Separation of Binary Mixtures of Propylene and Propane by Facilitated Transport through Silver Incorporated Poly(Ether-Block-Amide) Membranes

R. Surya Murali, K. Yamuna Rani, T. Sankarshana, S. Sridhar

To cite this version:
R. Surya Murali, K. Yamuna Rani, T. Sankarshana, S. Sridhar. Separation of Binary Mixtures of Propylene and Propane by Facilitated Transport through Silver Incorporated Poly(Ether-Block-Amide) Membranes. Oil & Gas Science and Technology - Revue d’IFP Energies nouvelles, Institut Français du Pétrole, 2015, 70 (2), pp.381-390. 10.2516/ogst/2013190. hal-01931335

HAL Id: hal-01931335
https://hal.archives-ouvertes.fr/hal-01931335
Submitted on 22 Nov 2018

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
This paper is a part of the hereunder thematic dossier published in OGST Journal, Vol. 70, No. 2, pp. 215-390 and available online here.

Cet article fait partie du dossier thématique ci-dessous publié dans la revue OGST, Vol. 70, n°2, pp. 215-390 et téléchargeable ici.

**DOSSIER Edited by/Sous la direction de : B. Dewimille**

**Fluids-Polymers Interactions: Permeability, Durability**

**Interactions fluides polymères : perméabilité, durabilité**

*Oil & Gas Science and Technology – Rev. IFP Energies nouvelles, Vol. 70 (2015), No. 2, pp. 215-390*

Copyright © 2015, IFP Energies nouvelles
Separation of Binary Mixtures of Propylene and Propane by Facilitated Transport through Silver Incorporated Poly(Ether-Block-Amide) Membranes

R. Surya Murali1,4, K. Yamuna Rani2, T. Sankarshana3, A.F. Ismail4 and S. Sridhar1*

1 Membrane Separations Group, Chemical Engineering Division, Indian Institute of Chemical Technology, Hyderabad 500 607 - India
2 Process Dynamics and Control Group, Chemical Engineering Division, Indian Institute of Chemical Technology, Hyderabad 500 607 - India
3 College of Technology, Osmania University, Hyderabad 500 607 - India
4 Advanced Membrane Technology Research Centre (AMTEC), Universiti Teknologi Malaysia, 81310 Skudai, Johor - Malaysia

e-mail: suryaracha@yahoo.co.in - kyrani@iict.res.in - tsankarshana@yahoo.com - afauzi@utm.my - sridhar11in@yahoo.com

* Corresponding author

Abstract — The separation of propylene and propane is a challenging task in petroleum refineries due to the similar molecular sizes and physical properties of two gases. Composite Poly(ether-block-amide) (Pebax-1657) membranes incorporated with silver tetra fluoroborate (AgBF4) in concentrations of 0-50% of the polymer weight were prepared by solution casting and solvent evaporation technique. The membranes were characterized by Scanning Electron Microscopy (SEM), Fourier Transform InfraRed (FTIR) and wide-angle X-Ray Diffraction (XRD) to study surface and cross-sectional morphologies, effect of incorporation on intermolecular interactions and degree of crystallinity, respectively. Experimental data was measured with an indigenously built high-pressure gas separation manifold having an effective membrane area of 42 cm2. Permeability and selectivity of membranes were determined for three different binary mixtures of propylene-propane at pressures varying in the range 2-6 bar. Selectivity of C3H6/C3H8 enhanced from 2.92 to 17.22 and 2.11 to 20.38 for 50/50 and 66/34 C3H6+C3H8 feed mixtures, respectively, with increasing loading of AgBF4. Pebax membranes incorporated with AgBF4 exhibit strong potential for the separation of C3H6/C3H8 mixtures in petroleum refineries.

Résumé — Séparation de mélanges binaire de propylène et de propane par transport au travers des membranes de poly(éther-bloc-amide) incorporant de l’argent — La masse moléculaire et les propriétés physiques similaires du propylène et du propane rendent problématique la séparation des deux gaz dans les raffineries de pétrole. Des membranes composites de poly (éther-bloc-amide) (Pebax-1657) incorporant du tétrafluoroborate d’argent (AgBF4) dans des concentrations massiques allant de 0 à 50 % ont été préparées par enduction et évaporation du solvant. Les membranes ont été caractérisées par Microscopie Électronique à Balayage (MEB), par spectroscopie InfraRouge à Transformée de Fourier (IRT) et par diffractométrie de rayons X aux grands angles (DRX) afin d’étudier respectivement la morphologie de surface et transversale, l’impact de l’intégration d’AgBF4 sur les interactions intermoléculaires et le degré de cristallinité. Les données expérimentales ont été collectées dans une chambre de séparation à haute pression conçue spécialement ayant une surface effective de membrane de 42 cm2. La perméabilité et la sélectivité des membranes sont déterminées pour trois mélanges...
INTRODUCTION

Propylene/propane separation is one of commercially important process in petrochemical industry and petroleum refinery. Propylene is main resource for the production of polymers, acids, alcohols, esters, ethers and some other intermediates. Propane (C₃H₈) and propylene (C₃H₆) have similar molecular sizes and physical properties; hence separation of these two gases is complex. Many processes like absorption, adsorption and membrane techniques have been developed as an alternative to this separation [1-3]. Among all these, membrane separation, especially use of facilitated transport membranes, could be an alternative to distillation or complement the same by reducing the number of stages [4, 5].

