

Supporting Conversion Section of Hydrogen Production Unit by Membrane Modules to Produce Extra-Pure Hydrogen and Improving Equilibrium Conversion

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Abstract: The main object of this research is the modification of an industrial hydrogen production unit with palladium-based membrane modules to produce extra-pure hydrogen and shift reactions toward the hydrogen production side. The considered hydrogen production unit includes steam reformer, high and low temperature shift converters, carbon dioxide absorption tower, and methanator. The membrane modules are applied in the catalytic reactors and hydrogen is simultaneously penetrated from the reaction zone toward the sweep gas. In the first step, both conventional and membrane-supported processes are heterogeneously models based on the mass and energy balance equations at steady state condition. Then, the simulation results of conventional process are compared with the plant data to prove the validity of the developed model. Finally, the simulation results of conventional and membrane-supported processes are compared under the same operating condition. In general, applying the membrane module on the system increases hydrogen production rate from 63.95 to 67.21 mole s⁻¹. Based on the simulation results, supporting the conventional with the membrane module increases hydrogen production rate by 5.1%.

Keywords: hydrogen, steam methane reforming, heterogeneous modeling, palladium-based membrane

Nomenclature

a_v	Specific surface area (m ² m ⁻³)
A_C	Cross-sectional area (m ²)
C	Concentration (mole m ⁻³)
D_b	Bed diameter (m)
d	Particle diameter (m)
$D_{\it ef}$	Effective diffusion coefficient (m s ⁻²)
D_{ij}	Binary diffusion coefficient (m s ⁻²)
E_{j}	Activation energy of reaction j (kJ mol ⁻¹)
F_p	Shape factor
h_f	Heat transfer across particle (W m ⁻² K ⁻¹)
h_1	Liquid hold up on each tray (m ³)
h_f	Heat transfer in furnace (W m ⁻² K ⁻¹)
H_n	Enthalpy of stream exit n th tray (J mol ⁻¹)
ΔH_{i}	Heat of reaction j (J)
K_{eq}	Equilibrium constant
K_i	Adsorption constant of species i
k_{i}	Rate constant of reaction j (kmol kg _{cat} - l h ⁻¹)
k_{gi}	Mass transfer diffusion in porous media (m s ⁻¹)
K_{value}	Phase equilibrium constant
L_n	Liquid flow rate (mole s ⁻¹)

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Molecular weight (kg kmol<sup>-1</sup>)
M_{\rm w}
P
                 Pressure (bar)
                 Rate of reaction (kmol kg<sub>cat</sub><sup>-3</sup> h<sup>-1</sup>)
r_i
                 Reynolds number
Re
T
                 temperature (K)
                 Superficial velocity (m s<sup>-1</sup>)
u_{\varphi}
                 Overall heat transfer in fire box (W m<sup>-2</sup> K<sup>-1</sup>)
U_o
                 Vapor flow rate (mole s<sup>-1</sup>)
V_n
                 Liquid composition
X_{i,n}
                 Vapor composition
y_{i,n}
                 Axial coordinate of bed (m)
Greek Letter
                 Gas density (Kg m<sup>-3</sup>)
ρ
                 Bulk density (Kg m<sup>-3</sup>)
\rho_B
                 Bed porosity
\varepsilon_B
                 Viscosity (Pa s)
μ
                 Effectiveness factor
η
                 Emissivity
\in
λ
                Gas thermal conductivity (W m<sup>-1</sup> K<sup>-1</sup>)
                 Thiele modulus
φ
Subscript
                 Furnace
                 Gas
g
                 Catalyst surface
                 Wall
Abbreviate
                 Low Temperature Shift Reactor
LTSR
HTSR
                 High Temperature Shift Reactor
SMR
                 Steam Methane Reformer
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1. Introduction

Typically, hydrogen as the most abundant element on the earth plays a key role in the development of human societies. Ammonia, dimethyl ether, methanol, gasoline, and various aldehydes are catalytically produced from a mixture of hydrogen and carbon monoxide. In the oil refinery units, hydrogen is used as the hydrogenation agent in the hydrocracking, reforming, and hydrotreating units. Besides, hydrogen is directly applied in the power plants^[1], fuel cells^[2], aerospace^[3], semiconductors^[4], and automotive industries^[5]. Due to the high global demand for hydrogen, different routes, feedstocks, and technologies have been developed to produce hydrogen^[6]. Reforming and partial oxidation of hydrocarbons, coal gasification, electrolysis of alkaline solutions, and fermentation of organic substrates are common technologies developed to produce hydrogen. In the reforming process, the hydrocarbons react with a reforming agent in the presence of a heterogeneous catalyst, and hydrogen, carbon monoxide, and carbon dioxide are produced. Since pure hydrogen is applied in the fossil fuel upgrading units in an oil refinery, the steam reforming of methane is more attractive compared to other reforming types due to high hydrogen yield^[7]. In the old technologies, the produced carbon monoxide through reforming is converted to hydrogen in the high and low temperature shift reactors, and then the produced carbon dioxide is separated from products in an absorption column^[8]. In the new technologies, a part of carbon monoxide is converted to hydrogen in the high temperature shift converter, and the outlet stream is purified in a PSA unit to produce pure hydrogen^[9]. Dincer and Acer presented a good review on the hydrogen production routes and investigated the environmental impact, operating and investment costs, and efficiency of each method^[10].

