) STREAM ID S-111 S-112 S-113 S-114 S-115 ENTROPY: -40.2390 -40.2366 -40.2390 -74.0968 -73.9371 BTU/LBMOL-R BTU/LB-R -2.1942 -2.1940 -2.1942 -1.4458 -1.4426 DENSITY: 3.3410 3.3409 3.3410 1.4063 1.4035 LBMOL/CUFT 61.2712 61.2688 61.2712 72.0743 71.9322 LB/CUFT 18.3391 18.3391 18.3391 51.2513 51.2513 AVG MW ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 52 STREAM SECTION S-116 S-117 S-118 S-119 S-120 _____ S-116 S-117 S-118 S-119 S-120 STREAM ID FROM : HX-102 ---- T-101 M-101 HX-103 T-101 M-101 TO : R-101 HX-103 R-101 MAX CONV. ERROR: 0.0 0.0 0.0 2.5376-04 0.0 SUBSTREAM: MIXED PHASE: LIQUID VAPOR VAPOR MIXED VAPOR COMPONENTS: LBMOL/HR GLYCEROL 168.4566 0.0 0.0 0.0 0.0 0.0 0.0 0.0 ACETOL 0.0 0.0 WATER 206.9758 0.0 0.0 17.0791 17.0791 0.0 171.8376 171.8376 665.5687 665.5687 HYDROGEN 0.0 0.0 0.2401 0.2401 PG 0.0 N-PROP 0.0 0.0 0.0 0.9993 0.9993 0.0 0.0 0.0 ETHYL-01 0.0 0.0 ISO-PROP 0.0 0.0 0.0 3.7252 3.7252 0.0 0.0 SODIU-01 0.0 0.0 0.0 METHA-01 6.7587-02 0.0 0.0 1.1240-02 1.1240-02 COMPONENTS: MOLE FRAC GLYCEROL 0.4486 0.0 0.0 0.0 0.0 ACETOL 0.0 0.0 0.0 0.0 0.0 0.0 2.4838-02 2.4838-02 WATER 0.5512 0.0 **HYDROGEN** 0.0 1.0000 1.0000 0.9679 0.9679 PG 0.0 0.0 3.4915-04 3.4915-04 0.0 N-PROP 0.0 0.0 0.0 1.4533-03 1.4533-03 ETHYL-01 0.0 0.0 0.0 0.0 0.0 5.4175-03 5.4175-03 ISO-PROP 0.0 0.0 0.0 0.0 0.0 0.0 SODIU-01 0.0 0.0 METHA-01 1.7999-04 0.0 0.0 1.6346-05 1.6346-05 COMPONENTS: MASS FRAC 0.0 GLYCEROL 0.8061 0.0 0.0 0.0 ACETOL 0.0 0.0 0.0 0.0 0.0

WATER 0.1938 0.0 0.0 0.1576 0.1576 1.0000 1.0000 0.6874 0.6874 HYDROGEN 0.0 PG 0.0 0.0 0.0 9.3594-03 9.3594-03 0.0 3.0767-02 3.0767-02 N-PROP 0.0 0.0 ETHYL-01 0.0 0.0 0.0 0.0 0.0 0.1147 0.1147 0.0 **ISO-PROP** 0.0 0.0 SODIU-01 0.0 0.0 0.0 0.0 0.0 METHA-01 1.1253-04 0.0 0.0 1.8451-04 1.8451-04 TOTAL FLOW: LBMOL/HR 375.5000 171.8376 171.8376 687.6236 687.6236 LB/HR 1.9245+04 346.4039 346.4039 1951.9440 1951.9440 289.9531 1010.0551 1479.7311 8233.9153 1.0946+04 CUFT/HR STATE VARIABLES: TEMP F 410.0000 70.0000 11.7339 202.4134 416.4896 PRES PSIA 581.0365 1000.0000 600.0000 600.0000 598.9147 1.0000 VFRAC 0.0 1.0000 0.9943 1.0000 LFRAC 1.0000 0.0 0.0 5.6867-03 0.0 SFRAC 0.0 0.0 0.0 0.0 0.0 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 53