Burns and Koros [6] had conducted a comprehensive review by evaluating all of the literature available until that period to define the upper bound relationship for C₃H₆/C₃H₈ separation. Nowadays, facilitated transport membranes utilize silver salt in appropriate polymer matrices, which have exhibited high performance for olefin/paraffin separation [7-10]. In our earlier work, C₃H₆ recovery using metal ion incorporated PolyPhenylene Oxide (PPO) and EthylCellulose (EC) membranes were reported [11, 12]. Silver salts like AgBF₄, AgNO₃, AgClO₄ and AgCF₃SO₃ were dissolved in the polymer solution for synthesizing silver embedded membranes to enhance the separation properties. Mixtures of silver salts were also incorporated for achieving better separation [7]. Sunderrajan et al. [13] reported that the order of olefin solubility in blends containing 1 mol of silver ions per mole of Ethylene Oxide (EO) units is:

AgBF₄ > AgCF₃SO₃ > AgNO₃

Kim et al. [14] had investigated the role of the polymer matrix in facilitated olefin transport through polymer/silver salt complex membranes, particularly its influence on the structure and interactions of polymer/silver salt complexes and hence on the interactions of silver ions with olefin molecules. It was found that the threshold concentration of silver salt for facilitated olefin transport depends on the polymeric ligand and decreases in the following order of amide > ketone > ester. Several researchers have been working using pure C₃H₆ and C₃H₈ gases to study the performance of polymeric membrane by incorporating metal complexes which facilitate the transport of C₃H₆.

Poly(ether-block-amide) resin is best known under the trademark Pebax and is a thermoplastic elastomer combining linear chains of the hard polyamide segments that provide mechanical strength with interspacing of flexible polyether segments that offer high permeability due to greater chain mobility of the ether linkage [15]. Crystalline amide block in Pebax functions as an impermeable phase, whereas the ether block acts as a permeable phase because of its high chain flexibility. Many researchers had worked with different grades of Pebax and different silver salts by varying degree of loading [16, 17]. Pebax-2533 consists of 80% amide and 20% ether groups, which allow high permeability and less resistance to plasticization. Pebax-4011 gives lower permeability due to the presence of only 57% soft ether block segment.

To counter these problems, Pebax-1657, constituted by 40% amide groups and 60% ether linkages, was selected as the base polymer for AgBF₄ incorporation to evaluate propylene-propane separation performance in the present study. Silver salt was expected to undergo adequate solubilization within the matrix of this block copolymer in order to allow enhanced interaction with the unsaturated olefin gas. Membranes with different AgBF₄ concentration in the polymer matrix were synthesized and characterized using SEM, FTIR and XRD studies. Experiments were conducted using these membranes in a permeability cell with propylene-propane mixtures of different compositions at varying feed pressures to determine the permeability and selectivity of each membrane for these cases, and also to find the AgBF₄ concentration most suitable for the desired separation.

1 MECHANISM OF FACILITATED TRANSPORT

The facilitated transport membranes involve reversible complexation reaction in addition to the simple solution-diffusion mechanism. The penetrant dissolves in the membrane and then diffuses down the concentration gradient or it can react with the complexing agent or a carrier and then diffuse down the concentration gradient of the carrier-gas complex. The latter transport
mechanism is not available to the saturated paraffin molecules such as C$_3$H$_8$, as they do not react with the complexing agent.

Figure 1 shows the metal-olefin bonding described by the Dewar-Chatt model and hypothetical structure of the AgBF$_4$ complex with C$_3$H$_6$ molecule in the polymeric matrix. The complex is formed by double bonding of a silver atom with the olefin [18, 19]. Both the metal and alkenes act as an electron donor and acceptor respectively in the complexation interaction. A $\pi$ component of the bond results from overlap of the vacant outermost $s$ atomic orbital of the metal with the full $\pi$ (bonding) molecular orbital of the olefin. This new molecular orbital, formed by donation of electrons from olefin to metal, has electron density concentrated between the bond members. In Ag$^+$ ions, the outermost $s$ orbital is empty because the single electron present in the metal is lost upon ionization to $a+1$ valence. In nonionizing facilitators, the metal is often bound to be an electronegative atom that withdraws electron density from the metal, resulting in a partial positive charge and a substantially vacant outermost $s$ orbit. A $\pi$ component of the metal-olefin bond is formed by back donation of electrons from the full outer $d$ atomic orbital of the metal to the $\pi^*$ (anti-bonding) molecular orbital of the olefin. The new molecular orbital has a nodal plane of electron density between the membranes of the bond.