Between developed technologies to integrate reaction and separation units including sorption enhanced and membrane separation, the membrane modules are more efficient due to low operating cost, complexity, and high flexibility^[11]. Many researchers have focused on the effects of supporting reforming and shift reactors by membrane modules on the operating conditions, hydrogen production, purity, and methane conversion. Roses et al. supported a steam methane reformer with a hydrogen permselective membrane module and investigated the effect of operating conditions on the process

performance^[12]. Although increasing operating pressure decreases the conversion based on Le Chatelier's principle, it enhances the equilibrium conversion by decreasing the hydrogen concentration in the system. Silva et al. formulated a multi-objective optimization problem to calculate the optimal operating conditions of a membrane supported reformer considering methane conversion and hydrogen recovery as the objective functions^[13]. Applying the optimal condition on the reformer increased the methane conversion and hydrogen recovery to 99.9% and 98.2%, respectively. Farsi and Shahosseini modified a membrane supported steam reformer by considering some quench points on the reactor body^[14]. The results showed that the proposed strategy could decrease pressure drop and increase the hydrogen production rate. Shigarov et al. reviewed the experimental and theoretical efforts on the application of Pd based membranes in catalytic reactors to produce hydrogen through steam methane reforming^[15].

In the hydrogen unit, produced carbon monoxide is catalytically converted to hydrogen and carbon dioxide in the shift converter. Mendes et al. presented a comprehensive review on the shift reaction and investigated the effects of supporting shift reactors with the membrane module on the process performance^[16]. Boutikos and Nikolakis proved the superior performance of carbon dioxide removal compared to the hydrogen separation in the shift reactor when the feed stream is carbon dioxide-rich^[17]. In the old technologies, the outlet product from the separation section feeds to the methanator to control hydrogen purity. Ohya et al investigated the effect of steam removal in the methanator by membrane module on the equilibrium conversion of reactions. The results showed that the steam removal by using the membrane module enhanced the equilibrium conversion when the inlet stream to the methanator was CO and CO₂ diluted^[18].

The main object of the present research is supporting the reformer, low and high temperature shift reactors, and methanator in an industrial hydrogen production unit with the palladium-based membrane modules to produce ultrapure hydrogen and increasing production capacity by shifting the equilibrium limitations. In general, the palladium-based membranes are used in hydrogen extraction because of their high permeability, selectivity, and good surface properties^[19]. In this regard, the proposed membrane supported and conventional processes are modeled based on the mass and energy balance equation at steady state condition. Then, the performance of the membrane supported configuration is compared with the conventional process.

2. Process description

2.1 Conventional design

Figure 1 illustrates the process flow diagram of the hydrogen production unit in a domestic oil refinery^[8]. In the first step, the chlorine and sulfur contaminants in the feed stream are removed in the desulphurization and dechlorination guard reactors. Then, the preheated feed stream mixed with superheated steam at the desired ratio of 7.06. The prepared mixture feeds to the coil placed in the radiation part of furnace. During steam reforming, methane is converted to carbon dioxide, carbon monoxide, and hydrogen over the Ni-Al₂O₃ catalyst. In the second step, the temperature of outlet product from the reformer is regulated in a cooler and feeds to the shift converters. In the shift reactors, the carbon monoxide and steam are catalytically converted to hydrogen and carbon dioxide. Afterward, the outlet stream from shift converters feeds to an amine absorption column to regulate carbon dioxide concentration in the product. Finally, the concentrations of carbon dioxide and carbon monoxide in the product stream decrease to an acceptable level in the methanator. Table 1 shows the reactor design data, catalyst specification, and operating conditions in the hydrogen unit.