STREAM SECTION

S-116 S-117 S-118 S-119 S-120 (CONTINUED)

STREAM ID S-116 S-117 S-118 S-119 S-120

ENTHALPY:

BTU/LBMOL -1.8274+05 -30.5191 -441.0296 -2648.5083 -1001.3455-3565.6165 -15.1393 -218.7777 -933.0067 -352.7503 BTU/LB BTU/HR -6.8620+07 -5244.3222 -7.5785+04 -1.8212+06 -6.8855+05 ENTROPY: -67.2210 -8.5047 -8.2835 -6.5588 -4.3807 BTU/LBMOL-R BTU/LB-R -1.3116 -4.2188 -4.1091 -2.3105 -1.5432 DENSITY: 1.2950 0.1701 0.1161 8.3511-02 6.2821-02 LBMOL/CUFT LB/CUFT 66.3723 0.3430 0.2341 0.2371 0.1783 51.2513 2.0159 2.0159 2.8387 2.8387 AVG MW ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 54

STREAM SECTION

S-121 S-122 S-123 S-124 S-125

| STREAM ID | S-1 | 21 S-12 | 22 S-123 | S-124 | S-125 |
|-----------|--------------|---------|----------|--------|-------|
| FROM : | R-101 | F-101 | F-101 | HX-102 | F-102 |
| TO : | F-101 | M-103 | HX-102 | F-102 | M-103 |

MAX CONV. ERROR: -2.0148-04 0.0 0.0 0.0 0.0 SUBSTREAM: MIXED LIQUID VAPOR PHASE: MIXED MIXED LIQUID COMPONENTS: LBMOL/HR **GLYCEROL** 0.0 0.0 0.0 0.0 0.0

ACETOL 0.0 0.0 0.0 0.0 0.0 395.0385 27.2438 367.7947 367.7947 42.1524 WATER **HYDROGEN** 494.5904 0.3463 494.2442 494.2442 0.3565 166.1698 50.0037 116.1662 116.1662 53.7172 PG N-PROP 2.2628 0.1430 2.1199 2.1199 0.1974 0.0 0.0 0.0 ETHYL-01 0.0 0.0 **ISO-PROP** 4.9896 0.1972 4.7924 4.7924 0.2646 SODIU-01 0.0 0.0 0.0 0.0 0.0 7.8827-02 3.8275-03 7.4999-02 7.4999-02 5.8603-03 METHA-01 COMPONENTS: MOLE FRAC 0.0 0.0 GLYCEROL 0.0 0.0 0.0 ACETOL 0.0 0.0 0.0 0.0 0.0 WATER 0.3716 0.3496 0.3733 0.3733 0.4359 HYDROGEN 0.4652 4.4429-03 0.5017 0.5017 3.6870-03 0.1563 0.6416 0.1179 0.1179 0.5555 PG N-PROP 2.1285-03 1.8346-03 2.1517-03 2.1517-03 2.0419-03 0.0 0.0 0.0 ETHYL-01 0.0 0.0 ISO-PROP 4.6933-03 2.5308-03 4.8644-03 4.8644-03 2.7370-03 SODIU-01 0.0 0.0 0.0 0.0 0.0 7.4146-05 4.9110-05 7.6126-05 7.6126-05 6.0607-05 METHA-01 COMPONENTS: MASS FRAC 0.0 GLYCEROL 0.0 0.0 0.0 0.0 ACETOL 0.0 0.0 0.0 0.0 0.0 0.3357 0.1137 0.3925 0.3925 0.1557 WATER HYDROGEN 4.7037-02 1.6169-04 5.9026-02 5.9026-02 1.4740-04 0.5965 0.8814 0.5237 0.5237 0.8384 PG 6.4155-03 1.9905-03 7.5472-03 7.5472-03 2.4336-03 N-PROP ETHYL-01 0.0 0.0 0.0 0.0 0.0 1.4146-02 2.7458-03 1.7062-02 1.7062-02 3.2619-03 ISO-PROP 0.0 SODIU-01 0.0 0.0 0.0 0.0 1.1916-04 2.8408-05 1.4237-04 1.4237-04 3.8513-05 METHA-01 TOTAL FLOW: LBMOL/HR 1063.1301 77.9378 985.1922 985.1922 96.6940 LB/HR 2.1197+04 4317.1178 1.6880+04 1.6880+04 4875.6888 1.6500+04 89.2728 1.6587+04 1.4834+04 97.9093 CUFT/HR STATE VARIABLES: TEMP F 455.8376 455.1614 455.1614 429.0464 429.0460 PRES PSIA 566.0365 560.0000 560.0000 559.1403 559.1403 VFRAC 1.0000 0.9019 0.0 0.9262 0.0 7.3771-02 $1.0000 \quad 0.0$ LFRAC 9.8148-02 1.0000 0.0 SFRAC 0.0 0.0 0.0 0.0 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 55 STREAM SECTION S-121 S-122 S-123 S-124 S-125 (CONTINUED) STREAM ID S-121 S-122 S-123 S-124 S-125 ENTHALPY: BTU/LBMOL -6.5193+04 -1.6354+05 -5.7413+04 -5.9433+04 -1.5820+05 BTU/LB -3269.7482 -2952.3758 -3350.9178 -3468.8219 -3137.3276 BTU/HR -6.9308+07 -1.2746+07 -5.6563+07 -5.8553+07 -1.5297+07

