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Effect of SiO₂ Concentration and Time on Stability of TiO₂ Zeolite Nanocomposite Membrane in Light Gas Dehumidification

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ABSTRACT

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Keywords: Dehumidification SiO₂ Concentration Nanocomposite Membrane TiO₂ Zeolite Stability The main goal of this research is to use zeolite TiO₂ nanocomposite membranes in order to remove moisture from gas. For this reason, a certain TiZ-V membrane was selected and manufactured as a standard membrane, which was the result of the initial assessment of a suitable membrane for gas dehumidification, and this membrane was used as a standard to measure the effect of manufacturing parameters. The findings showed that increasing the concentration of SiO₂ had greatest effect on increasing the water flux of the membrane due to the effect of increasing the reaction time of the vapor phase carrier and reducing the selectivity drop at higher pressures. Also, the experiments of changing the relative humidity of the feed have shown the improving efficiency of the membrane in relative humidities lower than 80%, so that under the conditions of lower relative humidity, the selectivity of the membrane has increased. Another positive point found is a slight change in the selectivity efficiency of the membrane with respect to different relative humidities. This case showed the stability of membrane performance under the different conditions of humidity of the feed gas. Next, in order to increase the performance of membrane as much as possible, the sweeper gas was added from the inside of membrane. Increase in the sweeper gas, which increases the water concentration gradient and decreases the gas concentration gradient on the sides of membrane wall, increases the selectivity of membrane to the highest level of 543.

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NOM	ENCLATURE		Greek Symbols		
Р	bar	Sw	mL/s		
V	mL	$RH_R\%$	relative humidity of residual		
Т	°C	$RH_F\%$	relative humidity of feed		
Q	L/h	GPU	permeability		
а	selectivity	Р	mol/m².s.pa		
P_{avg}	ра	ρ	Density (kg/m ³)		
g	Gravity(m/s ²)	τ	Lattice relaxation time		
n	mol	Y^G	mole fraction of gas		
Y^w	mole fraction of water	t	time (min)		

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1. INTRODUCTION

Because natural gas is one of the sources of energy production in the transmission industry and domestic use, its production and distribution is very important (1). Natural gas is mainly composed from methane (80 to 90 percent), a small amount of ethane-propane and butane. In addition, gas compounds often include other compounds such as H₂O and CO₂ that the presence of these compounds in gas causes problems during transportation and consumption (2-4). Because natural gas often leaves the reservoirs as saturated from water, A slight change in gas pressure or temperature causes water condensation or formation of hydrates (5). Hydrates are compounds very similar to ice which are obtained from the bonds between water and ice molecules (6). The formation of hydrates during gas transmission causes blockage of the pipeline and consequently, permanent interruption of gas flows (7). In addition to hydrate formation, the presence of liquid water in pipelines can cause pipe corrosion; therefore, it is necessary to be removed from the composition of natural gas (8). Removal of moisture from natural gas is due to the need to prevent from reducing the calorific value of gas and purifying gas in order to achieve the necessary standards for delivering gas to Petrochemical industries and household uses (9, 10). Also, in the facilities department in the gas industry, the presence of water in gas pipelines can cause two main problems; the presence of small amounts of water along with the presence of some sour gases causes widespread acid corrosion in gas transmission pipelines. This issue would cause the imposition of costs and the possibility of risks due to gas leakage (11, 12).

Also, water is one of the main bases for the formation of gas hydrates (13), and formation of solid methane hydrates in pipes containing gas in different parts of the facility can change the flow pattern, pressure changes and also, with its rapid growth, it can stop the gas transmission operation due to the blockage of the pipe and the possibility of its bursting (14). As a result, dehumidification of natural gas from the formation of gas hydrates helps proper transmission and reduction of corrosion. There are various methods for dehumidifying natural gas on an industrial scale (15, 16), the most important of which are as follows:

- 1. Direct cooling
- 2. Surface adsorption
- 3. Absorption by liquid

Dehumidification of natural gas is done in two main approaches: Towers filled with surfactants and the wider method that is ethylene glycol dehumidification units (17).

The first method includes towers filled with solid surface absorbents such as silica gel or alumina (18), which have the problems as follows: The need for several towers (on average 3 or 4 towers); restriction of use due to weight and large dimensions, the need for permanent and accurate operation controls, extensive destruction and high price of absorbent, also this method often has the problems of ethylene glycol dehumidification units (19).