Table 1. The reactor, catalyst design data and feed operating conditions in the hydrogen production unit

		Read	etor	Catalyst	Fee	d
	Reactions (Number)	Diameter (m)	Length (m)	Diameter (mm)	Temperature (K)	Pressure (bar)
Reformer	Reforming (1-3)	0.12	11.43	16	809	19
HTSR	Shift conversion (9)	2.45	3.2	8.47	623	15
LTSR	Shift reactions (9)	2.45	3.440	4.36	487	12
Methanator	Methanation (1-3)	1.7	2.895	5.4	523	15

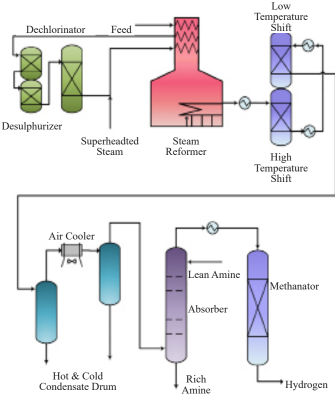


Figure 1. The schematic diagram of considered hydrogen production unit in Shiraz Oil Refinery

2.2 Modified design

Typically, the performance of industrial reformers and shift converters could be improved through the separation of hydrogen from the reaction zone^[20-21]. To support the conventional reformer with the membrane modules, a hydrogen permselective tube is placed in the coil tube and the annulus space between tubes is packed with the catalyst. The sweep gas flows in the inner tube and hydrogen penetrates from the reaction zone to the sweep gas gradually. Also, the low and high temperature shift reactors are equipped with membrane tubes. The considered low and high temperature shift reactors are two shell and tube exchangers that the catalyst is packed in the shell side of reactors, while the sweep gas flows in the hydrogen permselective tubes. Although steam removal from the reaction zone has a significant effect on the equilibrium conversion in the methanator^[18], the main idea to support the methanator by hydrogen permselective membrane module is producing the extra-pure hydrogen.

3. Reactions and kinetics

3.1 Reforming reactions

Typically, the nickel oxide supported alumina catalyst is used in the industrial reformer to convert methane to syngas. Kinetics of reforming and water gas shift reactions are as:

$$CH_{4(g)} + H_2O_{(g)} \leftrightarrow CO_{(g)} + 3H_{2(g)} \tag{1}$$

$$CH_{4(g)} + 2H_2O_{(g)} \leftrightarrow CO_{2(g)} + 4H_{2(g)}$$
 (2)

$$CO_{(g)} + H_2O_{(g)} \leftrightarrow CO_{2(g)} + H_{2(g)} \tag{3}$$

The rate of reforming reactions on the commercial catalyst are as [14]:

$$r_{1} = \frac{\left(\frac{k_{1}}{P_{H_{2}}^{2.5}}\right) \left(P_{CH_{4}}P_{H_{2}O} - \frac{P_{H_{2}}^{3}P_{CO}}{K_{1}}\right)}{\left(1 + K_{CO}P_{CO} + K_{H_{2}}P_{H_{2}} + K_{CH_{4}}P_{CH_{4}} + \frac{K_{H_{2}O}P_{H_{2}O}}{P_{H_{2}}}\right)^{2}}$$
(4)

$$r_{2} = \frac{\left(\frac{k_{2}}{P_{H_{2}}^{3.5}}\right) \left(P_{CH_{4}}P_{H_{2}O}^{2} - \frac{P_{H_{2}}^{4}P_{CO_{2}}}{K_{2}}\right)}{\left(1 + K_{CO}P_{CO} + K_{H_{2}}P_{H_{2}} + K_{CH_{4}}P_{CH_{4}} + \frac{K_{H_{2}O}P_{H_{2}O}}{P_{H_{2}}}\right)^{2}}$$
(5)

$$r_{3} = \frac{\left(\frac{k_{3}}{P_{H_{2}}}\right)\left(P_{CO}P_{H_{2}O} - \frac{P_{H_{2}}P_{CO_{2}}}{K_{3}}\right)}{\left(1 + K_{CO}P_{CO} + K_{H_{2}}P_{H_{2}} + K_{CH_{4}}P_{CH_{4}} + \frac{K_{H_{2}O}P_{H_{2}O}}{P_{H_{2}}}\right)^{2}}$$
(6)

3.2 Shift reaction

Industrially, the iron-chrome oxide and copper-zinc oxide catalysts are used in the high and low temperature shift converters, respectively. In the shift converter, carbon monoxide and steam react over the catalyst, and hydrogen is produced. The kinetic of shift reaction is as:

$$CO_{(g)} + H_2O_{(g)} \leftrightarrow CO_{2(g)} + H_{2(g)}$$
 (7)