2 EXPERIMENTAL

2.1 Materials

Commercially available Pebax-1657 was purchased from Atofina Chemicals, France, whereas PolyEtherSulfone (PES) powder was procured from Radel, USA. AgBF$_4$ salt was supplied by Aldrich Chemical, USA. Ethanol, DiMethylFormamide (DMF) and propionic acid were purchased from Loba Chemie, Mumbai, India. Gas mixture cylinders of C$_3$H$_6$ and C$_3$H$_8$ (50/50, 66/34 and 90/10 mole%) were supplied by Bhoruka Gases Ltd., Bangalore, India, at filling pressures of 9 bar each. Deionized water was produced with a laboratory reverse osmosis system.

2.2 Membrane Synthesis

PES substrate of approximately 30 000 molecular weight cutoff (MWCO) was prepared by procedure described elsewhere [20]. PES was dissolved in DMF for making 18% w/v solution and 3% propionic acid was added as additive. Homogeneous polymer solution was cast on a non-woven fabric, which was fixed on a glass plate. After casting, the glass plate was immersed in ice-cold water bath for 5-10 min to obtain ultra porous thin substrate. AgBF$_4$ loaded Pebax membranes were prepared on PES substrate by solution casting and solvent evaporation technique.

A 4% Pebax solution was prepared by adding polymer pellets to solvent mixture of 70% ethanol and 30% v/v water. The polymer was dissolved at 90$^\circ$C with rigorous stirring maintained under constant reflux over a period of 6 h. AgBF$_4$ in proportion of 35, 40 and 50% of the polymer weight was added to Pebax solution and kept for stirring. A small quantity of an oxidizing agent, commonly hydrogen peroxide, was added to stabilize the solution and prevent silver poisoning and silver loss. Without the addition of a stabilizing agent, polymer solutions containing silver salts turned black within a few hours, indicating that silver ions were being reduced to uncharged silver atoms. Hydrogen peroxide was added for slowing down the reduction of silver ions to uncharged silver atoms. The bubble free solution was cast on PES substrate using a doctor’s blade. The obtained Thin Film Composite (TFC) membrane was dried at ambient temperature (30$^\circ$C) for 3 h followed by vacuum drying for 24 h to remove residual solvent. Since it is difficult to accurately determine the exact thickness of the skin layer of a composite membrane, the authors carefully peeled off the top layer from the PES substrates in order to measure the effective thickness.

2.3 Membrane Characterization

The surface and cross-sectional morphology of Pebax and Ag-Pebax membranes were studied by SEM using a Hitachi S2150 microscope. To obtain smooth cross-section, the samples were cut in liquid nitrogen. FTIR spectra of Pebax and Ag-Pebax composite membranes...
were scanned between 400 and 3700 cm\(^{-1}\) using Perkin-Elmer-283B FTIR Spectrometer. A Siemens D 5000 powder X-ray diffractometer was used to study the solid-state morphology of Pebax and Ag-Pebax composite membranes. X-rays of 1.5406 Å wavelengths were generated by a CuK\(\alpha\) source. The angle (2\(\theta\)) of diffraction was varied from 0° to 65° to identify the crystal structure and intermolecular distances between the intersegmental chains.

2.4 Permeability Studies

A schematic diagram of the experimental setup to carry-out permeability studies is shown in Figure 2. The gas permeation tests were conducted using an indigenously designed and fabricated permeation test cell made of Stainless Steel (SS) 316. The detailed description of test cell was given in our earlier publication [20]. Gas inlet and outlet ports were provided in the test cell for the transport of feed, permeate, retentate and carrier streams. The test cell contained a circular perforated plate affixed with a mesh to support membrane of effective area 42 cm\(^2\). All the tubes were made of 1/4" SS 316. Needle valves (Swagelok, USA) with 1/4" SS 316 end connections were used to regulate the flow of inlet and outlet streams. Membrane is supported on a metallic disc across which feed flows. During permeability studies, permeate line is evacuated to remove air or residual gases present. However, evacuating feed line may tear/damage the membrane or make the edges of the membrane to get displaced from the rubber ‘O’ ring used for sealing the gas tight arrangement. Therefore, feed/permeate lines were flushed with the feed gas to remove the residual gas/air present inside. Permeability values were recorded only after steady state was reached or constant flux was attained during the experiments.

