

Hydrogen Distribution Network of Medor Refinery Plant by Using Two Different Optimization Techniques

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Abstract

Hydrogen consumption of refineries is rising sharply with the increase of conversion units such as hydrocracking and hydrotreating processes, since these processes play an important role in modern and integrated refineries to upgrade the heavy crude oil to more valuable products. Therefore hydrogen distribution network becomes unavoidable inspite of its complexity. So the refinery hydrogen management techniques are developed in recent years to determine maximum hydrogen recovery networks which do not account for varying operating conditions of hydrogen consuming processes and assume constant operating conditions. This paper addresses the problem of actual hydrogen distribution at the Medor refinery plant at Alexandria-Egypt in which different optimization techniques were applied to minimize the consumption of hydrogen. Additionally, an efficient network design has been achieved that reduces the hydrogen utilities usage by 14.9%, hydrogen discharge by 33% and 3.242 MUS\$/year saving in the operating costs.

Keywords

H₂ integration, H₂ distribution network, Optimization techniques, H₂ management.

Introduction

There are reasons for refineries to increase their use of hydrogen in recent years. The first reason is the move to heavier crud oils made refineries increase the use of hydrocracking processes as a means of upgrading heavy oils to middle distillates. Another reason is the regulations on the sulphur content of fuels made refineries increase the use of hydrotreating processes. Catalytic reforming process is considered as a major source of hydrogen for refinery industry catalytic reforming process produces amount of hydrogen as a by-product and this amount of

hydrogen is affected by the need for the low aromatic gasoline. As a result, the overall availability of hydrogen in the refinery is decreased [1, 2]. Some specific technologies are applied on hydrogen network such as purification technologies to increase the amount of recycled hydrogen [3, 4]. If the amount of recycled hydrogen is increased, the amount of hydrogen that is sent to the furnaces with the off gas will decrease. As a result, the amount of fresh hydrogen obtained from hydrogen plant is decreased and can be removed from the network [5, 6]. There are many processes in the refinery dealing with

hydrogen if we separate these processes and hydrogen plant from other refinery processes, a refinery hydrogen network can be formed. Fig (1) [7] gives us an example of refinery hydrogen network with two hydrogen producers and six hydrogen consumers. In this hydrogen network, the two main hydrogen producers are the catalytic reforming and hydrogen plant. Catalytic reforming increases the octane number of heavy naphtha by cyclization and dehydrogenation of aliphatic hydrocarbon molecules into aromatic compounds and at the same time generates large amounts of hydrogen at 70-90% purity as a by-product [8]. The second hydrogen producer is the hydrogen plant which produces hydrogen with purity of exceeding 99% [8, 9]. The hydrogen consumers are hydrocracker (HCU), diesel hydrotreater (DHU), kerosene hydrotreater (KHU), cracked naphtha hydrotreater (CNHU), naphtha hydrotreater (NHU), and hydrodealkylation (HDA). After hydrogen consumption, the purge gas will be sent from hydrogen consumers outlets to the fuel system. In this paper, the application of a mass integration procedure techniques hydrogen appropriate purifier from pressure swing adsorption processes (PSA), and membranes for recovering hydrogen from refinery off-gases is applied to minimize the hydrogen utility.

Literature Review

Hydrogen management was considered to be important in both design and operation. Simpson 1984 [9] proposed the work over hydrogen management that is based on the

analysis of the hydrocarbon thermodynamic. Appropriate operating conditions and strategy of using catalyst are two main factors of

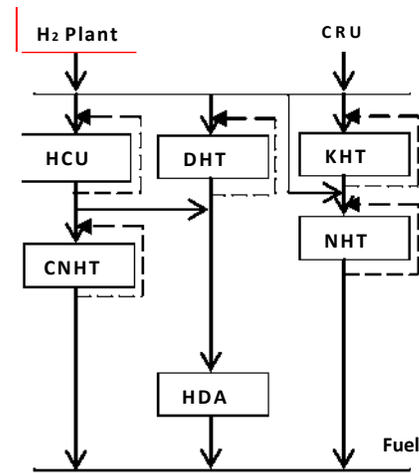


Figure 1 A typical hydrogen network..

Towler et al., 1996 [10] developed the first systematic approach for hydrogen management. Economics analysis of hydrogen recovery against added values in product by hydrogen is proposed as the main feature in this method. Hydrogen is recovered for a cost and brings extra value to fuel products. When the extra value brought by hydrogen can not compensate the cost of hydrogen recovery, it is not preferred to recover hydrogen because no profit can be made. Under this concept, the cost and value composite curves can be plotted for either hydrogen producers or consumers. The value added to products can be calculated as the value of products minus the summation of the value of feedstock, operating cost and capital cost. The cost of hydrogen recovery is represented by the cost of hydrogen purification units. The proposed methodology can be used not only for an economic analysis of a refinery hydrogen network, but also for refinery operation management, sensitivity analysis and in

examining retrofit design options. However, the essential economic data to the analysis such as the added value by adding hydrogen will not be always available for refineries, bringing difficulties in applying the method. Another limitation of this method is the lack of hydrogen purifier selection and placement strategies. Linnhoff et al., 1979 [11] proposed the pinch technology for heat exchanger network synthesis. By plotting cold streams and hot streams data into a composite curve, the overall heat exchanger network's pinch point can be found leading a theoretical optimal solution. Alves, 1999 [6] utilized Linnhoff's work and extended the pinch technology into the hydrogen network field. Hydrogen sinks and sources are introduced similarly to the cold and hot exchanger networks. With observation on the balance between hydrogen sinks and sources, hydrogen pinch analysis gives a general overview of the hydrogen usage situation of a specific hydrogen network. In order to apply the pinch technology on hydrogen networks, hydrogen sources and sinks must be defined in a simplified hydrogen consumer model [6].

Fig (2) shows a simplified hydrogen consumer model. The figure illustrates how hydrogen flows and is used through a process. The hydrogen sink, located at the inlet of the consumer, is defined as the mix of the makeup hydrogen and the recycle stream. The makeup hydrogen mainly comes from a H₂ plant or a catalytic reformer. F_{sink} and Y_{sink} are used to denote the flow rate and purity of a sink. On the other hand, a hydrogen source locates at the outlet of a hydrogen consumer, containing a purge stream and a recycle stream. A hydrogen source is a hydrogen-rich stream that can be utilized by hydrogen

consumers. It can be off-gas from other hydrogen consumers. In the hydrogen consumer model the hydrogen source would be the mixture of purge and recycle stream. F_{source} and Y_{source} are used to denote the flow rate and purity of a source. Fig (2) also demonstrates how a hydrogen consumer unit works. Make-up hydrogen will be mixed with liquid hydrocarbon feed. The mixture is then sent into a reactor for reaction under certain operating conditions. The after reaction stream goes into the flash separation unit and gets stripped into vapour and liquid. The vapour phase portion can be recycled or purge, while the liquid phase becomes a fuel product afterwards.