The rates of shift reactions at high and low temperatures are as [22-23]:

$$r_{High} = k \left(P_{CO} P_{H_2O} - \frac{P_{CO_2} P_{H_2}}{K_{eq}} \right) \tag{8}$$

$$r_{Low} = k \frac{\left(P_{H_2O} - \frac{P_{CO_2} P_{H_2}}{K_{eq} \times P_{CO}}\right)}{\left(1 + \frac{K_{CO_2} P_{CO_2}}{P_{CO}}\right)}$$
(9)

3.3 CO₂ absorption

As expressed in the literature, CO_2 absorption in the aqueous solution of monoethanolamine (MEA) started with the formation of carbamate according to the zwitterion mechanism, followed by the hydration of CO_2 to form HCO_3^-/CO_3^{2-} , and accompanied by the hydrolysis of carbamate^[24]. The elementary rate of reaction is as [25]:

$$r_{CO_2} = k \ C_{CO_2} C_{MEA} \tag{10}$$

3.4 Methanation reaction

In the methanator, the reforming reactions occur in the reverse direction on the nickel oxide supported alumina catalyst. Thus, the kinetic and intrinsic rate of reactions in the methanator and reformer are similar^[26].

4. Process modeling

In this section, both conventional and membrane supported reactors are modeled based on the mass and energy

balance equations at steady state condition. The applied assumptions in the considered model are:

- The ideal gas law is acceptable because the compressibility factor is in range 0.99-1.0.
- The flow regime is plug due to the high Reynolds number.
- Mass and energy diffusions are negligible compared to convections^[27].
- The temperature gradient in the catalyst is negligible due to the small Biot Number^[28].
- The Palladium based membrane module is completely selective to hydrogen

4.1 Reforming reactor

In this part, the considered model to simulate the membrane-supported reformer is presented at steady state condition. Based on the considered assumptions, the mole balance equation in the gas and solid phases are as:

$$-u_g \frac{dC_i^g}{dz} - a_v k_{gi} \left(C_i^g - C_i^s \right) - \frac{P_m}{A_c} J_i = 0 \tag{11}$$

$$a_{\nu}k_{gi}(C_i^g - C_i^s) + \rho_B \sum_{j=1}^N \eta_j r_{ij} = 0$$
(12)

The Ergun equation is applied to calculate the pressure drop along the coil tube^[29]:

$$\frac{dp}{dz} = \frac{150\mu_g (1 - \varepsilon_b)}{d_p^2 \varepsilon_b^3} u_g + \frac{1.75(1 - \varepsilon_b)\rho_g}{d_p \varepsilon_b^3} u_g^2 \tag{13}$$

The energy balance equations in the gas and solid phases of the coil are as:

$$-u_{g}C_{P}C_{t}\frac{dT_{g}}{dz}-a_{v}h_{f}(T_{g}-T_{s})+\frac{U_{f}P}{A_{c}}(T_{f}-T_{g})-\frac{U_{m}P_{m}}{A_{c}}(T_{g}-T_{m})=0$$
(14)

$$a_{v}h_{f}(T_{g}-T_{s})+\rho_{B}\sum_{j=1}^{N}\eta_{j}r_{ij}(-\Delta H_{j})=0$$
(15)

4.2 Shift converters and methanator

In this part, the developed model to simulate the shift and methanation reactors are presented at steady state condition. The mole balance equations in the gas and solid phases are as follows:

$$-u_g \frac{dC_i^g}{dz} - a_v k_{gi} \left(C_i^g - C_i^s \right) - \frac{P_m}{A_c} J_i = 0$$

$$\tag{16}$$

$$a_{\nu}k_{gi}(C_i^g - C_i^s) + \rho_B \sum_{j=1}^N \eta_j r_{ij} = 0$$
(17)

The energy balance equations in both gas and solid phases are as follows:

$$-u_{g}C_{p}C_{t}\frac{dT_{g}}{dz}-a_{v}h_{f}(T_{g}-T_{s})-\frac{U_{m}P_{m}}{A_{c}}(T_{g}-T_{Perm})=0$$
(18)

$$a_{\nu}h_{f}(T_{g}-T_{s})+\rho_{B}\sum_{j=1}^{N}\eta_{j}r_{ij}(-\Delta H_{j})=0$$
(19)