In the present work, experiments were conducted with three binary feed mixtures for testing, in which C\(_3\)H\(_6\) is 50%, 66% and 90% and the remaining is C\(_3\)H\(_8\). Since the permeate flow rate was considerable, carrier gas was not employed. Before each experiment, the permeate line was evacuated with a vacuum pump (Hindhivac, Bangalore, India) with the perforated plate support ensuring no membrane rupture. The feed and retentate lines cannot be evacuated and hence were flushed with the feed to remove residual air and ensure maximum purity of the gas inside the cell. Experiments were conducted over a feed pressure range of 2-6 bar at ambient temperature (30 ± 2°C). Feed gas was introduced slowly into the upper chamber of the test cell by means of a needle valve with the outlet valve kept partially closed until the gauge indicated the desired pressure. Continuous

\[ P_1 \quad V_1 \quad V_2 \quad P_2 \quad V_3 \quad T-C \quad P_3 \quad S-P \quad V_4 \quad V_5 \quad V_6 \quad V_7 \quad \text{Permeate} \]

\[ \text{T-C} \quad \text{Membrane test cell}, \text{V-P} \quad \text{Vacuum pump}, \text{P1, P2, P3, P4} \quad \text{Pressure gauges} \]

\[ \text{S-B} \quad \text{Soap bubble meter}, \text{S-P} \quad \text{Sample point}, \text{V1, V2, V3, V4} \quad \text{Needle valves} \]

Figure 2
Schematic of high-pressure gas permeability manifold.
flow of the gas in the feed chamber was maintained. Once steady state was attained, the flow rate of permeate and retentate were measured with the help of a soap bubble meter. Permeate side valve was kept open and flow rate was measured at atmospheric pressure (1 bar). Provision was made for online analysis by throttling respective individual valve of feed, permeate and retentate with a gas chromatograph equipped with a 30ft SP-1700 Chromosorb P-AW column (stationary phase: methyl cyan silicone) which gave complete resolution of C$_3$H$_6$ and C$_3$H$_8$ peaks. The chromatograph was operated with a thermal ionization detector and Helium was used as the carrier gas. The oven temperature was maintained at 60°C. The degree of error while collecting experimental data is about ±2% which also accounts for deviation in analytical method. As long as the membrane was not exposed to sunlight the permeability results obtained with different samples of the Ag loaded membranes were very consistent (reproducible) as long as the substrate porosity and selective layer thickness were kept fairly constant.

A binary feed gas, $F$ (cm$^3$/s), with a mole fraction of $x_f$ entered the membrane test cell giving two streams as outlet with gas which passed through the membrane denoted as permeate, $V$ (cm$^3$/s), with a moles fraction of $y_p$; and non-permeating gas, $R$ (cm$^3$/s), with a mole fraction of $x_0$.

The rate of permeation of species $A$, in binary mixture of $A$ and $B$, is given by:

$$V_y_p = \left(\frac{P'_A A_m}{t}\right) (p_h x_0 - p_l y_p)$$  \hspace{1cm} (1)

where $P'_A$ is the permeability of $A$ in the membrane, cm$^3$(STP).cm/(s.cm$^2$.cmHg); $A_m$ is membrane area, cm$^2$; $t$ is membrane thickness, cm; $p_h$ is the pressure in feed side, cmHg; $p_l$ is the pressure in the permeate side, cmHg.

A similar equation can be written for the component B:

$$V(1 - y_p) = \left(\frac{P'_B A_m}{t}\right) (p_h (1 - x_0) - p_l (1 - y_p))$$  \hspace{1cm} (2)

Permeability of species $i$, $P'_i$, in the gas mixture is given by:

$$P'_i = \frac{y_p l}{A_m (x_f p_h - y_r p_l)} \left(\frac{273}{T}\right) \left(\frac{P_{atm}}{76}\right) \left(\frac{dv}{dt}\right)$$  \hspace{1cm} (3)

where $P_{atm}$ is the atmospheric pressure, $(dv/dt)$ is the volumetric flow rate, cm$^3$/s.

The actual separation factor $\alpha$ is given by:

$$\alpha = \frac{P'_A}{P'_B}$$  \hspace{1cm} (4)

3 RESULTS AND DISCUSSION

3.1 Membrane Characterization

3.1.1 SEM

Figures 3a and c show the surface morphology of pristine Pebax and 50 wt% Ag-Pebax membranes which are clear and homogeneous. Surface morphology of Ag-Pebax clearly shows a regular polymer surface without any particle clusters. It can be concluded that the added salt is completely solubilized in the polymer matrix. Cross-section of Pebax is shown in Figure 3b, which clearly reveals the presence of three layers in the TFC membrane. Top layer is the active Pebax, second layer is ultraporous PES substrate and the last layer is the non-woven polyester fabric support. Cross-sectional views of Ag-Pebax is illustrated in Figure 3d which reveals that Pebax containing AgBF$_4$ partially penetrates into the ultraporous PES support and forms a thin but uniform selective dense layer over it.