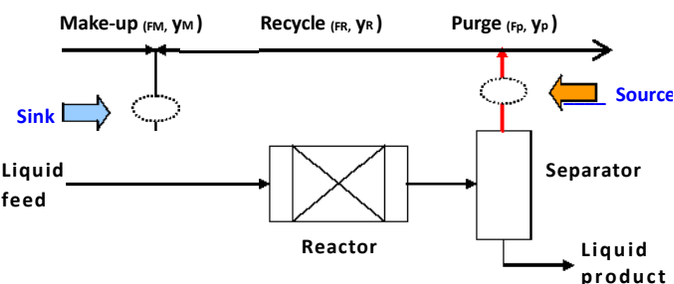


Figure 2. Simplified representation of a hydrogen consuming process in hydrogen pinch analysis [13].

Alves, 1999 [6] proposed a linear programming (LP) approach for optimizing H₂ network. Hallale et al., 2001 [1] and Liu, 2002 [12] developed the methodology of automated hydrogen network design using a mixed integer non-linear programming (MINLP) method. Liu has taken the pressure into consideration as well as the hydrogen purifier placement strategy. El Halwagi, 2003 [18] proposed a mathematical linear program for optimizing H₂ network. Ahmed, 2011 [7] extended the problem formulation to address the multi-period problem faced by H₂ network. All these design methods have a

common feature of representing a hydrogen network design problem as a superstructure, and then use mathematical programming algorithms to obtain optimal solutions.

Hydrogen purification methods

There are three hydrogen purification methods, pressure swing adsorption (PSA), membranes, and cryogenic units. Selecting the most appropriate hydrogen purification technology, depends upon both procedural requirements (hydrogen recovery, feed, and product) and operational requirements (flexibility, reliability, feed pre-treatment, and by-product recovery) [13, 14, 15].

- (I) Pressure swing adsorption (PSA) units are well suited for purifying catalytic reformer hydrogen used in hydro-processing units. Feed purity of the PSA units is up to 40 vol. %. PSA units produce hydrogen with recovery up to 90 % and with 99.9 vol. % purity.
- (II) Membrane units are suited for recovering hydrogen from high pressure purge gases. Membrane is used when the feed purity is up to 25 vol. %. The hydrogen product is with recovery up to 95% and with 98 vol. % purity.
- (III) Cryogenic units used when the hydrogen content of the feed is 15-80 vol. %. Cryogenic units require feed pre-treatment and give by-products such as methane and ethane. In the cryogenic units, the hydrogen product is with recovery up to 98% and 97 vol. % purity [12].

By adding a purifier to the hydrogen network, a new sink (the feed stream) and two new sources (product and residue streams) have been actually added to the network. Fig (3) shows a schematic diagram of the purifier. Mass balance around the purifier gives:

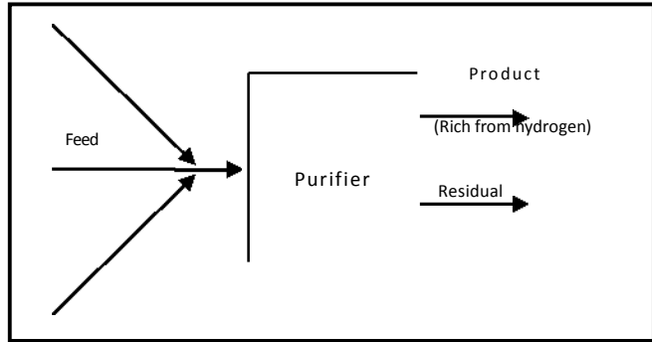


Figure 3. Schematic diagram of purifier [12]

- Viscous
- fluids Foam

$$\sum_i F_{i,pur} = \sum_j F_{prod,j} + F_{residual} \quad (1)$$

Hydrogen balance around the purifier gives:

$$y_{prod} * \sum_j F_{prod,j} = R * \sum_i F_{i,pur} * y_i \quad (2)$$

The low-impurity product stream can either be reused in another hydrogen consuming process or recycled to the same process. A high-impurity by-product stream is either purged or used as fuel. The works applied in this paper are:

- (i) Study and apply the cascade analysis technique.
- (ii) Study and apply mathematical programming technique.
- (iii) Apply the two techniques to industrial case study of Medore refinery plant at Alexandria-Egypt.
- (iv) Study two different purification units, membrane and pressure swing adsorption (PSA).
- (v) Comparison between the costs of the hydrogen network of Medore refinery plant with the two purification units.

Cascade analysis technique

The first step in conducting the cascade technique is to locate the various sinks and sources at their respective concentration levels. As shown in Table 4, the concentration levels (C_k) are arranged in an ascending order and the flow rate of each sink (F_j) and source (F_i) is located individually at its respective concentration level k in columns 3 and 4. Column 5 is the net flow rate $\sum_i F_i - \sum_j F_j$ between sources and sinks at each concentration level k ; positive values indicate a surplus, negative values indicate a deficit [16]. The next step is to cascade the net flow rate surplus/deficit down the concentration levels to yield the cumulative surplus/deficit flow rate (F_c) shown in column 6 of Table 4 and, the overall net surplus/deficit for the network. A zero fresh flow rate is assumed at this stage to facilitate the search for the minimum utility gas flow rate. The two final steps in getting the minimum utility targets involve setting up the cumulative impurity load cascade (cum Δm , column 8), and finally calculating the flow rate of fresh hydrogen needed at each concentration level, or the interval fresh gas flow rate, ($F_{F,k}$). The largest negative $F_{F,k}$, which actually represents the minimum fresh gas target that cascaded down the cumulative gas flow rate column (F_c), yields the minimum discharge flow rate. The impurity load (Δm) shown in Column 7 of Table 4 is obtained from the product of cumulative flow rate (F_c) and the concentration difference across two concentration levels (ΔC).

$$\Delta m = F_c * \Delta C / 1000 \quad (10)$$

Cascading the impurity load down the concentration levels of column 8 yields the cumulative load (cum Δm). The amount of interval fresh gas flow rate ($F_{F,k}$) is obtained by dividing cum Δm by the difference between the concentration at level k (C_k) and the fresh feed concentration (C_F).

$$F_{F,k} = \frac{\text{cum } \Delta m * 1000}{C_k - C_F} \quad (11)$$

The absolute value of the largest negative ($F_{F,k}$) is the minimum fresh gas flow rate target (F_F). This amount is cascaded down the F_c column of the feasible cascade as described in Table 5, to produce the minimum discharge flow rates (F_D) [16]. The changes in the cascade tables in case of adding a purifier is the location of the flow rate of fresh hydrogen F_F as described in the cascade technique in the reference [16]. F_F is located in the column 5 instead of column 6 at its concentration as described in Table 6 and Table 7. Also the feed flow at the pinch point is decreased by the amount of the discharge flow rate which purified by the purifier. In the cascade technique, the hydrogen network is divided into two design regions above the pinch and below the pinch. Above the pinch region, the fresh hydrogen is used and no fuel generation. In each interval above the pinch, the sink takes its requirements from the source in that interval. Fresh hydrogen is added to any sink in any interval if the source flow rate is less than the sink flow rate in that interval and there is not any source residual from the above intervals. Below the pinch region, there is not fresh hydrogen to be used and only fuel is generated. Each sink in any interval below the pinch takes its requirements from the source in that interval. If the source flow rate in any interval is less than the sink flow rate in that interval, any source residual from the above intervals must be added to the sink. The unused source residual from each interval is discharged as a fuel [19].