A more widespread method of dehumidification unit is tri-ethylene glycol (TEG) (20). This method only involves the use of two towers that is contacted with ethylene glycol which is a liquid of water absorbent in the gas absorption tower in the form of counter flow on the bubble cap trays and then in the solvent recycling operations disposal tower, is required to separate moisture from tri-ethylene glycol (21). Some of the main problems with glycolic dehumidifiers are:

- 1. Absorbent wide flammability; Result: safety issues.
- Complex unit operation in absorption and disposal towers; Result: high investment, operating and maintenance costs.
- 3. Energy supply of towers and facilities; Result: consuming part of gas as fuel.
- 4. Requiring solvent warehouse and storage, replacing, neutralizing and reproducing it; Result: the need for a large space.
- 5. Emission of hazardous volatile organic compounds (VOCs), especially benzene, toluene, ethylbenzene, and xylenes (BTEX); Result: release of 40 tons of VOCs per year from a typical dehumidification unit that half of which is BTEX (22-24).

If the goal is to dehumidify of large volume of gas to reach a dew point of 40 to 140 degrees Fahrenheit, liquid absorption systems are more economical (25). But if the dew point is above 180 degrees Fahrenheit, membrane absorption units are more suitable (26). Due to the disadvantages of moisture, extensive studies are carried out in the field of investigating the effect of various parameters such as the type of adsorbent, feed flow intensity, temperature and process pressure on the moisture absorption performance of the gas flow (27). Therefore, temperature, pressure, moisture concentration in feed, Intensity of feed flow (independent variable) as variables affecting moisture absorption performance in a nanostructure membrane system were investigated (23). In terms of the growing trend of problems caused by the method of dehumidification conventionally, the necessity of conducting research by promising technologies such as Nano is more apparent.

Among the various techniques, membrane filtration is a more effective and efficient one for separating water from gas, especially oil emulsions. The advantages of membrane separation are high selectivity, low maintenance cost, low required space, high efficiency, and no need for chemical additives (28).

Among the most important advantages of Nano composites are better mechanical properties compared to normal composites (reinforcement in Nano composites is more bidirectional than unidirectional), increasing thermal stability and self-extinguishing, preventing the penetration of gases such as oxygen, steam He mentioned water, helium, carbon dioxide and organic vapors such as ethyl acetate (due to the formation of a porous and tortuous path due to the presence of clay) (29).

Also, nanocomposites are lighter than normal composites, and this feature makes them competitive with other materials with special applications (29).

This study was conducted regarding the possibility of using TiO₂ zeolite nanocomposite membranes to remove moisture from gas with natural gas and nitrogen approach and about gas membrane dehumidification. In this way, the present work has been completely innovative in terms of the selected method and the type of surface material used in the fabrication of TiZ-V membrane on NAA zeolite hollow fibers.

2. MATERIALS AND METHODS

2.1. Synthesis of Preparation of Titanium Dioxide Nanowire by Eutectic Method Due to the methods of preparation of nanostructures and also checking the condition of the equipment and available facilities, eutectic method has been used for fabrication of titanium nano oxide. Because the selected equipment used in this method is relatively cheap, the need for low temperature in the process, low energy consumption and compatible with the environment. To prepare titanium dioxide One-dimensional nano structures (nanowire) by eutectic method, a mixture of disodium phosphate salts (Na₂HPO₄), sodium chloride (NaCl) and titanium dioxide raw material (TiO₂) with specific weight ratios (1:4:1)was prepared and ground in a mortar, then transferred to the furnace in the porcelain crucible and the reaction was performed at temperature 825 °C for 8 hours. After cooling, the crucible is removed from the furnace and then several times washing was done with boiled distilled water. During this step, the excess salts in the products were washed. At finally, the sediments from washing were dried in the furnace (Figure 1).



The numerical value of HLB plays an important role in deciding to choose the type of surfactant and indicates the polarity and non-polarity ratio of surfactant groups. To form a water-in-oil emulsion, HLB is needed between 5.3 to 8. But, SDS has a general mode and is used for the most tests. In order to prepare micellar solution, surfactant (SDS) and auxiliary surfactant (butanol) with a weight ratio of 2:1 are mixed with solvent (Hexane). Auxiliary surfactant is used to neutralize electrostatic charges caused by side co-positioning of the charged heads of the ionic surfactant and stabilizing the micelles. To dissolve SDS in hexane, the resulting mixture is kept night and day in a closed glass container and mixed by a magnetic stirrer to achieve complete uniformity. Then the composition of zeolite is prepared with the molar ratio of Al₃O₃, 2.1SiO₂, 3Na₂O, 150H₂O. In short, 2g of aluminum sulfate and 0.58g of sodium hydroxide is dissolved in 6.22g of deionized water and will be poured into syringe A. 1.41g of sodium silicate solution is poured into syringe B.