3.1.2 FTIR

Figure 4 represents Fourier Transform InfraRed (FTIR) spectra of Pebax and Ag-Pebax membranes. In Pebax (Fig. 4a), the characteristic peaks at 1599 and 1058 cm$^{-1}$ represent –C=C and –C=O– stretching vibrations respectively. Two more peaks at 1 641 and 3 299 cm$^{-1}$ indicate the presence of H–N–C=O and N–H groups respectively. The stretching vibration at 2 940 cm$^{-1}$ indicates the presence of aliphatic –C–H. The C=O stretching in amide group of Pebax is constituted of two types, the free amide C=O peak at 1 657 cm$^{-1}$ and the hydrogen bonded amide peak at 1 640 cm$^{-1}$. The characteristic peaks of Ag-Pebax membranes are similar to pristine Pebax, which may be probably due to the suppression of vibration effects caused by silver salt with polymer matrix. Vibrations at 1 599 cm$^{-1}$ and 3 060 cm$^{-1}$, caused by C=C stretching, differ a little from plane Pebax, steepness of peak decreases with increasing salt loading. This could be due to temporary bonding between Ag$^+$ and C=C. The metal ligand interaction bands are seen distinctly in the low frequency region (600-400 cm$^{-1}$). It also reveals that the C=O vibrations of HN–C=O groups at 1 641 cm$^{-1}$ decrease with increasing AgBF$_4$. This decrease indicates the loosening of the
C=O bond due to the electron donation from the C=O oxygen to the silver ions. Thus we can conclude that Ag have interaction with the electron donating sites present in the polymer backbone. The vacant $d$ orbital present in silver is available for binding with the unpaired electrons present in the double bonded C$_3$H$_6$ olefin leading to II-complexation.

3.1.3 XRD

Among the different degrees of loading investigated, 50% Ag-Pebax membrane was chosen for comparison with the pristine polymer. X-ray diffractograms of Pebax and 50% Ag-Pebax composite are shown in Figure 5. Both membranes exhibit semicrystalline nature. Pebax (a) is a semicrystalline polymer, which shows narrow diffraction peaks at 14°, 27°, 54° and 56° of 2θ and the corresponding XRD peaks obtained for Ag-Pebax (b) are broader. This indicates an increase in intersegmental spacing, which may lead to a reduction in degree of crystallinity.

3.2 Experimental Studies

3.2.1 Effect of % AgBF$_4$ Loading

Composite membranes were prepared by varying the AgBF$_4$ incorporation in extents of 35, 40 and 50 wt% of the Pebax polymer. As long as the membrane is not exposed to sunlight the permeability results obtained with different samples of the Ag loaded membranes are
very consistent (reproducible). Table 1 shows the effect of silver loading on permeability and selectivity for 50/50 $\text{C}_3\text{H}_6 + \text{C}_3\text{H}_8$ feed mixture at a feed pressure of 4 bar. The authors have carefully peeled off the top layer from the substrates in order to measure the effective thickness for permeability calculation. With increasing AgBF$_4$ loading from 0-50 wt% of the polymer weight, the permeability enhanced from 7.67 to 99.78 Barrer for $\text{C}_3\text{H}_6$ and 2.62 to 5.79 Barrer for $\text{C}_3\text{H}_8$. On the other hand, 50 wt% Ag-Pebax exhibited higher selectivity of 17.22, which is nearly five times higher than that of the pristine Pebax membrane. Table 2 exhibits the permeability of Ag-Pebax membranes for 66/34 $\text{C}_3\text{H}_6 + \text{C}_3\text{H}_8$ feed mixture at a feed pressure of 2 bar. Selectivity of $\text{C}_3\text{H}_6/\text{C}_3\text{H}_8$ enhances with increasing AgBF$_4$ loading from 2.11 to 20.38, which is 8.6 folds higher than pristine Pebax. The permeability for 66/34 is greater than that of 50/50 $\text{C}_3\text{H}_6/\text{C}_3\text{H}_8$ mixture since the higher concentration of the preferentially permeating olefin in the former would lead to greater partial pressure gradient which is the driving force for mass transfer by solution-diffusion mechanism. Moreover, higher numbers of olefin molecules are now available for permeation by carrier mediated transport which is the more prominent route for passage through AgBF$_4$ incorporated membranes.