Mathematical Programming Technique

A source-sink representation as described in Figure 4 is the first step in the application of the mathematical technique. Each source is split into fractions (of unknown flow rate) that are allocated to the various sinks. A waste sink is an additional sink placed to account for unrecycled / unreused material. Also, the fresh resource is allowed to split and is allocated to all process sinks except the waste sink [17, 18].

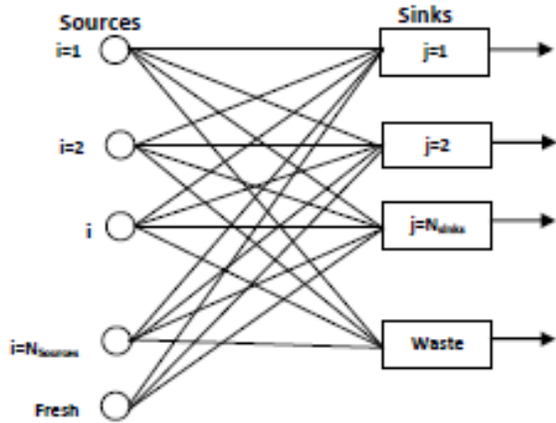


Figure 4. Source/ Sink allocation [18].

The objective function is to minimize the consumption of fresh resources.

Minimize consumption of fresh resource

$$= \sum_{j=1}^{N_{sinks}} F_j \quad (3)$$

Each source i , is split into N_{sinks} fractions as described in fig (5). The flow rate of each split is j . Also, one split is forwarded to the waste sink which is denoted by $w_{i,waste}$ [18].

$$w_i = \sum_{j=1}^{N_{sink}} w_{i,j} + w_{i,waste} \quad \text{for } i = 1,2,3, \dots, N_{source} \quad (4)$$

The following step is the mixing of the split fractions into a feed to the j th sink as described in Figure 6. The split fractions come from the process sources and the fresh stream [18].

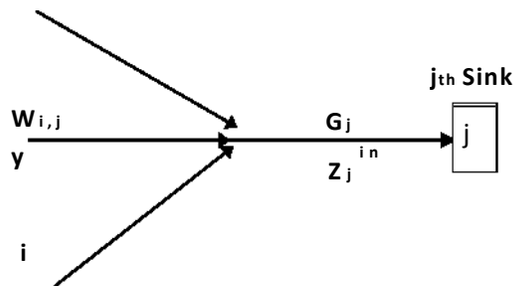


Figure 6. Mixing of Sources at inlets of Sinks [18].

$$G_j = F_j + \sum_{l=1}^{N_{sources}} w_{l,j} \quad \text{for } j = 1,2,3, \dots, N_{sinks} \quad (5)$$

$$G_j z_j^{in} = \sum_{l=1}^{N_{sources}} w_{l,j} y_l \quad \text{for } j = 1,2,3, \dots, N_{sinks} \quad (6)$$

Where:

$$z_j^{min} \leq z_j^{in} \leq z_j^{max} \quad \text{for } j = 1,2,3, \dots, N_{sinks} \quad (7)$$

To ensure the non-negativity of the fresh flows and the fractions of sources allocated to a sink, the following constraints are applied:

$$w_{i,j} \geq 0 \quad \text{for } i = 1,2,3, \dots, N_{sources} \text{ and } j = 1,2,3, \dots, N_{sinks} \quad (8)$$

$$F_j \geq 0 \quad \text{for } j = 1,2,3, \dots, N_{sinks} \quad (9)$$

The above formulations are a linear program that can be solved to obtain the optimal target and source-sink matches [18]. In this work Lingo program v.13 is used. By applying the mathematical technique, The network design is obtained through the program. The total annual cost is calculated for all cases, existing H₂ network, integrating H₂ network by mathematical or cascade technique, integrating H₂ network with adding a Membrane unit, and integrating H₂ network with adding a PSA unit. The total annual cost is the summation of the annual operating costs and annualized capital costs for new equipments.

$$TAC = Cost_{operating} + Af (Cost_{equipment}) \quad (12)$$

Operating costs

Operating costs are the operating costs of hydrogen production, the operating costs of electricity of the compressors, and the operating costs of the fuel value.

$$Cost_{operating} = OC_{H2} + OC_{ELC} - OC_{fuel} \quad (13)$$

$$OC_{H2} = OCH + FHP \quad (14)$$

OC_{H2} is the hydrogen production operating cost which is function of the FHP, hydrogen plant flow rate, multiplied by the unit cost of production, OCH.

$$OC_{ELC} = OCE \left(\sum_K PWT_k \right) \quad (15)$$

OC_{ELC} is the electricity operating cost, which is a function of the P_{wr}, compressor's power, multiplied by the unit cost of electricity, OCE.

$$Power = 160.376 \left[\left(\frac{P_{out}}{P_{in}} \right)^{0.1857} - 1 \right] F_{in} \quad (16)$$

- Power calculated from equation (16) is in KW, and the flow rate into the compressor (F_{in}) is in MMSCFD. Discharge (P_{out}) and suction pressure (P_{in}) of compressor are in psi. Fuel gas is assumed to be a binary mixture of hydrogen and methane and both are functions of the fuel gas flow rate and the purity [20].

$$OC_{Fuel} = OCF * F_{Fuel} * (LHV_{H_2} * Y_{H_2} + LHV_{CH_4} * Y_{CH_4}) \quad (17)$$

- OC_{Fuel} is the fuel gas value operating cost, which is function of the summation of fuel gas heating value LHV for hydrogen and methane multiplied by unit heat cost of fuel, OCF.

Equipment Cost

Equipment costs are the cost of the PSA or membrane purifier added to the H₂ network. The cost of a PSA unit has been correlated as a simple linear function of the feed flow rate [20].

$$Cap_{PSA} = a_{PSA} + b_{PSA} * F_{in} \quad (18)$$

- Cap_{PSA} is the capital cost of the PSA unit, which is in *10³\$. The capital cost coefficients a_{PSA} and b_{PSA} are constants as 503.8 and 347.4, respectively and the PSA feed flow rate, F_{in} is measured in MMSCFD. - Membrane costs are more complex to estimate as those that depend on product purity as well as the membrane pressure drop. Ratan, 1994 [21] provides some data [1].

-Af means annualizing factor, and is addresses

$$as: \quad Af = \frac{f * (1+f)^n}{(1+f)^n - 1} \quad (19)$$

Where f_i is the fractional interest and n_y is the number of years.