The contents of two syringes were added drop by drop to the micellar solution under ultra sonication. The weight ratio of mixing materials was 3H:1S:1Z, where H, S and Z represent solvent, the mixture of surfactants and aluminosilicate gel (the total weight of aluminum solution and silicate solution) respectively. Mixing of the ingredients continued for 30 minutes and then the resulting mixture was transferred to a steel reactor which its inner body was covered with Teflon and placed in an oven at a temperature of 75°C for 6 hours. After this period, the contents of the reactor were removed and centrifuged to separate zeolite particles from the water and oil mixture, which are two separated phases. Then zeolite particles were washed several times with water and acetone to reduce the pH of the water after washing. Then zeolite particles were placed inside the oven to dry at a temperature of 120°C for 12 hours. In order to calcine and remove the remaining surfactants, zeolite particles are placed in furnace at a temperature of 550°C for 5 hours.

2. 3. SiO₂ Deposition on TiO₂ In this step, 1g of particles (TiO₂) obtained from step 1 is dispersed in deionized water by a homogenizer and the resulting





Figure 1. Different stages of synthesis by Eutectic method

Figure 2. The chemical structure of sodium di-decyl sulfate

mixture is placed in a water bath at a temperature of 90°C. Then the amount of 3.7 g of sodium silicate (the weight ratio of $\frac{SiO_2}{TiO_2} = \frac{1}{1}$) is added to the solution. At this

time, 0.1M hydrochloric acid solution is added drop by drop to the solution until the pH of the solution reaches 0.5. Stirring of the solution continues for 1h and after this period, the solution is cooled at room temperature and the covered particles (TiO₂) with SiO₂ are separated with a sieve. (TiO₂) @SiO₂ particles were washed and dried at 50°C. These particles were called P–TiS.

2.4. P-TiS Covering with NaA Zeolite In order to coat P-TiS particles by zeolite, the vapor phase transport method (VPT) will be used, which is a method in synthesis of high silica zeolites. Matsukata et al. (31) reported that dry alum inosilicate gel was converted to MFI when exposed to water vapors and volatile amines. In fact, in this method, amorphous alum inosilicate gel is crystallized in contact with vapors of a structure-forming material without direct contact with it. In this research, this method was used to integrate the SiO₂ layer in zeolite. In short, the synthesized zeolite particles in step 2 were mixed with 30mL of toluene and placed in a closed container on a magnetic stirrer for one day and night. Then P-TiS particles were added to the mixture under vigorous stirring by a homogenizer. The weight ratio of P-TiS to zeolite is 1:1. Stirring was continued for 10 minutes. Then the mixture was placed statically to deposit TiO₂ particles covered with zeolite. The resulting particles were placed under vacuum and at ambient temperature for 2 hours to evaporate the remaining toluene. The resulting solid was transferred to a steel reactor with Teflon inner body. An overview of the used reactor can be seen in Figure 3. The particles were placed on the cylinder inside the reactor and a solution containing 4g of deionized water and 2g of trimethylamine was poured around it. Reactor was kept at a temperature of 140°C for 3 days. Then the resulting particles were removed from the reactor and dried at 40°C. These particles are named with TiZ-V (32).



Figure 3. Design of gas test device of the hollow fiber membrane

The entering gas from the cylinder enters to the chamber with a certain pressure. This gas goes from the side of the shell by passing through membrane wall to the inside of the hollow fiber and from there is directed to the bubble flow meter. According to the amount of permeated gas in the bubble flowmeter, gas permeability will be calculated (see Figure 4).

In order to determine the selectivity of membrane, the data obtained from the experiments are used as input for the calculations include; different pressures of the inlet gas (P[bar]), a certain volume of the bubble flow meter (V[mL]), The gas flow time in the bubble flow meter for a given volume (t[s]), ambient temperature and vapor generator tank temperature (T [°C]), relative humidity of feed RH_F, relative humidity of penetrant flow RH_P, relative humidity of residual RH_R and feed flow (L/h) which are used to measure the dehumidification performance of the constructed membrane. It should be noted that the considered standard operational conditions in performance tests include relative humidity of feed, 80%, the gas flow is 50 [L/h] and without sweep flow which, these conditions in all operations except those mentioned in the measurement section of the effect of change operating parameters are followed by default.

3. RESULTS AND DISCUSSIONS

3. 1. SiO₂ Concentration The effect of change the concentration of SiO_2 on the results of gas, water and performance tests are shown in Figures 5, 6 and 7, respectively.