4.3 Membrane module

The applied hydrogen permselective membrane modules in the processes could be classified into three main groups including ceramic, polymer, and metal membranes^[30]. Although the ceramic membranes present acceptable stability and strength, the low selectivity is the main disadvantage of those modules. Besides, the polymeric membranes are highly

selective and applied at low-temperature conditions due to low thermal and mechanical stability. Between synthesized membranes, the metal modules are wildly recommended to separate hydrogen from a gas mixture due to high thermal, mechanical and chemical stability and selectivity. In this research, due to high temperature and severe conditions, the Pd-Ag membrane module is selected to remove hydrogen from the reaction zone in the reformer, shift converter, and methanator. The mass and energy balance equations on the membrane modules are as:

$$-u_m \frac{dC_i^m}{dz} + \frac{J_i}{A_m} P_m = 0 \tag{20}$$

$$-u_{m}C_{p}C_{t}\frac{dT_{m}}{dz} + \frac{U_{m}P_{m}}{A_{m}}(T_{g} - T_{Perm}) = 0$$
(21)

The rate of hydrogen permeation through the Pd-Ag membrane module is calculated by [31]:

$$J_h = 6.33 \times 10^{-8} \frac{2\pi e^{\frac{-E_p}{RT}}}{\ln(R_o/R_i)} (\sqrt{p_h^R} - \sqrt{p_h^S})$$
(22)

4.4 CO₂ absorption column

Generally, the equilibrium and rate-based models are used to simulate the absorption columns^[32]. In the equilibrium models, it is assumed that the rate of mass transfer between gas and liquid streams are high and the outlet streams are at the thermodynamic equilibrium. In the rate-based models, the rate of mass transfer between phases is low and calculated with the mass transfer correlations. In this work, the CO₂ absorber column is simulated based on the mass and energy balance equations considering the equilibrium approach. It is confirmed that due to the non-ideality of the liquid phase, the activity coefficient of components in the liquid is calculated with the Wilson model^[33]. The developed equation to simulate the absorption column are:

$$L_{n-1}x_{i,n-1} + V_{n+1}y_{i,n+1} - V_ny_{i,n} - L_nx_{i,n} + r_{ij}h_l = 0$$
(23)

$$L_{n-1}H_{n-1} + V_{n+1}H_{n+1} - L_nH_n - V_nH_n + \Delta H_j = 0$$
(24)

$$y_{i,n} = K_{value} x_{i,n} \tag{25}$$

$$\sum_{i=1}^{C} y_{i,n} = 1 \tag{26}$$

$$\sum_{i=1}^{C} x_{i,n} = 1 \tag{27}$$

5. Solution method

Generally, the produced heat in the furnace is indirectly transferred to the gas stream in the coil tubes. To simplicity, it is assumed that the conductive heat resistance in the coil material is negligible. In the CO₂ absorber, the mass and energy balance equations, solubility, and considered auxiliary equations develop a set of nonlinear algebraic equations. Also, the applied mass and energy balance equations on the reformer, shift converter, and methanator result in a set of ordinary differential equations. In this research, the proposed procedure by Sarkarzadeh et al. is applied to solve the developed equations^[8].

6. Results and discussion

6.1 Process performance

In this section, the simulation results of conventional process are compared with the plant data to prove the validity of the developed model. Table 2 presents the relative absolute error of simulation results. It appears that the obtained model is

capable to predict the performance of considered industrial hydrogen units.

Table 2. Comparison	between sir	nulation resu	ilts and plant dat	a

		Simulation	Plant Data	Absolute Relative Error
Reformer	H ₂ Mole Fraction	74.95	74.7	0.33 %
	CH ₄ Mole Fraction	10.42	10.33	0.86 %
	HTSR Temperature [K]	647	654	1.10 %
	LTSR Temperature [K]	491.3	491	0.06 %
	Methanator Temperature	545.1	550	0.89 %

Based on the simulation results supporting the hydrogen production unit by the membrane module increases hydrogen production capacity by about 5.1% compared to the conventional process at the same feed condition. Typically, the hydrogen production rate in the conventional and membrane-supported units are 63.95 and 67.21 mole s⁻¹, respectively. Generally, the outlet hydrogen from the methanation reactor and separated hydrogen by membrane modules are 65.01 and 26.19 mole s⁻¹, respectively. Also, supporting the catalytic reactors by membrane modules increases methane conversion about 5.15% compared to the conventional process.