Case study

This case study is representative of a real refinery system. Fig (7) shows the existing hydrogen network in the Medor Refinery Plant at Alexandria-Egypt. There are four consuming units, naphtha hydrotreating, isomerisation, diesel hydrotreating, and hydrocracking unit. The hydrogen is supplied by catalytic reforming unit and hydrogen plant. All the consuming units have recycle compressors except the isomerization unit. Currently 2263.67 kmole/hr hydrogen is produced from hydrogen plant. The data related to the flow rate and purity is represented in Table 1. PSA unit is used to purify some amount of hydrogen flow rate from the catalytic reforming unit and produced purified hydrogen with 99.9% purity. All purified hydrogen from PSA unit mixed with the hydrogen produced from hydrogen plant and sent to the hydrocracking unit. Table 2 represents the sink streams and the source streams of the existing network in the refinery.

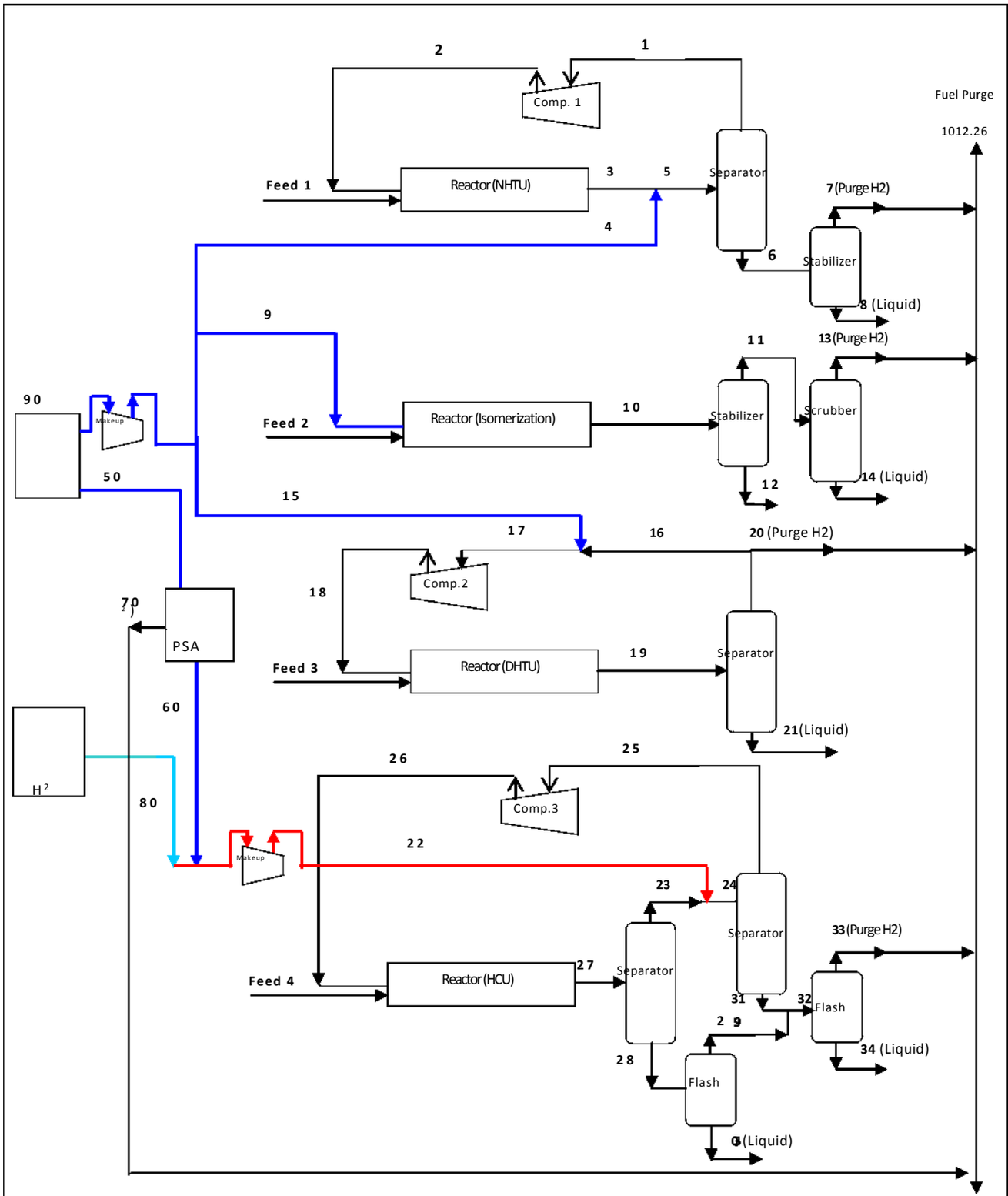


Figure 7. The Existing Network in the Case Study.

Table 1: Streams data for the H₂ network in the case study.

Stream No.	Flow Rate (Kg mole/hr)	H ₂ Purity (Mole %)	Pressure bar	Temp. C°
1	2024.99	95.62	41.4	55.0
2	2024.99	95.62	52.4	77.0
3	3472.2	53.56	45.0	343.0
4	135.56	90.21	44.8	38.0
5	3607.76	54.94	42.1	134.0
7	97.9	46.84	7.0	48.0
9	241.3	90.22	44.8	38.0
10	1120.37	4.57	15.38	124.0
11	121.96	41.99	7.58	35.0
13	122.6	41.78	4.50	36.0
15	476.54	90.22	44.8	38.0
16	2396.94	83.04	34.5	52.0
17	2873.48	84.23	34.5	50.0
18	2873.48	84.23	50.5	84.0
20	Zero	Zero	Zero	Zero
22	3276.9	99.9	166.03	123.0
23	20088.76	83.86	164.99	198.0
24	23365.66	86.11	164.99	157.0
25	22434.72	89.21	164.65	60.0
26	22434.72	89.21	187.79	76.0
33	515.15	73.48	31.03	56.0
50	1287.8	90.164	22.0	38.0
60	1011.19	99.9	20.0	38.0
70	276.61	54.57	2.0	38.0
80	2263.67	99.99	20.0	38.0
90	2141.2	90.141	22.0	38.0

Table 2: Sinks and Sources streams in the case study.