ENTROPY:

BTU/LBMOL-R-20.1361-76.6788-15.6425-17.8791-71.3337BTU/LB-R-1.0099-1.3843-0.9130-1.0435-1.4147DENSITY:LBMOL/CUFT6.4431-020.87305.9395-026.6413-020.9876LB/CUFT1.284648.35871.01761.137949.7980AVG MW19.938255.391817.133517.133550.4239ASPEN PLUSPLAT:WIN32VER: 24.004/11/2011PAGE 56

STREAM SECTION

S-126 S-127 S-128 S-129 S-130

-----S-126 S-127 S-128 S-129 STREAM ID S-130 FROM : F-102 HX-103 C-102 V-101 V-101 TO : HX-103 C-102 V-101 M-103 CP-101 SUBSTREAM: MIXED PHASE: VAPOR MIXED MIXED LIQUID VAPOR COMPONENTS: LBMOL/HR GLYCEROL 0.0 0.0 0.0 0.0 0.0 0.0 0.0 ACETOL 0.0 0.0 0.0 WATER 325.6423 325.6423 325.6423 308.5626 17.0797 493.8876 493.8876 493.8876 0.1372 493.7505 HYDROGEN PG 62.4490 62.4490 62.4490 62.2089 0.2401 N-PROP 1.9224 1.9224 1.9224 0.9230 0.9994 ETHYL-01 0.0 0.0 0.0 0.0 0.0 ISO-PROP 4.5277 4.5277 4.5277 0.8016 3.7261 0.0 0.0 0.0 SODIU-01 0.0 0.0 6.9139-02 6.9139-02 6.9139-02 5.7899-02 1.1240-02 METHA-01 COMPONENTS: MOLE FRAC $0.0 \quad 0.0 \quad 0.0 \quad 0.0$ GLYCEROL 0.0 ACETOL 0.0 0.0 0.0 0.0 0.0 0.3665 0.3665 0.3665 0.8279 3.3113-02 WATER 0.5559 0.5559 0.5559 3.6804-04 0.9572 HYDROGEN PG 7.0286-02 7.0286-02 7.0286-02 0.1669 4.6548-04 N-PROP 2.1637-03 2.1637-03 2.1637-03 2.4765-03 1.9376-03 ETHYL-01 0.0 0.0 0.0 0.0 0.0 5.0959-03 5.0959-03 5.0959-03 2.1508-03 7.2239-03 ISO-PROP SODIU-01 0.0 0.0 0.0 0.0 0.0 7.7815-05 7.7815-05 7.7815-05 1.5535-04 2.1791-05 METHA-01 COMPONENTS: MASS FRAC GLYCEROL 0.0 0.0 0.0 0.0 0.0 0.0 ACETOL 0.0 0.0 0.0 0.0 WATER 0.4887 0.4887 0.4887 0.5346 0.1916 HYDROGEN 8.2940-02 8.2940-02 8.2940-02 2.6592-05 0.6199 PG 0.3959 0.3959 0.3959 0.4552 1.1379-02 9.6242-03 9.6242-03 9.6242-03 5.3342-03 3.7406-02 N-PROP ETHYL-01 0.0 0.0 0.0 0.0 0.0 2.2667-02 2.2667-02 2.2667-02 4.6327-03 0.1395 ISO-PROP SODIU-01 0.0 0.0 0.0 0.0 0.0 METHA-01 1.8455-04 1.8455-04 1.8455-04 1.7841-04 2.2431-04 TOTAL FLOW: 888.4982 888.4982 888.4982 372.6911 515.8070 LBMOL/HR