Regarding the 1SiO_2 membrane, the water permeability of the membrane has decreased with a decrease in concentration of SiO_2 , which is also



Figure 4. Image of the constructed gas dehumidification membrane device



Figure 5. Diagram of the effect of change the *SiO*₂ concentration on gas test results



Figure 6. Diagram of the effect of change the SiO_2 concentration on the results of water test



Figure 7. Diagram of the effect of change the *SiO*₂ concentration on performance test results

considered an unfavorable situation. The decrease in concentration of the silicate phase in the SiO_2 aqueous solution has caused a decrease in the hydrophilicity of the membrane and, as a result, a decrease in the water permeability of the membrane. The results of $1SiO_2$ membrane efficiency test are presented in Table 1.

TABLE 1. Permeability results of zeolite membrane with
variations of $1SiO_2$ concentration

P(bar)	$P_{avg}(pa)$	N (mol)	$P(mol/m^2.s.pa)$
0/1	106325	-06E2/2442	-06E1/766
0/3	116325	-06E6/7191	-06E1/762
0/5	126325	-05E1/1174	-06E1/758
0/7	136325	-05E1/56294	-06E1/757
0/9	146325	-05E2/00279	-06E1/751
1/2	161325	-05E2/66429	-06E1/747
1/6	181325	-05E3/54392	-06E1/743
2	201325	-05E4/4189	-06E1/738
3	251325	-05E6/61227	-06E1/734
5	351325	0/000109702	-06E1/726
7	451325	0/000153287	-06E1/723

Regarding the $4SiO_2$ membrane, it shows an increase in water permeability of the membrane with the increase of SiO_2 concentration. Increasing the concentration of the silicate phase in the mixture of SiO_2 has increased the hydrophilicity of the membrane and, as a result, increased the water permeability of the membrane. The results of the performance test of $4SiO_2$ membrane are presented in Table 2.

3. 1. 1. The Effect of Decrease in the SiO_2 Concentration According to the comparison between the concentrations of SiO_2 in (Figure 7), respect to TiZ-V membrane, it is concluded that the decrease in the concentration of SiO_2 increased the gas permeability. An increase in gas permeability is considered as an unfavorable state because membrane must have minimum gas permeability and maximum water permeability. The cause of this phenomenon is due to the decrease in the concentration of SiO_2 and as a result, the failure to complete the cross-linking process on the

TABLE 2. Permeability results of zeolite membrane with variations of 4SiO₂ concentration

$P_{avg}(pa)$	n(mol)	$P(mol/m^2.s.pa)$
126325	-07E2/08963	-08E4/078
136325	-07E2/90998	-08E4/056
146325	-07E3/78164	-08E4/100
161325	-07E5/11107	-08E4/156
181325	-07E6/86339	-08E4/185
201325	-07E8/60661	-08E4/199
251325	-06E1/33895	-08E4/355
351325	-06E2/33977	-08E4/566
451325	-06E3/39365	-08E4/730

surface of membrane, which causes the cross-linking layer of membrane, especially at higher pressures (according to Figure 5) loses the possibility of resistance to prevent penetration of nitrogen molecules and then, the permeability of nitrogen gas increases.

Perusing the diagram in (Figure 6), regarding membrane $1SiO_2$, shows that the decrease in water permeability of membrane have been with the decrease in the concentration of SiO_2 . This is also considered as an unfavorable state. The decrease in the concentration of the silicate phase in the aqueous solution of SiO_2 has caused a decrease in the hydrophilicity of membrane and consequently, a decrease in the water permeability of membrane. Membrane performance test results of $1SiO_2$ is presented in Table 3.

According to Table 3, regarding the state of penetrant, it indicates that water component is lower compared to gas even in the lowest measured pressure. This state means an increase in gas permeability from the penetrant side, which means an increase in gas loss from the penetrant side in the form of wet gas which, corresponds to the result of gas test of this membrane. Also, the lower water content in the penetrant corresponds to the result of water test of this membrane, i.e., the decrease in water permeability. These two unfavorable states have caused a general decrease in the selectivity of the mixed gas of membrane. Relative humidity change diagram of membrane $1SiO_2$ is also presented in Figure 8. Finally, Figure 9 presents the gas and water permeability of the 1SiO₂ membrane at different pressures and their corresponding selectivity.

3. 1. 2. The Effect of Increase of the SiO_2 Concentration ($4SiO_2$ Membrane) According to the comparison between the concentrations of SiO_2 in (Figure 7), relative to TiZ-V membrane, it is concluded that the increase in the concentration of SiO_2 has caused a slight decrease in gas permeability. The increase in SiO_2 provides the required values for the reaction with the existing value of P-TiS, but its higher values will not cause more crosslinking of the network due to the no reaction with P-TiS; therefore, as the results show, the major no change in permeability values of the $4SiO_2$ membrane was due to the major no change in the crosslinking of the selective layer compared to TiZ-V membrane.