Sink Stream s	Flow Rate (kgmole/hr)	H₂ Purity (Mole %)	Impurity (Mole %)
2(NHT)	2024.99	95.62	4.38
4(NHT)	135.56	90.22	9.78
9(ISO.)	241.3	90.22	9.78
18(DHT)	2873.48	84.23	15.77
22(HC)	3276.9	99.9	0.1
26(HC)	22434.72	89.21	10.79
50(PSA)	1287.8	90.164	9.836
Source Streams	Flow Rate (kgmole/hr)	H₂ Purity (Mole %)	Impurity (Mole %)
1(NHT)	2024.99	95.62	4.38
7(NHT)	97.9	46.84	53.16
13(ISO.)	122.6	41.78	58.22
16(DHT)	2396.94	83.04	16.96
25(HC)	22434.72	89.21	10.79
33(HC)	515.15	73.48	26.52
60(PSA)	1011.19	99.9	0.1
70(PSA)	276.61	54.57	45.43
90(CRU)	2141.2	90.186	9.814
80 (H2 Plant) to be determined	2263.67	99.99	0.01

Results and Discussion

Integration of H₂ network

The two techniques have been applied on the integration of Medore hydrogen network and the results showed that:

- By applying the cascade technique as described in Table 3 and Table 4, it is found that the minimum hydrogen utility would be 2257.9641 kmole/hr instead of 2263.67 kgmole/hr and the minimum discharge would be 1004.5141 kgmole/hr instead of 1012.26. Fig (8) shows the optimum design procedure without adding any new equipment to the network. Dotted lines illustrate the new connections.
- By applying the mathematical technique on the Medor hydrogen network by using lingo program v13 it is found that the minimum hydrogen utility would be 2257.964 kmole/hr instead of 2263.67 kgmole/hr and the minimum discharge would be 1004.514 kgmole/hr instead of 1012.26 kgmole/hr. Fig (9) shows the optimum design procedure without adding any new equipment to the network. Dotted lines illustrate the new connections.

Adding a purifier

For more recovery of hydrogen from off gases, two purification units have been added to the hydrogen network. Membrane and PSA.

Adding a membrane to the H₂ network

Given a gas separation membrane with a hydrogen recovery of 95% [1], one possible option is to purify the portion of stream (DHT) that lies at 16.96%. 1004.5141 kgmole/hr is purified to a product stream at 2% impurity. From material balance calculations, the flow rate of this high quality permeate stream is 808.61 kgmole/hr. The flow rate and impurity concentration for the

retentate stream are 195.9 kgmole/hr and 78.71%, respectively. Adding these two new hydrogen sources at their respective concentration levels yields a new cascade table shown in Table 5. From this table, the fresh hydrogen flow rate is reduced to 2163.216 kmole/hr, which corresponds to a reduced discharge flow rate of 909.756 kmole/hr. The saving in fresh hydrogen is 4.44% and the discharge is reduced by 9.43%. When applied the mathematical technique on the hydrogen network with adding a membrane to the hydrogen network, it is found that the fresh hydrogen flow rate is 2163.241 Kmole/hr and the hydrogen discharge is 909.791 kmole/hr.

Adding a PSA to the H₂ network

When another hydrogen purifier is used, a PSA unit with a hydrogen-recovery value of 90% and capability to purify the hydrogen stream up to 0.10% impurity [1]. From material balance calculations, the product and residue flow rates of 751.49 kmole/hr and 253.03 kmole/hr, respectively are obtained, with the residue stream impurity concentration at 67.03%. Resetting targets using the cascade technique gave a reduced fresh hydrogen flow rates of 1926.35 kmol/hr, which corresponds to a reduced discharge flow rate of 672.91 kmol/hr as shown in Table 6. The saving in fresh hydrogen is 14.9% and the discharge is reduced by 33.0%. Fig (10) shows the optimum design procedure for the Medor hydrogen network with adding the new PSA unit as a purifier. When applied the mathematical technique on the hydrogen network with adding a PSA, it is found that the fresh hydrogen is 1926.356 kmole/hr and the hydrogen discharge is 672.946 kmole/hr. Figure 11 shows the optimum design procedure for the Medor hydrogen network with adding the new PSA unit.

Cost Calculation Results

The utility prices used are hydrogen 2000 US\$/MSCF (0.075 US\$/Nm³), power 0.03 US\$/KWh, and fuel gas 2.5 US\$/MBTU (8.53 US\$/MWh) [20]. The results of cost calculation are showed in Table 7. As described in Table 7, it is noted that: The power costs for all cases are the same and that because the flow rates entering the compressors in all cases are the same but the sources of these amounts are different. In case of integration 112 network by cascade or mathematical technique, it is found that the operating cost saving is 26000 US\$/yr. The saving in the operating cost is low that because the minimum fresh hydrogen obtained by the integration of 112 network was decreased by 5.706 kmole/hr than the actual fresh hydrogen used without integration of the 112 network. Also, the hydrogen discharge was decreased by 7.746 kmole/hr than the actual hydrogen discharge. In case of integration 112 network by cascade or mathematical technique with adding a membrane unit, it is found that the operating cost saving is 1.246 MUS\$/yr. The saving in the operating cost is increased that because the minimum fresh hydrogen obtained by the integration of 112 network with adding a membrane unit was decreased by 100.454 kmole/hr by cascade technique or 100.429 kmole/hr by mathematical technique than the actual fresh hydrogen used without integration of the 112 network. Also, the hydrogen discharge was decreased by 102.504 kmole/hr by cascade technique or 102.469 kmole/hr by mathematical technique than the actual hydrogen discharge. In case of integration 112 network by cascade or mathematical technique with adding a PSA unit, it is found that the operating cost saving is 3.242 MUS\$/yr. The saving in the operating cost is increased than the previous cases that because the minimum fresh hydrogen obtained by the integration of 112 network with adding a PSA unit was decreased by 337.32 kmole/hr by cascade technique or 337.314 kmole/hr by

mathematical technique than the actual fresh hydrogen used without integration of the 112 network. Also, the hydrogen discharge was decreased by 339.35 kmole/hr by cascade technique or 339.314 kmole/hr by mathematical technique than the actual hydrogen discharge. The PSA purifier plays a more important role in reducing the fresh hydrogen target in the hydrogen network of Medore refinery plant, compared to the gas-separation membrane. This is because a PSA generates an additional amount of purified hydrogen with higher hydrogen purity which leads to minimization of the hydrogen utilities usage by 14.9% and hydrogen discharge by 33.0%. Also 3.242 MUS\$/year saving in the operating costs achieved by using the PSA purifier.

Table 3: Infeasible Cascade Table for the hydrogen network in the case study (Hydrogen Integration).

Level	Concentration Mole%	$F_{sources}$ Kmole/hr	F_{sinks} Kmole/hr	$\sum_i F_i - \sum_j F_j$ Kmole/hr	F_c Kmole/hr	Δm Kmole/hr	Cum Δm Kmole/hr	$F_{F,k}$ Kmole/hr
1	0.01	0.0	0.0	0.0	0.0			
2	0.1	1011.19	3276.9	-2265.71	-2265.71	-9.6972	0.0	
3	4.38	2024.99	2024.99	0.0	-2265.71	-12.2348	-9.6972	-2219.0481
4	9.78	0.0	376.86	-376.86	-2642.57	-0.0898	-21.9321	-2244.8380
5	9.814	2141.20	0.0	2141.20	-501.37	-0.0110	-22.0219	-2246.2178
6	9.836	0.0	1287.8	-1287.8	-1789.17	-1.7069	-22.0329	-2242.3112
7	10.79	22434.72	22434.72	0.0	-1789.17	-8.9101	-23.7398	-2202.2096
8	15.77	0.0	2873.48	-2873.48	-4662.65	-5.5486	32.6498	-2071.6935
9	16.96	2396.94	0.0	2396.94	-2265.71	-21.6602	-38.1984	-2253.5953
10	26.52	515.15	0.0	515.15	-1750.56	-33.1031	-59.8586	-2257.9641

Table 4: Feasible Cascade Table for the hydrogen network in the case study.