1.2004+04 1.2004+04 1.2004+04 1.0398+04 1605.6541 LB/HR 1.4736+04 1.3675+04 7127.7148 178.0009 6949.7139 CUFT/HR STATE VARIABLES: TEMP F 429.0460 409.0464 225.0000 225.0000 225.0000 PRES PSIA 559.1403 558.2795 553.2795 553.2795 553.2795 1.0000 0.9382 0.5805 0.0 VFRAC 1.0000 6.1755-02 0.4195 LFRAC 0.0 1.0000 0.0 SFRAC 0.0 0.0 0.0 0.0 0.0 ENTHALPY: BTU/LBMOL -4.8685+04 -4.9959+04 -5.8397+04 -1.3434+05 -3525.5544 BTU/LB -3603.4643 -3697.8180 -4322.3718 -4814.9205 -1132.5638 BTU/HR -4.3256+07 -4.4389+07 -5.1886+07 -5.0068+07 -1.8185+06 ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 57

STREAM SECTION

S-126 S-127 S-128 S-129 S-130 (CONTINUED)

STREAM ID S-126 S-127 S-128 S-129 S-130

ENTROPY:

| BTU/LBMOL-R | -12.0617 -13.5096 -24.1456 -49.0038 -6.1846 |
|--------------|---------------------------------------------|
| BTU/LB-R | -0.8928 -0.9999 -1.7872 -1.7564 -1.9868 |
| DENSITY: | |
| LBMOL/CUFT | 6.0293-02 6.4970-02 0.1247 2.0938 7.4220-02 |
| LB/CUFT | 0.8146 0.8778 1.6841 58.4178 0.2310 |
| AVG MW | 13.5105 13.5105 13.5105 27.9009 3.1129 |
| ASPEN PLUS P | LAT: WIN32 VER: 24.0 04/11/2011 PAGE 58 |

STREAM SECTION

S-131 S-132 S-133 S-134 S-135

| STREAM ID | S-13 | 1 S-132 | S-133 | S-134 | S-135 |
|-----------|--------|---------|-------|-------|-------|
| FROM : | CP-101 | M-103 | VALVE | D-102 | C-103 |
| TO : | M-101 | VALVE | D-102 | C-103 | |