TABLE 3. Membrane performance test results of $1SiO_2$

p(bar)	n_F	n_p	Y^w	Y ^G	а
2	5.797E-04	1.120E-05	0.460	0.540	8.32
3	5.827E-04	1.41973E-05	0.395	0.605	0.25
4	5.877E-04	1.92646E-05	0.320	0.680	1.18
5	5.981E-04	1.96823E-05	0.251	0.749	8.12
6	6.197E-04	5.12518E-05	0.169	0.831	7.7



Figure 8. Diagram of changes in relative humidity of residue versus relative humidity of feed in the 1*SiO*₂ membrane



Figure 9. Diagram of gas and water permeability values of 1*SiO*₂ membrane and its corresponding selectivity

The comparison of gas permeability values in $1SiO_2$ membrane compared to $4SIO_2$ was done and data are summarized in Table 4.

Perusing the diagram in Figure 6, regarding to the $4SiO_2$ membrane, shows an increase in membrane water

TABLE 4. Comparison of gas permeability values of 1SiO2

 membrane compared to 4SiO2

P _{avg} (pa)	1SiO ₂ , P(mol /m ² .s.pa)	4SiO ₂ , P(mol /m ² .s.pa)	1SiO ₂ / 4SiO ₂ , P(mol/ m ² .s.pa)
126325	-06E1/758	-08E4/078	+01E4/31
136325	-06E1/757	-08E4/056	+01E4/33
146325	-06E1/751	-08E4/100	+01E4/27
161325	-06E1/747	-08E4/156	+01E4/20
181325	-06E1/743	-08E4/185	+01E4/16
201325	-06E1/738	-08E4/199	+01E4/14
251325	-06E1/734	-08E4/355	+01E3/98
351325	-06E1/726	-08E4/566	+01E3/78
451325	-06E1/723	-08E4/730	+01E3/64

permeability with increasing SiO_2 concentration. Increasing the concentration of the silicate phase in SiO_2 mixture has increased the hydrophilicity of membrane and as a result then increased the permeability of water through membrane. The test results of $4SiO_2$ membrane performance are presented in Table 5.

Examining Table 5, regarding to the state of penetrant, it shows that water component is higher than the gas from low pressures up to 4 bar pressure, which compared to $1SiO_2$ membrane, it is concluded that an increase in the concentration of SiO_2 has improved the state of permeability by increasing its water content. According to the results of relative humidity change diagram of 4SiO₂ membrane presented in Figure 10, the same improvement is clear compared to membrane 1SiO₂. But according to the diagram in Figure 7, regarding to the comparison of performance with standard TiZ-V membrane, a slight improvement has been achieved in selectivity which is similar to relative humidity change graph data in Figure 10. This means that increasing the concentration of SiO_2 increases the water permeability in water test. Finally, Figure 11 presents the gas and water permeability of the 4SiO₂ membrane at different pressures and their corresponding selectivity.

3. 2. The Effect of Operating Conditions After measuring the effect of different parameters in membrane construction, in this section, this membrane is measured under changes in some operating conditions. For this

TABLE 5. 4SiO₂ membrane performance test results

P (bar)	n_F	n_p	Y^w	Y ^G	а
2	5.845E-05	1.601E-05	0.674	0.326	3.79
3	5.863E-05	1.78387E-05	0.628	0.372	6.63
4	5.893E-05	2.08073E-05	0.536	0.464	2.44
5	5.952E-05	2.6619E-05	0.427	0.573	6.2 8
6	6.032E-05	3.46833E-05	0.345	0.655	0.20



Figure 10. Diagram of changes in the residue relative humidity relative to feed in the 4MPD membrane



Figure 11. Diagram of gas and water permeability values of 4*SiO*₂ membrane and its corresponding selectivity

purpose, the 0.4 TiZ-V membrane has been subjected to the measurement of relative humidity change of feed, the change of the flow rate of dry gas, the effect of the sweeper gas and the test of increasing the operation time.

3. 2. 1. Change Relative Humidity of Feed In this experiment, by change the element temperature in the steam generator, it changes relative humidity of the inlet wet gas to the module. The results of performance tests in relative humidity values of about 50%, 65% and 80% are presented in Table 6. It should be noted that relative humidity data were 100% in this experiment as well as all experiments in the present study which indicates the high humidity of the penetrant flow in TiZ-V membranes and it has been due to the high absorption of water from the humid gas.

According to Table 6, it can be seen that at any given relative humidity of feed, selectivity has decreased with increase of pressure which, is due to the decrease of water component relative to the gas with increase of pressure. It is also seen that decrease of relative humidity of feed from 80% to 50% increased the selectivity value. This is due to the fact that the incoming gas is drier which means less moisture content with the gas.