6.2 Steam methane reformer

In this part, the effect of supporting reformer by membrane module on the rate of hydrogen production is presented at steady state condition. Figure 2(a) and (b) show the molar flow rate of hydrogen and methane along the membrane supported and conventional reformers. According to Le Chatelier's principle, the equilibrium conversion of reforming reactions increases by decreasing hydrogen concentration in the reaction zone. Permeation of produced hydrogen in the reformer toward the sweep gas decreases hydrogen concentration in the reaction zone and shifts the reforming reactions toward the right side. Based on the simulation results, the produced hydrogen in the membrane supported and conventional reformers are 62.50 and 59.33 mole s⁻¹, respectively. Also, the methane conversion in the membrane supported and conventional reformers are 61.33% and 58.51%, respectively. Typically, supporting the steam methane reformer by membrane modules increases hydrogen production and methane conversion by about 5.34% and 4.82% at the same operating condition, respectively. Also, the molar flow rate of hydrogen at the outlet sweep gas from the membrane module is 6.53 mole s⁻¹.

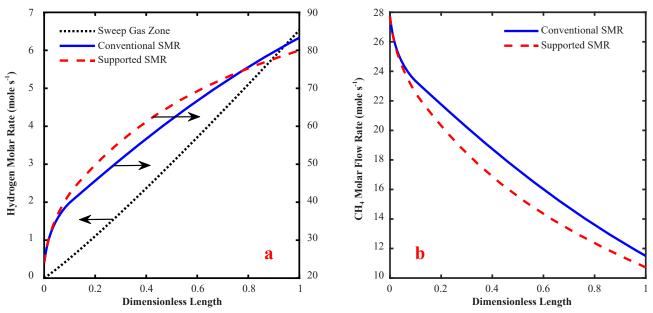


Figure 2. (a) Hydrogen and (b) methane flow rate along the reformer tube

Figure 3 shows the temperature trajectories along the membrane supported and conventional reformers at the same operating condition. Typically, the considered membrane reactor consists of two coaxial tubes that the annulus space

between tubes is packed with the catalyst. In the membrane configuration, the membrane module is surrounded by the reaction zone and the coil is placed in the firebox. As a base case, the sweep gas temperature is selected to be upper than the feed temperature. In this regard, the heat is transferred from sweep gas to the reaction zone in the first half of the membrane reactor and sweep gas temperature decreases gradually. While the heat transfer direction is reversed in the second part of the membrane supported reformer and sweep gas temperature increases gradually due to low heat consumption in the reaction zone. In general, decreasing methane concentration in the second part of reformer decreases the rate of methane conversion, hydrogen production, and heat consumption.

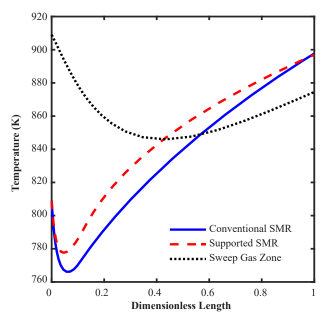
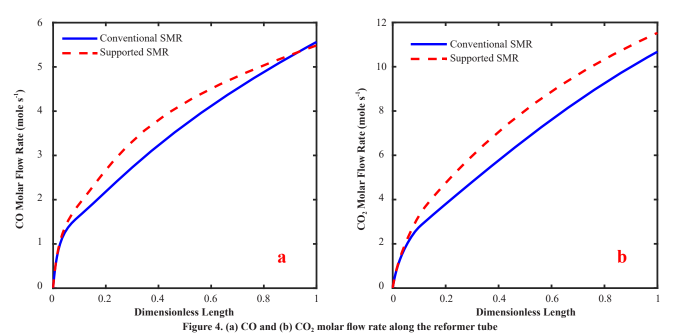


Figure 3. Temperature profile along the coil at the conventional and supported membrane conditions

Figure 4(a) and (b) show the molar flow rate of carbon monoxide and carbon dioxide along the membrane and conventional reformers, respectively. Based on the simulation results, increasing methane conversion in the membrane supported reformer increases carbon dioxide production from 10.68 to 11.53 mole s⁻¹.



6.3 High-temperature shift converter

In the industrial hydrogen unit, the outlet effluent from reformer is cooled down and feeds to the high temperature

shift reactor. In the shift converter, the carbon monoxide is converted to hydrogen and carbon dioxide. According to Le Chatelier's principle, hydrogen removal with the membrane module shifts the water gas shift reaction toward the carbon monoxide conversion. Figure 5(a-c) shows the profile of hydrogen flow rate, carbon monoxide flow rate, and operating temperature along the membrane supported and conventional high temperature shift converters. Based on the simulation results, carbon monoxide conversion in the membrane supported and conventional high temperature shift reactors are 81.32% and 73.60%, respectively. It appears that the heat transfer from sweep gas toward the reaction zone increases reaction temperature and enhances the reaction rate and conversion. Based on the simulation results, the produced hydrogen in the membrane supported and conventional converters are 4.46 and 4.09 mole s⁻¹, respectively. Besides, the molar flow rate of hydrogen at the outlet sweep gas from the membrane module is 7.55 mole s⁻¹.