SUBSTREAM: MIXED

| PHASE: | VAPOR | MIXE | ED 1 | MIXED | LIQUID | LIQUID |
|------------|------------|----------|---------------|----------|------------|-------------|
| COMPONENTS | S: LBMOL/H | IR | | | | |
| GLYCEROL | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | |
| ACETOL | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | |
| WATER | 17.0797 | 377.958 | 39 37' | 7.9589 | 2.1127 | 2.1127 |
| HYDROGEN | 493.7 | 505 0.3 | 8399 | 0.8399 | 1.5652-20 | 1.5652-20 |
| PG | 0.2401 165 | 5.9297 | 165.92 | 97 162. | 1873 162. | 1873 |
| N-PROP | 0.9994 | 1.2634 | 1.26 | 534 3.63 | 66-06 3.63 | 66-06 |
| ETHYL-01 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | |
| ISO-PROP | 3.7261 | 1.2635 | 5 1.2 | 2635 9.1 | 415-10 9.1 | 415-10 |
| SODIU-01 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | |
| METHA-01 | 1.1240-0 | 02 6.758 | 6-02 6 | 5.7586-0 | 2 1.6305-0 | 6 1.6305-06 |
| COMPONENTS | S: MOLE FR | AC | | | | |
| GLYCEROL | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | |
| ACETOL | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | |

3.3113-02 0.6906 0.6906 1.2859-02 1.2859-02 WATER 0.9572 1.5346-03 1.5346-03 9.5267-23 9.5267-23 HYDROGEN PG 4.6548-04 0.3032 0.3032 0.9871 0.9871 1.9376-03 2.3084-03 2.3084-03 2.2134-08 2.2134-08 N-PROP ETHYL-01 0.0 0.0 0.0 0.0 0.0 7.2239-03 2.3085-03 2.3085-03 5.5639-12 5.5639-12 ISO-PROP SODIU-01 0.0 0.0 0.0 0.0 0.0 2.1791-05 1.2349-04 1.2349-04 9.9242-09 9.9242-09 METHA-01 COMPONENTS: MASS FRAC GLYCEROL 0.0 0.0 0.0 0.0 0.0 ACETOL 0.0 0.0 0.0 0.0 0.0 0.1916 0.3476 0.3476 3.0745-03 3.0745-03 WATER 0.6199 8.6428-05 8.6428-05 2.5488-24 2.5488-24 HYDROGEN PG 1.1379-02 0.6445 0.6445 0.9969 0.9969 N-PROP 3.7406-02 3.8755-03 3.8755-03 1.7653-08 1.7653-08 ETHYL-01 0.0 0.0 0.0 0.0 0.0 0.1395 3.8757-03 3.8757-03 4.4376-12 4.4376-12 ISO-PROP SODIU-01 0.0 0.0 0.0 0.0 0.0 METHA-01 2.2431-04 1.1054-04 1.1054-04 4.2203-09 4.2203-09 TOTAL FLOW: LBMOL/HR 515.8070 547.3230 547.3230 164.3000 164.3000 1605.6541 1.9591+04 1.9591+04 1.2380+04 1.2380+04 LB/HR 6597.3326 357.4023 1.7899+04 233.5486 193.9751 CUFT/HR STATE VARIABLES: TEMP F 244.4171 317.2634 254.7511 364.0271 90.0000 PRES PSIA 600.0000 548.2795 30.0000 20.0000 15.0000 VFRAC 1.0000 1.7993-04 0.1278 0.0 0.0 1.0000 LFRAC 0.0 0.9998 0.8722 1.0000 SFRAC 0.0 0.0 0.0 0.0 0.0 ENTHALPY: -3384.1109 - 1.4271 + 05 - 1.4271 + 05 - 1.9740 + 05 - 2.1311 + 05BTU/LBMOL BTU/LB -1087.1259 -3986.9863 -3986.9863 -2619.7765 -2828.3357 BTU/HR -1.7455+06 - 7.8110+07 - 7.8110+07 - 3.2432+07 - 3.5014+07ASPEN PLUS PLAT: WIN32 VER: 24.0 04/11/2011 PAGE 59

STREAM SECTION

S-131 S-132 S-133 S-134 S-135 (CONTINUED)