As it can be seen in Figure 12, the selectivity of membrane has increased with decrease in relative humidity of feed and the remarkable case is small change of the selectivity values with the change of relative humidity of feed. This matter shows the stability of membrane function under the operating conditions of change relative humidity, so that with a decrease in relative humidity, we see an increase in selectivity of membrane and with an increase in relative humidity, decrease in selectivity is slightly changes. This matter can also be seen in Figure 13, which shows the change in relative humidity of the residue.

3. 3. Change the Gas Flow Rate In this section, by adjusting the flow meter placed on the residue side,

TABLE 0. Test results of relative number of feed								
а	Y ^G	Y ^w	n_p	n_F	RH _R %	P(bar)	<i>RH_F</i> %	
9.376	0.144	0.856	8.944E-06	5.774E-04	5.7	2	8.49	
3.303	0.170	0.830	8.47891E-06	5.779E-04	2.7	4	6.50	
0.225	0.218	0782	0.0867E-05	5.786E-04	8.6	6	3.50	
5.371	0.144	0.886	1.157E-05	5.800E-04	9.0	2	3.65	
2.295	0.140	0.860	1.21519E-05	5.806E-04	9.7	4	3.65	
4.211	0.185	0.815	1.29161E-05	5.814E-04	4.7	6	1.65	
3.367	0.096	0.904	1.411E-05	5.826E-04	4.10	2	2.80	
4.288	0.118	0.882	1.48941E-05	5.834E-04	3.9	4	1.81	
2.195	0.165	0.835	1.59014E-05	5.844E-04	2.8	6	6.80	

TABLE 6. Test results of relative humidity change of feed



Figure 12. Diagram of selectivity with change of relative humidity of feed

we set and check the total flow rate of the gas flow in different values. It should be noted that the gas flow rate was 50 [L/h.] in other tests except for this test which is related to the change of gas flow rate. In addition to the flow rate of 50[L/h], values of 100 and 150 have also been investigated. Performance test results for the gas flow rate change test are presented in Table 7.



Figure 13. Relative diagram of the residue with change of relative humidity of feed

Relative humidity of feed in this experiment is set at 65%.

The data in Table 7 show the decrease in membrane selectivity with increasing gas flow rate. This matter which has been occurred due to the reduction of the molecules stopping time in the vicinity of membrane in

		IADLE	7. Results of the gas f	low face change test	·		
а	Y ^G	Y ^w	n_p	n_F	$RH_R\%$	P(bar)	QF
5.371	0.114	0.886	1.157E05	5.800E-04	9.0	2	50
2.295	0.140	0.860	1.21519E05	5.806E-04	9.7	4	50
4.211	0.185	0.815	1.29161E05	5.814E-04	4.7	6	50
4.336	0.125	0.875	2.230E05	1.159E-03	6.11	2	100
2.255	0.158	0.842	2.38157E05	1.161E-03	2.10	4	100
8.188	0.203	0.797	2.56262E05	1.163E-03	3.9	6	100
8.304	0.136	0.864	3.219E05	1.738E-03	2.14	2	150
8.231	0.172	0.828	3.4146E05	1.740E-03	3.13	4	150
1.170	0.220	0.780	3.74936E05	1.743E03	9.11	6	150

TABLE 7. Results of the gas flow rate change test

order to separate the moisture from the gas is shown in Figure 14. Also, Table 7 shows maintaining the superiority of the difference between water component and the gas in all the pressures and all the investigated flow rates, which is a completely favorable state. Another result obtained from the gas flow rate change test is the possibility of membrane performance with a selectivity of 170 to 304 at investigated high flow rate.

3. 4. The Effect of Sweeper Gas Test In this section, the effect of adding sweeper gas to the inner side of the hollow fibers is investigated on increasing the selectivity. For this purpose, dry nitrogen is inserted with pressure close to zero at different flow rates from the end side of the module that we blocked by placing the cap in the previous tests. Also, regarding the other operating conditions of this experiment, it should be mentioned that the flow rate and the humidity level of feed are set at 50 [L/h] and 80% respectively. The results of the sweep gas effect test are presented in Table 8.