In pristine Pebax membrane gas transport is solely a function of solubility and diffusivity of the molecules through the polymer matrix. In contrast, gas transport in Ag-Pebax membrane is the combined effect of facilitated transport and solution-diffusion phenomenon. AgBF$_4$ forms temporary bond with $\text{C}_3\text{H}_6$ and carriers it through the membrane which results in increase of permeability. Upon increasing the AgBF$_4$ loading, the selective nature of Pebax also increases, which is consistent with the concept that Ag$^+$ ions increase the cohesive energy density of the matrix. It is believed that Ag$^+$ ions could form weak complexes with polar ether groups in the polymer backbone [21]. These complexes act as transient crosslinks that restrict polymer chain mobility, which results in the reduction of diffusion coefficients of non-interacting saturated species like $\text{C}_3\text{H}_8$. In the specific and reversible Ag$^+$-$\text{C}_3\text{H}_6$ molecular interaction, Ag$^+$ ions act as carriers for facilitated transport in the membrane and cause carrier mediated transport in addition to the normal transport process. As the Ag$^+$ ion concentration increases, large numbers of unsaturated $\text{C}_3\text{H}_6$ molecule form reversible bonds and pass through the membrane via facilitated transport. On the other hand $\text{C}_3\text{H}_8$ is rejected by the nonporous AgBF$_4$ filler and therefore it has to follow a tortuous path unlike $\text{C}_3\text{H}_6$ to reach the downstream side of the membrane. Increase in permeability of $\text{C}_3\text{H}_8$ might also be anticipated due to an increase in the membrane free volume.

![Figure 5](image_url)

**Figure 5**
XRD of a) Pebax b) 50% Ag-Pebax membranes.

| % AgBF$_4$ | Membrane thickness (µm) | Permeability (Barrer$^*$) | Selectivity ($\text{C}_3\text{H}_6/\text{C}_3\text{H}_8$) |
|------------|------------------------|--------------------------|----------------------------------|
|            |                        |                          | $\text{C}_3\text{H}_6$  | $\text{C}_3\text{H}_8$  |
| 0          | 15                     | 7.7                      | 2.6                             | 2.9                      |
| 35         | 20                     | 31.6                     | 6.5                             | 4.9                      |
| 40         | 25                     | 31.8                     | 3.1                             | 10.1                     |
| 50         | 25                     | 99.9                     | 5.8                             | 17.2                     |

1 Barrer = $10^{-10}$ (cm$^3$ (STP).cm$^2$.s.cmHg)
by the gas molecules permeating first which in turn make the flow of the next kind relatively easier.

Researchers had developed facilitated transport membranes with different grades of Pebax for olefin-paraffin separation. Muller et al. [16] had studied performance of Pebax 2533 (constituted by 20% amide groups) and Pebax 4011 (constituted by 43% amide groups) embedded with AgClO₄ salt, for which selectivities of 110 to 190 were observed with humidified equimolar ethane/ethylene gas mixture. Wang et al. [17] synthesized facilitated transport membrane by coating AgBF₄ over defect free Pebax 2533 membrane for C₃H₆/C₃H₈ separation. In the present work, AgBF₄ loaded Pebax 1657 (constituted by 40% amide groups) shows mixture selectivity of 17.2 with equimolar C₃H₆/C₃H₈ gas mixture. Lesser selectivity in mixture gas is due to the coupling effect and absence of any humidification. Table 3 compares performance of different membrane materials for C₃H₆/C₃H₈ binary gas mixture separation.

### Table 3

| Membrane                              | C₃H₆/C₃H₈ binary mixture | Pressure (bar) | Selectivity | References |
|---------------------------------------|---------------------------|---------------|-------------|------------|
| EC                                    | 55/45                     | 3.9           | 3.2         | [6]        |
| CA                                    | 55/45                     | 3.9           | 2.6         | [6]        |
| PSF                                   | 55/45                     | 3.9           | 1.4         | [6]        |
| POZ/AgNO₃/AgCF₃SO₃                    | 50/50                     | 0.02          | 28          | [6]        |
| POZ/AgBF₄/AgCF₃SO₃                    | 50/50                     | 0.02          | 23          | [6]        |
| Poly(ethylene-co-propylene) / 62% AgBF₄ | 50/50                     | 55            |             | [9]        |
| PAEK/Azide (carbonaceous membrane)    | 50/50                     | 6             | 32          | [22]       |
| ZIF-8 (zeolite membrane)              | 50/50                     | 1             | 28          | [23]       |
| ZIF-8/6FDA-DAM                        | 50/50                     | 0.037         | 15.5        | [24]       |
| Pebax 1657 / 50% AgBF₄               | 50/50                     | 4             | 17.2        | This study |
| Pebax 1657 / 50% AgBF₄               | 66/34                     | 2             | 20.2        | This study |