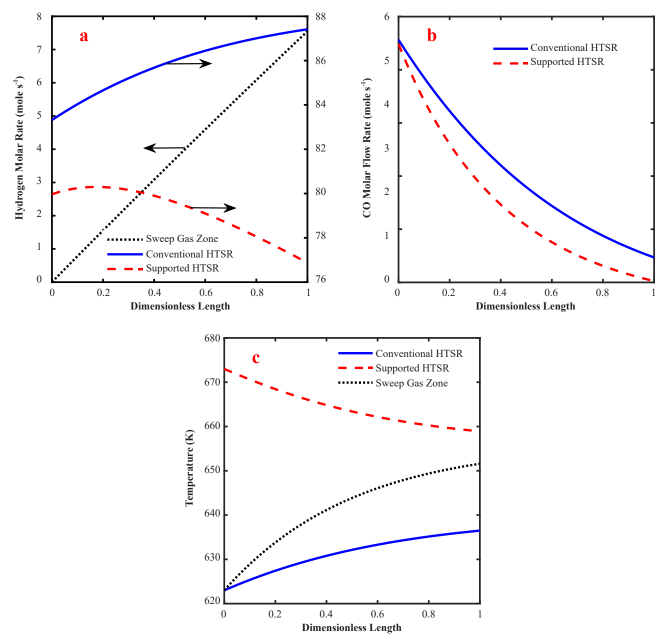


Figure 5. (a) Hydrogen, (b) CO molar flow rate (c) Temperature along the high temperature shift reactor

6.4 Low-temperature shift converter

The outlet stream from high temperature converter is cooled down in a heat exchanger and feeds to the low temperature water gas shift reactor. It appears that the low-temperature converter is under equilibrium control and decreasing operating temperature increases equilibrium conversion. As a base case, the temperature of inlet sweep gas to

the reactor is set to be lower than the feed temperature. Thus, heat transfer from the reaction zone toward the sweep gas could decrease the temperature of reaction zone and shift the reaction toward the hydrogen production. Figure 6(a-c) shows the profiles of hydrogen, carbon monoxide, and temperature along the membrane supported and conventional converters. Based on the simulation results, the molar flow rate of carbon monoxide in the outlet stream from the low shift reactor is 0.03 and 0.06 mole s⁻¹ in the membrane supported and conventional configurations, respectively. Typically, hydrogen production is dominant over the hydrogen permeation rate, and hydrogen concentration increases in the first half of membrane converter. Since the shift reaction approaches toward the equilibrium in the second half, the hydrogen permeation rate is dominant over the hydrogen production and hydrogen concentration decreases along the membrane reactor. Based on the simulation results, the hydrogen molar flow rate increases along the sweep gas and approaches to 6.602 mole s⁻¹.

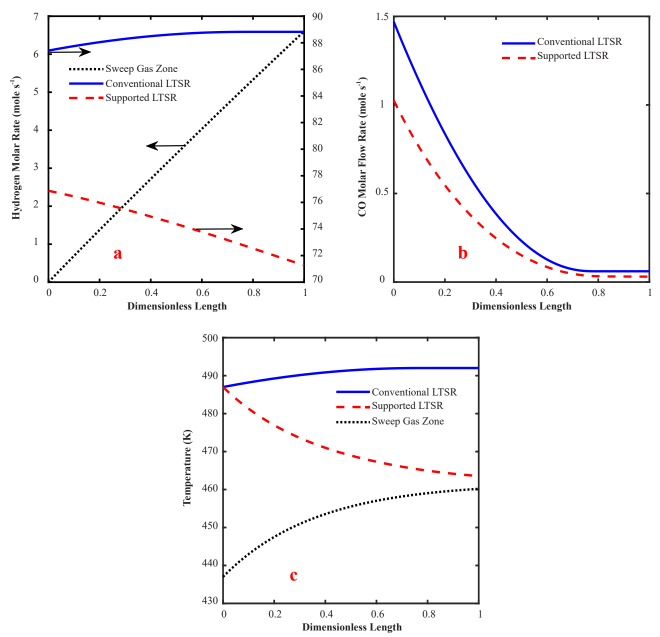


Figure 6. (a) Hydrogen, (b) CO molar flow rate (c) Temperature along the low temperature shift reactor

6.5 CO₂ absorber column

In the absorber, carbon dioxide is absorbed by the aqueous solution of monoethanolamine in a sieve tray column, and outlet stream from the top of absorber feeds to the methanator. Based on the simulation results, the molar flow rate of carbon dioxide in the inlet stream to the absorber is 16.18 and 16.98 mole s⁻¹ in the conventional and membrane supported

configurations, respectively. Besides, the amine-circulating rate in the membrane-supported configuration is 144.37 mole s⁻¹. Since the inlet stream to the absorption column in the membrane configuration is CO₂ rich, increasing the amine-circulating rate is necessary to control CO₂ concentration at the acceptable level. Figure 7 shows the molar flow rate of carbon dioxide along the absorption column in the conventional and membrane-supported processes.