Figure 14. Membrane selectivity graph with change gas flow rate

As can be seen in Table 8, the entry and increase of sweeper gas has a significant effect on increase of membrane selectivity. This phenomenon is due to the role of the sweeper gas in reducing the gas concentration gradient on the outer and inner sides of membrane, so that due to the presence of gas inside the hollow fibers, the external gas has less possibility and tendency to penetrates into membrane. As can be seen in the table about, gas component in penetrant has decreased with the increase in the sweeper gas flow rate, which indicates to this matter. Another positive role of the sweeping gas is in moving the resulting moisture inside the fibers which causes an increase of the water gradient pressure inside and outside of membrane by faster movement and exit of the amount of water inside the fiber. And as a result, the water in the wet flow has a greater possibility and tendency to penetrate into membrane which is a very favorable role. The factor that limits the use of high amounts of sweep gas in industries is the economic debate about the cost of dry gas especially the use of the same type of under operation gas.

Figure 15 shows the changes in relative humidity of the residue relative to feed for this case which shows the decrease of relative humidity of the wet gas from 80% to an average of about 7% during only one membrane phase.

3. 5. Increase the Operation Time Test In this section, the changes in the performance of membrane selectivity are investigated during the increase in the duration of membrane operations in such a way that operational data has recorded every 30 minutes during 7 hours of continuous operations to obtain the graph of selectivity changes during this period. The operational conditions include feed relative humidity equal to 80%, feed gas pressure equal to 4 bar, gas flow rate equal to 50 [L/h.] and without using sweeper gas.

TABLE 8. The results of the sweeper gas effect test

а	Y ^G	Y ^w	n_p	n_F	RH _R %	p(bar)	Sw(mL/s)
3.367	0.096	0.904	1.411E-05	5.826E-04	4.10	2	0
4.288	0.118	0.882	1.448941E-05	5.834E-04	3.9	4	0
2.195	0.165	0.835	1.59014E-05	5.844E-04	2.8	6	0
1.416	0.086	0.914	1.421E-05	5.827E-04	9.8	2	1
5.323	0.107	0.893	1.47108E-05	5.832E-04	5.8	4	1
6.216	0.152	0.848	1.55579E-05	5.840E-04	2.8	6	1
8.494	0.073	0.927	1.409E-05	5.826E-04	6.8	2	5
7.381	0.093	0.907	1.44097E-05	5.829E-04	3.8	4	5
9.276	0.123	0.877	1.50901E-05	5.836E-04	9.7	6	5
1.543	0.067	0.933	1.413E-05	5.826E-04	1.8	2	10
5.418	0.085	0.915	1.44652E-05	5.829E-04	7.7	4	10
1.315	0.110	0.890	1.49611E-05	5.834E-04	4.7	6	10



Figure 15. Diagram of changes in relative humidity of the residue relative to feed in Sw10 mode

The results of the increase the operations time test and its changes are presented in Table 9 and Figure 16, respectively. These results indicate the stability of membrane performance over time. As can be seen, this stability of the results is also seen in the removal of moisture from the gas which in all results, the gas moisture has decreased from about 80% to about 90% during only one membrane phase and also regarding the superiority of water component in the penetrant relative to the gas component, there is this stability which indicates the performance of membrane to operate at higher pressures. In addition, it can be seen that the rate of selectivity with an average of 288 in the investigated time period according to the above graph has also had a

IABLE 9. lest results of increase the operations time								
а	Y ^G	Y^w	n_p	n_F	$RH_R\%$	t		
6.287	0.119	0.881	1.476E-05	5.831E-04	5.9	30		
9.288	0.118	0.882	1.47808E-05	5.832E-04	1.9	60		
8.287	0.119	0.881	1.47096E-05	5.832E-04	2.9	90		
0.288	0.119	0.881	1.46857E-05	5.832E-04	4.9	120		
6.288	0.119	0.881	1.46873E-05	5.832E-04	1.9	150		
6.287	0.119	0.881	1.47496E-05	5.832E-04	1.9	180		
0.288	0.118	0.882	1.47595E-05	5.832E-04	2.9	210		
4.288	0.119	0.881	1.47439E-05	5.832E-04	2.9	240		
6.288	0.119	0.881	1.48048E-05	5.833E-04	9.8	270		
8.287	0.119	0.881	1.46909E-05	5.832E-04	2.9	300		
3.289	0.119	0.881	1.47404E-05	5.832E-04	1.9	330		
5.287	0.119	0.881	1.467E-05	5.831E-04	4.9	360		
3.279	0.118	0.882	1.47591E-05	5.832E-04	1.9	390		
0.288	0.119	0.881	1.47657E-05	5.832E-04	2.9	420		



Figure 16. Diagram of selectivity changes with increase the operations time

relatively increasing trend. In particular, as shown in Figure 16, the changes in selectivity of membrane in all

investigated times has been varied only between 287.5 and 289.5 which shows the stability of membrane performance with increasing time of operations.