Level	Concentration Mole%	$F_{sources}$ Kmole/hr	F_{sinks} Kmole/hr	$\sum_i F_i - \sum_j F_j$ Kmole/hr	F_c Kmole/hr	Δm Kmole/hr	Cum Δm Kmole/hr
1	0.01	0.0	0.0	0.0	$F_p = 2257.9641$ 2257.9641		0.0
2	0.1	1011.19	3276.9	-2265.71	-7.7459	0.2032	0.2032
3	4.38	2024.99	2024.99	0.0	-7.7459	-0.0332	0.1701
4	9.78	0.0	376.86	-376.86	-384.6059	-0.04183	0.1282
5	9.814	2141.20	0.0	2141.20	1756.5941	-0.0131	0.1151
6	9.836	0.0	1287.8	-1287.8	468.7941	0.0387	0.1538
7	10.79	22434.72	22434.72	0.0	468.7941	0.4472	0.6010
8	15.77	0.0	2873.48	-2873.48	-2404.6859	2.3346	2.9356
9	16.96	2396.94	0.0	2396.94	-7.7459	-2.8616	0.0740
10	26.52	515.15	0.0	515.15	507.4041	-0.0741	0.000
11	45.43	276.61	0.0	276.61	784.0141	9.5950	9.5950
12	53.16	97.90	0.0	97.90	881.9141	6.0604	15.6550
13	58.22	122.60	0.0	122.60	FD = 1004.5141	4.4625	20.1179
14	100					41.9686	62.0860

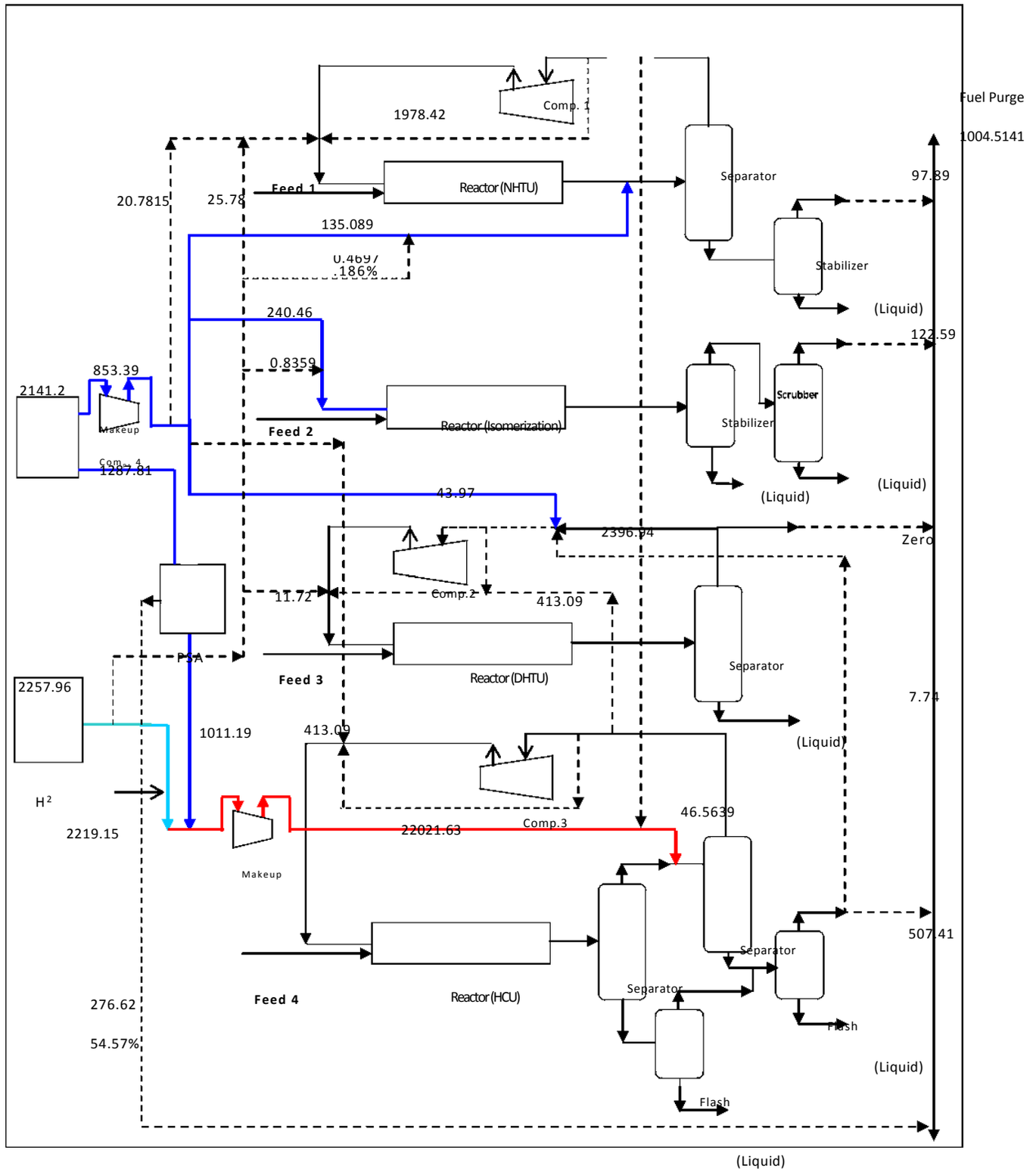


Figure 8. Hydrogen Integration Network in the Case Study by Cascade Technique.