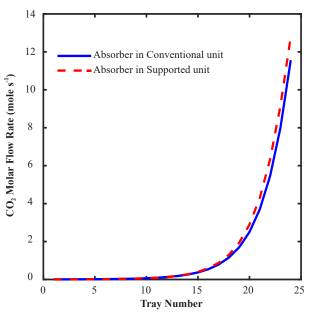


Figure 7. The carbon dioxide molar flow rate along the absorption column

6.6 Methanation reactor

In the methanation reactor, decreasing temperature and steam concentration shift the reforming reactions toward the left side and results in the hydrogenation of carbon dioxide and carbon monoxide. Typically, the main idea to support the methanator by hydrogen permselective membrane is the production of the extra-pure hydrogen. Since the inlet stream to the methanator is hydrogen-rich, hydrogen removal has an insignificant effect on the equilibrium conversion. Figure 8(a-c) shows the flow rate of hydrogen, carbon monoxide, and carbon dioxide along the conventional and membrane methanation reactors. Based on the simulation results, the hydrogen molar flow rate increases along the sweep gas and approaches to 5.50 mole s⁻¹. The CO molar flow rate in the outlet stream from the membrane supported reactor is 7.5×10^{-6} mole s⁻¹, while it approaches 9.4×10^{-6} in the conventional methanator. The CO₂ concentration in the outlet stream from both conventional and membrane methanation reactors is negligible. Generally, the lower carbon monoxide concentration in the inlet stream to the membrane-supported methanator decreases the rate of hydrogen consumption in the system and results in higher hydrogen concentration in the outlet stream.

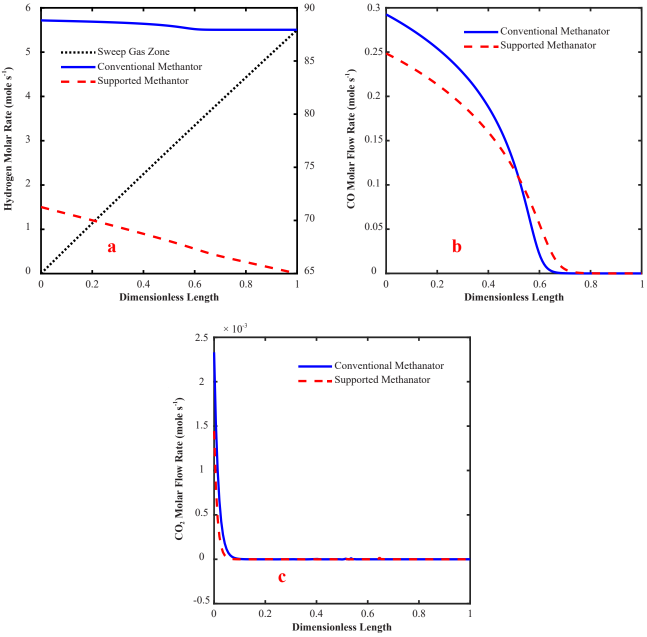


Figure 8. The molar flow rate of (a) hydrogen, (b) carbon monoxide and (c) carbon dioxide along the methanation reactor

7. Conclusion

In this paper, a hydrogen production unit in a domestic oil refinery, which comprises steam methane reformer, high and low temperature shift reactors, CO₂ absorber and methanator, was modified by hydrogen membrane modules to produce ultra-pure hydrogen. In this regard, palladium-based membrane modules were applied in the reactors. Both membrane supported and conventional processes were heterogeneously modeled based on the mass and energy balance equations at steady state condition. The simulation results showed that supporting the reactors by the membrane module improves the hydrogen production rate and methane conversion by 5.1% and 5.15%, respectively. The improvement in the hydrogen production rate and methane conversion in membrane supported reformer were 5.34% and 4.82%, respectively. Besides, supporting shift converters by membrane modules improved reaction rate and equilibrium conversion, and resulted in the higher carbon monoxide conversion to hydrogen. Briefly, the main benefits of proposed system were higher hydrogen production and methane conversion.

Conflict of interest

The authors declare no competing financial interest.

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