3.2.2 Effect of Feed Composition

Figure 6 illustrates the effect of C₃H₆ concentration in the binary feed mixture on the performance of the indigenously synthesized membrane. Since 50% Ag-Pebax membrane shows better performance, it was chosen to study the effect of feed composition at a constant pressure of 2 bar. Compositions of feed, permeate and retentate streams were analyzed consecutively with an online gas chromatograph. With 50% Ag loaded Pebax membrane, 50% C₃H₆ feed mixture gives 78.5% C₃H₆ in permeate whereas 66% C₃H₆ feed mixture gives 92.9% C₃H₆ and 90% C₃H₆ feed mixture gives a permeate containing 98.6% C₃H₆. As the C₃H₆ composition in feed is increased from 50 to 90%, permeability decreases from 244.3 to 79.2 Barrer. Mixture gas selectivity varies from 13.4 to 16.9 and attains a maximum of 20.4 at 66% C₃H₆.
feed mixture. Reduction in permeability might be due to limited number of sorption sites and transport passage ways available in the membrane. At higher feed C\textsubscript{3}H\textsubscript{6} concentrations, the sorption sites and silver salt in the Ag-Pebax membrane might get saturated, which could result in a reduction in permeability due to absence of carrier mediate transport.

### 3.2.3 Effect of Pressure

Figure 7 shows the effect of pressure (2-4 bar) on the performance of 50% Ag-Pebax composite membrane with the feed mixture of 50% C\textsubscript{3}H\textsubscript{6}. Increasing feed pressure results in gradual increase of selectivity from 13.4 to 17.2 whereas permeance decreases from 244.3 to 99.8 Barrer. Presence of fixed sorption sites in the membrane enhances the uptake of pure gas molecules with rising feed pressure. However, the existence of even a small amount of C\textsubscript{3}H\textsubscript{8} in the feed alters the permeability due to coupling effect. During permeation of binary gas mixture, the presence of one gas may affect the permeation of the other through transfer of momentum leading to dragging of less permeable gas through the barrier, which is called coupling effect [25]. For pure propylene permeation, the membrane permeability is determined by the interactions between propylene molecules and the membrane material. Compared to propylene, propane is a slow gas. When propane is present, the permeation is complicated by competitive sorption and coupled transport between the two feed components resulting in greater free volume and reduced selectivity.

### CONCLUSIONS

The present study demonstrated the feasibility of preparing TFC membrane of Pebax as well as Ag-Pebax on PES support for C\textsubscript{3}H\textsubscript{6}/C\textsubscript{3}H\textsubscript{8} separation. Experiments were conducted with three feed mixtures, in which C\textsubscript{3}H\textsubscript{6} is 50%, 66% and 90% and the remaining is C\textsubscript{3}H\textsubscript{8}. Membranes were physically modified by incorporating 35, 40 and 50% of AgBF\textsubscript{4} to facilitate the transport of C\textsubscript{3}H\textsubscript{6}. SEM revealed complete dissolution of silver salt within the polymer solution. Characterization by XRD and FTIR showed Ag\textsuperscript{+} ions induced weak physical interactions within the polymer matrix. Increasing the percentage of Ag\textsuperscript{+} incorporation improved the separation performance of membranes. It was found that the selectivity and permeability of the membrane with respect to C\textsubscript{3}H\textsubscript{6} improved with increasing feed concentration of the gas due to greater partial pressure gradient. A hybrid process combining membrane separation and distillation would make production of polymer grade propylene in refineries technically feasible, more economical and safe due to reduction in number of distillation trays in the lengthy C\textsubscript{3} splitter.

### ACKNOWLEDGMENTS

The first author R. Surya Murali is grateful to the Council of Scientific and Industrial Research (CSIR), New Delhi, India for funding his Ph.D. research programme under GATE-JRF scheme. The corresponding author acknowledges the financial support from CSIR under their XII Five Year Plan MATES project.

### REFERENCES

1. Maryam A., Tahereh K., Mohammad R. (2008) A review on olefin/paraffin separation using reversible chemical complexation technology, *Journal of Industrial and Engineering Chemistry* **14**, 5, 622-638.

2. Grande C.A., Rodrigues A.E. (2005) Propane/propylene separation by pressure swing adsorption using zeolite 4A, *Industrial and Engineering Chemistry Research* **44**, 8815-8829.

3. Faiz R., Li K. (2012) Olefin/paraffin separation using membrane based facilitated transport/ chemical absorption techniques, *Chemical Engineering Science* **73**, 261-284.

4. Pettersen T., Argo A., Noble R.D., Koval C.A. (1996) Design of combined membrane and distillation processes, *Separation Technology* **6**, 3, 175-187.

5. Stephan W., Noble R.D., Koval C.A. (1995) Design methodology for a membrane/distillation column hybrid process, *Journal of Membrane Science* **99**, 3, 259-272.