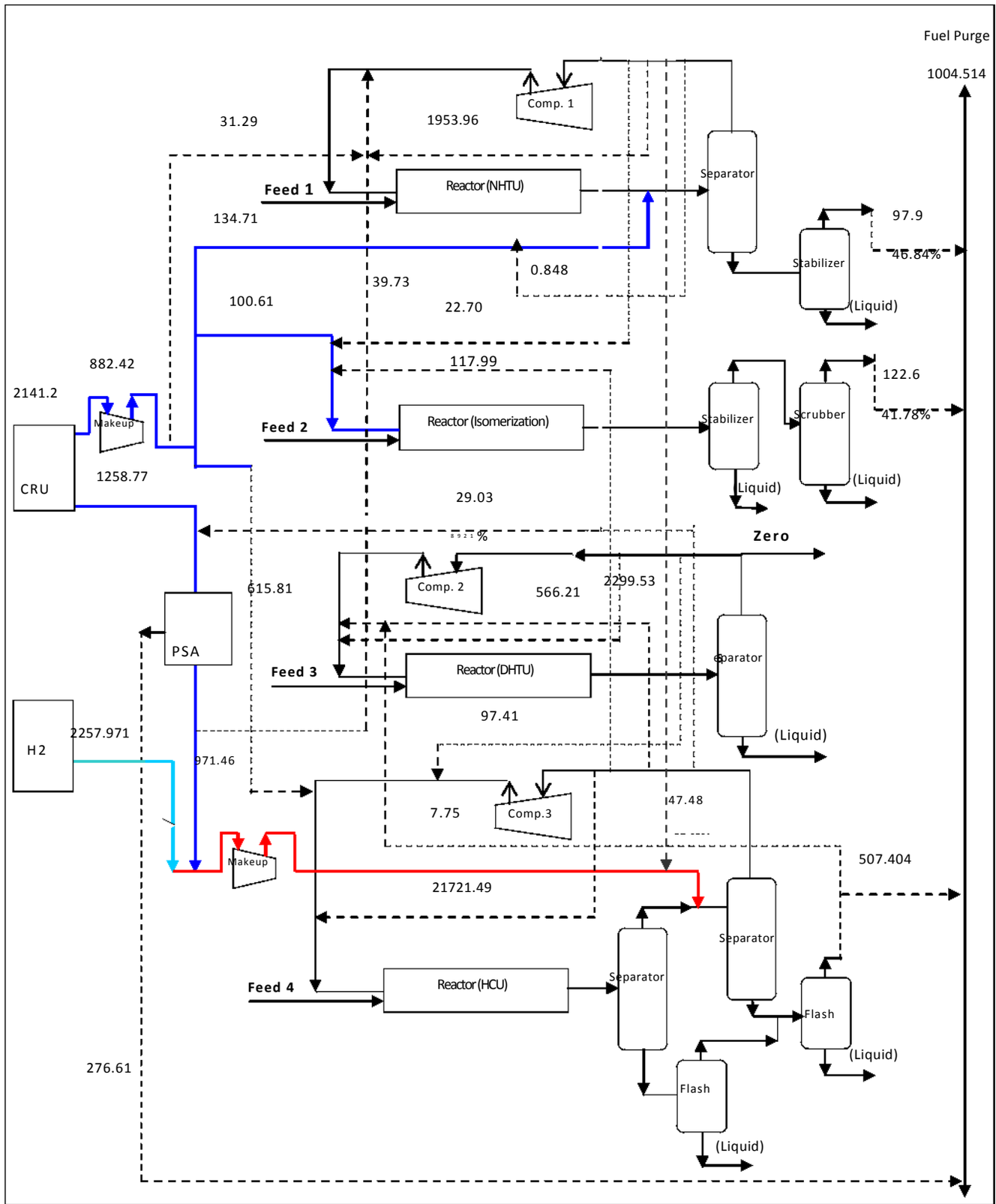


Figure 9. Hydrogen Integration Network in the Case Study by Mathematical Technique.

Table 5: Cascade Table for the hydrogen network in the case study (Regeneration with Membrane).

Level	Concentration Mole%	$F_{\text{in,mem}}$ Kmole/hr	$F_{\text{out,mem}}$ Kmole/hr	$\sum F_i - \sum F_j$ Kmole/hr	F_c Kmole/hr	Δm Kmole/hr	Cum Δm Kmole/hr
1	0.01	0.0	0.0	$F_i = 2163.216$	0.0 2163.216		0.0
2	0.1	1011.19	3276.9	-2265.71	-102.4940	0.1947	0.1947
3	2	808.61	0.0	808.61	706.1160	-0.1947	0.000
4	4.38	2024.99	2024.99	0.0	706.1160	1.6806	1.6806
5	9.78	0.0	376.86	-376.86	329.2560	3.8130	5.4936
6	9.814	2141.2	0.0	2141.2	2470.4560	0.0112	5.5048
7	9.836	0.0	1287.8	-1287.8	1182.6560	0.0544	5.5592
8	10.79	22434.72	22434.72	0.0	1182.6560 -	1.1283	6.6875
9	15.77	0.0	2873.48	-2873.48	-1690.8240	5.8896 -	12.5771
10	16.96	1392.43	0.0	1392.42	-298.4040	-2.0121	10.5650
11	26.52	515.15	0.0	515.15	216.7460	2.8527	13.4177
12	45.43	276.61	0.0	276.61	493.3560	4.0987	17.5164
13	53.16	97.9	0.0	97.9	591.2650	3.8136	21.3300
14	58.22	122.6	0.0	122.6	713.8560	2.9918	24.3218
15	78.71	195.9	0.0	195.9	FD = 909.7560	14.6269	38.9487
16	100					19.3687	58.3174

Table 6: Cascade Table for the hydrogen network in the case study (Regeneration with PSA).

Level	Concentration Mole%	F_{inflow} Kmole/hr	F_{outlet} Kmole/hr	$\sum F_i - \sum F_j$ Kmole/hr	F_c Kmole/hr	Δm Kmole/hr	Cum Δm Kmole/hr
1	0.01	0.0	0.0	$F_1 = 1926.35$	0.0 1926.35		0.0
2	0.1	1762.68	3276.9	-1514.22	412.13	0.1737	0.1737
3	4.38	2024.99	2024.99	0.0	412.13	1.7639	1.9376
4	9.78	0.0	376.86	-376.86	35.27	2.2255	4.1631
5	9.814	2141.2	0.0	2141.2	2176.47	0.0012	4.1643
6	9.836	0.0	1287.8	-1287.8	888.67	0.0479	4.2122
7	10.79	22434.72	22434.72	0.0	888.67	0.8478	5.0600
8	15.77	0.0	2873.48	-2873.48	-1984.81	4.4256	9.4856
9	16.96	1392.43	0.0	1392.43	-592.38	-2.3619	7.1237
10	26.52	515.15	0.0	515.15	-77.23	-5.6632	1.4605
11	45.43	276.61	0.0	276.61	199.38	-1.4605	0.000
12	53.16	97.9	0.0	97.9	297.28	1.5412	1.5412
13	58.22	122.6	0.0	122.6	419.88	1.5042	3.0454
14	67.03	253.03	0.0	253.03	FD = 672.91	3.6991	6.7445
15	100					22.1858	28.9303