 STREAM ID
 S-131
 S-132
 S-133
 S-134
 S-135

 ENTROPY:
 BTU/LBMOL-R
 -6.1443
 -56.4712
 -56.2361
 -111.9337
 -135.0044

 BTU/LB-R
 -1.9738
 -1.5776
 -1.5711
 -1.4855
 -1.7917

 DENSITY:
 LBMOL/CUFT
 7.8184-02
 1.5314
 3.0578-02
 0.7035
 0.8470

 LB/CUFT
 0.2434
 54.8156
 1.0945
 53.0072
 63.8214

 AVG MW
 3.1129
 35.7946
 35.7946
 75.3485
 75.3485

 ASPEN PLUS
 PLAT: WIN32
 VER: 24.0
 04/11/2011
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STREAM SECTION

S-136 S-137

STREAM ID S-136 S-137 FROM : D-102 D-102 TO : SUBSTREAM: MIXED PHASE: LIQUID VAPOR COMPONENTS: LBMOL/HR GLYCEROL 0.0 0.0 ACETOL 0.0 0.0 WATER 375.5238 0.3223 **HYDROGEN** 7.0112-05 0.8399 PG 3.7416 8.8865-04 N-PROP 0.3473 0.9162 ETHYL-01 0.0 0.0 4.0319-02 **ISO-PROP** 1.2232 SODIU-01 0.0 0.0 METHA-01 6.6709-02 8.7551-04 COMPONENTS: MOLE FRAC GLYCEROL 0.0 0.0 ACETOL 0.0 0.0 0.9889 9.7583-02 WATER HYDROGEN 1.8464-07 0.2543 9.8535-03 2.6902-04 PG N-PROP 9.1452-04 0.2773 0.0 ETHYL-01 0.0 **ISO-PROP** 1.0618-04 0.3703 SODIU-01 0.0 0.0 METHA-01 1.7568-04 2.6504-04 COMPONENTS: MASS FRAC GLYCEROL 0.0 0.0 ACETOL 0.0 0.0 WATER 0.9562 4.2649-02 1.9976-08 1.2435-02 HYDROGEN PG 4.0241-02 4.9664-04 N-PROP 2.9496-03 0.4044 ETHYL-01 0.0 0.0 3.4246-04 0.5399 **ISO-PROP** 0.0 0.0 SODIU-01 METHA-01 3.0211-04 2.0603-04 TOTAL FLOW: LBMOL/HR 379.7197 3.3033 LB/HR 7075.3126 136.1602 117.1993 1020.6045 CUFT/HR STATE VARIABLES: TEMP F 130.0000 130.0000 PRES PSIA 20.0000 20.0000 VFRAC 0.0 1.0000 1.0000 0.0 LFRAC 0.0 SFRAC 0.0 **ENTHALPY:** -1.2347+05 -8.3091+04 BTU/LBMOL BTU/LB -6626.6697 -2015.8168 BTU/HR -4.6886+07 -2.7447+05

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STREAM SECTION

S-136 S-137 (CONTINUED)

STREAM ID S-136 S-137

| ENTROPY: | |
|--------------|----------------------------|
| BTU/LBMOL-F | R -39.0904 -47.4015 |
| BTU/LB-R | -2.0979 -1.1500 |
| DENSITY: | |
| LBMOL/CUFT | 3.2399 3.2366-03 |
| LB/CUFT | 60.3699 0.1334 |
| AVG MW | 18.6330 41.2194 |
| ASPEN PLUS F | PLAT: WIN32 VER: 24.0 |

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PROBLEM STATUS SECTION

BLOCK STATUS

| *************************************** | *************************************** |
|----------------------------------------------|-----------------------------------------|
| * | * |
| * Calculations were completed with errors | * |
| * | * |
| * The following Unit Operation blocks were | * |
| * completed with warnings: | * |
| * HX-102 HX-103 | * |
| * | * |
| * All streams were flashed normally | * |
| * | * |
| * The following Convergence blocks were | * |
| * completed with errors: | * |
| * \$OLVER02 | * |
| * | * |
| * All Sensitivity blocks were completed norm | ally * |
| * | * |
| *************************************** | *************************************** |