6. Burns R.L., Koros W.J. (2003) Defining the challenges for C\textsubscript{3}H\textsubscript{6}/C\textsubscript{3}H\textsubscript{8} separation using polymeric membranes, *Journal of Membrane Science* **211**, 2, 299-309.
7 Kim J.H., Park S.M., Won J., Kang Y.S. (2005) Unusual separation property of propylene/propane mixtures through polymer/silver complex membranes containing mixed salts, *Journal of Membrane Science* 248, 1-2, 171-176.
8 Lee D.H., Kang Y.S., Kim H.J. (2009) Olefin separation performances and coordination behaviors of facilitated transport membrane based on poly(styrene-b-isoprene-b-styrene)/silver salt complexes, *Macromolecular Research* 17, 2, 104-109.
9 Kim J.K., Min B.R., Kim Y.W., Kang S.W., Won J., Kang Y.S. (2007) Novel composite membranes comprising silver salts physically dispersed in poly(ethylene-co-propylene) for the separation of propylene/propane, *Macromolecular Research* 15, 4, 343-347.
10 Joo S.H., Kim J.H., Kang S.W., Jang J., Kang Y.S. (2007) Propylene sorption and coordinative interactions for poly(N-vinyl pyrrolidone-co-vinyl acetate)/silver salt complex membranes, *Journal of Polymer Science Part B: Polymer Physics* 45, 16, 2263-2269.
11 Susheela B., Sridhar S., Khan A.A. (2000) Recovery of propylene from refinery off-gas using metal incorporated ethylcellulose membranes, *Journal of Membrane Science* 174, 1, 67-69.
12 Sridhar S., Khan A.A. (1999) Simulation studies for the separation of propylene and propane by ethylcellulose membrane, *Journal of Membrane Science* 159, 1-2, 209-219.
13 Sunderrajan S., Freeman B.D., Hall C.K., Pinnau I. (2001) Propylene and propylene sorption in solid polymer electrolytes based on poly(ethylene oxide) and silver salts, *Journal of Membrane Science* 182, 1-2, 1-12.
14 Kim J.H., Min B.R., Won J., Joo S.H., Kim H.S., Kang Y.S. (2003) Role of polymer matrix in polymer/silver complexes for structure, interaction, and facilitated olefin transport, *Macromolecules* 36, 16, 6183-6188.
15 PEBAX-Basis of Performance, Polyether Block Amide, Elf Atochem Technical Document, Paris, France.
16 Muller J., Peinemann K.V., Muller J. (2002) Development of facilitated transport membranes for the separation of olefins from gas streams, *Desalination* 145, 1-3, 339-343.
17 Wang Y., Ren J., Deng M. (2011) Ultrathin solid polymer electrolyte PEI/Pebax2533/AgBF₄ composite membrane for propylene/propane separation, *Separation and Purification Technology* 77, 1, 46-52.
18 Ferraz H.C., Duarte L.T., Alves M.D.T.L.M., Habert A.C., Borges C.P. (2007) Recent achievements in facilitated transport membranes for separation processes, *Brazilian Journal of Chemical Engineering* 24, 1, 101-118.
19 Hamouda S.B., Quang T.N., Dominique L., Sadok R. (2008) Facilitated transport of ethylene in poly(amide12-block tetramethylenoxide) copolymer/AgBF₄ membranes containing silver (I) and copper (I) ions as carriers, *Journal of Applied Sciences* 8, 7, 1310-1314.
20 Surya Murali R., Sridhar S., Sankarshana T., Ravikumar Y.V.L. (2010) Gas permeation behavior of Pebax-1657 nanocomposite membrane incorporated with multiwalled carbon nanotubes, *Industrial and Engineering Chemistry Research* 49, 14, 6530-6538.
21 Merkel T., Blanc R., Zeid J., Suwarlim A., Firtat B., Wijmans H., Asaro M., Greene M. (2007) Separation of olefin/paraffin mixtures with carrier facilitated membranes, *Project DE-FC36 04GO14151*.
22 Yeom C.K., Lee S.H., Lee J.M. (2000) Study of transport of pure and mixed CO₂/N₂ gases through polymeric membranes, *Journal of Applied Sciences* 78, 179-189.
23 Chng M.L., Xiao Y., Chung T.S., Toriida M., Tamai S. (2009) Enhanced propylene/propane separation by carbonaceous membrane derived from poly(aryl ether ketone)/2,6-bis(4-azidobenzylidene-4-methyl-cyclohexanoneinterpenetrating network, *Carbon* 47, 7, 1857-1866.
24 Pan Y., Li T., Lestari G., Lai Z. (2012) Effective separation of propylene/propane binary mixtures by ZIF-8 membranes, *Journal of Membrane Science* 390-391, 93-98.
25 Zhang C., Dai Y., Johnson J.R., Arvan O., Koros W.J. (2012) High performance ZIF-8/6FDA-DAM mixed matrix membrane for propylene/propane separations, *Journal of Membrane Science* 389, 34-42.