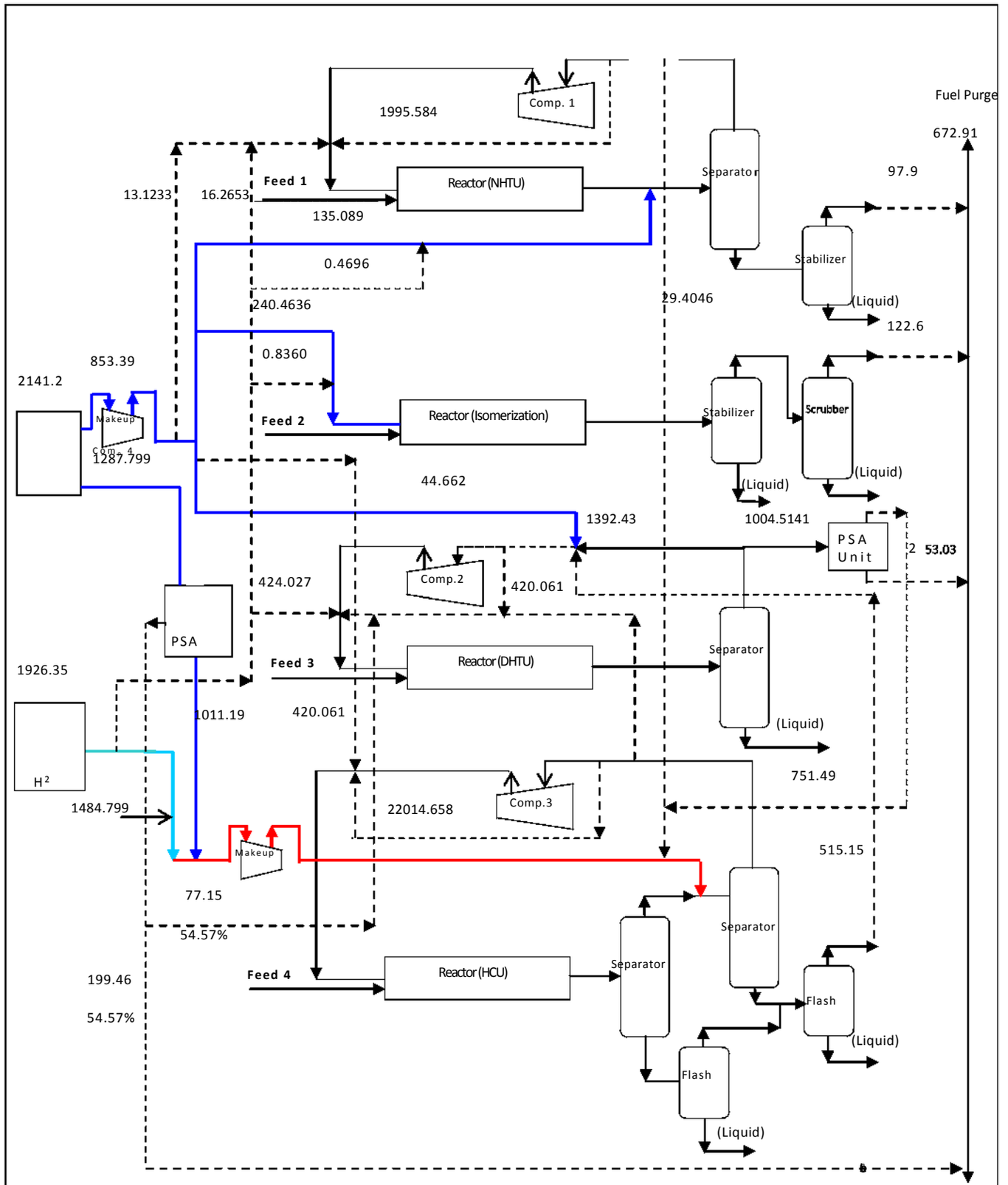


Figure 10. Hydrogen Integration Network with adding a PSA Unit by Cascade Technique.

Table 7: Cost Calculation Results.

Items	Existing Network Cost (MUS\$/yr)	Network integration Cost without adding any equipments	Network integration Cost with adding a membrane (MUS\$/yr)	Network integration Cost with adding a PSA (MUS\$/yr)
Hydrogen Cost	33.336	MUS\$/y 33.25	31.86	28.369
Power Cost	2.166	2.166	2.166	2.166
Fuel Cost	- 9.570	- 9.510	- 9.340	- 7.845
Total Operating Cost	25.932	25.906	24.686	22.69
Operating Cost saving	Base	0.026	1.246	3.242
Capital Cost	-	-	-	7.51 MUS\$
Payback period	-	-	-	2.316 year
Total Annual Cost	-	-	-	24.49 MUS\$/yr

Conclusion

- The two techniques, cascade and mathematical technique gave results very near to each other.
- In the integration of Medor hydrogen network without adding any new units and applying the cascade technique or the mathematical technique, it is found that, the saving in the fresh hydrogen is 0.25% and the saving in the hydrogen discharge is 0.77%.
- When adding a membrane as a purification unit and applying the cascade technique or mathematical technique, it is found that, the saving in the fresh hydrogen is 4.44% and the saving in the hydrogen discharge is 9.43%.

- When adding a PSA as a purification unit and applying the cascade or mathematical technique it is found that the saving in the fresh hydrogen is 14.9% and the saving in the hydrogen discharge is 33.0%.
- The saving in the operating cost is 3.242 MUS\$/year achieved by using the PSA purifier and the saving of the operating cost is 1.246 MUS\$/year achieved by using a membrane.

Nomenclature

CRU = Catalytic reforming unit
 HCU = Hydrocracking unit
 DHT = Diesel Hydrotreating
 KHT = Kerosene Hydrotreater
 CNHT = Cracked naphtha hydrotreater
 NHT = Naphtha hydrotreater
 HDA = Hydrodealkylation
 Fuel = Fuel gas system
 PSA = Pressure swing adsorption
 MMSCFD = Million standard cubic feet per day

F_{Fuel} = Fuel flow rate
 F_M = Hydrogen make-up flow rate
 Y_M = Hydrogen make-up composition
 F_R = Hydrogen recycle flow rate
 Y_R = Hydrogen recycle composition
 F_P = Hydrogen purge flow rate
 Y_P = Hydrogen purge composition
 F_D = Discharge hydrogen flow rate
 F_F = Fresh Hydrogen flow rate
 C_k = Concentration level
 C_F = Concentration of fresh hydrogen
 $F_{i,\text{pur}}$ = Flow rate of purifier
 $F_{\text{prod},j}$ = Flow rate of product stream of the purifier
 F_{residual} = Flow rate of residual stream of the purifier
 Y_{prod} = Purity of product stream of the purifier
 R = Hydrogen recovery of the purifier
 i,j = From source i to sink j
 w_i = Flow rate of source stream
 $w_{i,j}$ = Flow rate from source i to sink j

F_j = Flow rate of fresh hydrogen to sink j
 z_j = Composition of sink j after mixing of sources fractions and fresh fractions
 $N_{sources}$ = Total number of source streams
 N_{sinks} = Total number of sink streams
 G_j = Flow rate of sink j after mixing of sources fractions and fresh fractions
 $W_{i,waste}$ = Flow rate from source i to waste
 T_{AC} = Total annual cost
 Cap_{PSA} = Capital cost of PSA
 A_f = Annualizing factor
 f_i = the fractional interest
 n_y = number of the years
 LHV = Lower heating value
 $MUS\$$ = Million American dollar
 F_{in} = inlet flow rate to purifier
 P_{wr} = Compressor power
 $Comp.$ = Compressor

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