4. CONCLUSION

The study was conducted on the possibility of using TiO_2 zeolite nanocomposite membranes in order to remove moisture from gas with the approach of natural gas and nitrogen and with the aim of influencing membrane manufacturing parameters.

According to the comparison between the SiO_2 concentrations, compared to the TiZ-V membrane, it is concluded that the decrease in SiO_2 concentration has increased the gas permeability. An increase in gas permeability is considered an undesirable situation because the membrane must have minimum gas

permeability and maximum water permeability. The cause of this phenomenon is due to the decrease in the concentration of SiO_2 and as a result of not completing the cross-linking process on the surface of the membrane, which makes the cross-linking layer of the membrane, especially at higher pressures, the possibility of resistance to prevent the penetration of nitrogen molecules. lost and the permeability of nitrogen gas increases.

The findings showed that increasing the concentration of SiO₂ has the greatest effect on increasing membrane water flux. According to the other findings of this study, the effect of increasing the reaction time of the vapor phase carrier on reducing the selectivity reduction (drop trend) was at higher pressures. Although this measurement was not done due to the safety limitation of the equipment to measure the performance of membrane at pressures higher than 7 bar, but some graphs were presented regarding the state of penetrant which were a measure of the superiority of water component in the penetrant relative to gas component and the higher water component in the penetrant, is an indicator of the appropriateness of membrane's performance at its corresponding pressure, in such a way that the investigation of the difference trend of this two diagrams shows the possibility of using membrane at higher pressures.

According to the comparison between SiO_2 concentrations, compared to TIZ-V membrane, it is concluded that the increase in SiO_2 concentration has caused a slight decrease in gas permeability. An increase in SiO_2 provides the required amounts for the reaction with the existing amount of P-TiS, but its higher amounts will not cause more transverse connection of the network due to the lack of reaction with P-TiS, so if the results show that there is no major change in the amounts The permeability of the 4SiO2 membrane was due to the lack of major change in the transverse connection of the selective layer compared to the TiZ-V membrane.

Next, in order to increase the performance of membrane as much as possible, the sweeper gas was added from the inside of membrane. Increase in the sweeper gas, which increases the water concentration gradient and decreases the gas concentration gradient on the sides of membrane wall, increases the selectivity of membrane to the best level of 543. Finally, in order to investigate the stability of membrane in longer operation times, the test of increase the operation times was carried out and no change in relative humidity reduction data and insignificant changes in its selectivity indicate the stability of membrane with an increase in the dehumidification operation time.

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چکيده

Persian Abstract

هدف اصلی در این پژوهش استفاده از غشاهای نانو کامپوزیتی زئولیت TiO2به منظورحذف رطوبت از گاز است. به همین جهت انتخاب و ساخت یک غشا TiZ-V معین به عنوان غشاء معیار، که حاصل براورد اولیه از یک غشا مناسب جهت نم زدایی از گاز بوده صورت گرفت و این غشا به عنوان معیاری برای سنجش اثر پارامترهای ساخت قرارگرفت . یافته ها نشان داد که افزایش غلظتSiO2 دارای بیشترین اثر در افزایش شار آب غشا با توجه به تاثیر افزایش زمان واکنش حامل فاز بخار و کاهش روند افت گزینش پذیری در فشارهای بالاتر بوده است. همچنین آزمایشات تغیر رطوبت نسبی خوراک نشان دهنده کارایی بهتر غشا در رطوبت های نسبی پایین تر از 80٪ بوده است گزینش پذیری در فشارهای بالاتر بوده است. همچنین آزمایشات تغیر رطوبت نسبی خوراک نشان دهنده کارایی بهتر غشا در رطوبت های نسبی پایین تر از 80٪ بوده است به نحوی که تحت شرایط رطوبت نسبی کمتر ، گزینش پذیری غشا افزایش یافته است. مورد مثبت دیگر یافت شده تغییر اندک کارایی گزینش پذیری غشا نسبت به رطوبت های نسبی مختلف است. این مورد نشان دهنده پایداری عملکرد غشا افزایش یافته است. مورد مثبت دیگر یافت شده تغییر اندک کارایی گزینش پذیری غشا نسبت به رطوبت ها نسبی مختلف است. این مورد نشان دهنده پایداری عملکرد غشا تحت شرایط مختلف رطوبت گاز خوراک است. سپس به منظور افزایش هرچه بیشتر عملکرد غشا را سویپر از داخل ممبران اضافه شد. افزایش گاز سوئیر که باعث افزایش گرادیان غلظت آب و کاهش گرادیان غلطت گاز در طرفین دیواره غشا می شود، گزینش پذیری غشا را به بهترین سطح 543 افزایش می دهد.