PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

B01D 17/00

(11) International Publication Number:

WO 98/10852

(43) International Publication Date:

19 March 1998 (19.03.98)

(21) International Application Number:

PCT/US97/16386

A1

(22) International Filing Date:

16 September 1997 (16.09.97)

(30) Priority Data:

60/026,196 16 September 1996 (16.09.96) PCT/US97/07451 2 May 1997 (02.05.97)

(34) Countries for which the regional or

international application was filed: AT et al.

wo

US

(71) Applicant: COMMODORE SEPARATION TECHNOLOGIES, INC. [US/US]; Suite 200, 3240 Town Point Drive, Kennesaw, GA 30144 (US).

(72) Inventor: KILAMBI, Srinivas; Apartment 2616, 1049 Powers Ferry Road, Marietta, GA 30067 (US).

(74) Agents: FARLEY, Felipe, J. et al.; Jones & Askew, LLP, 37th floor, 191 Peachtree Street, N.E., Atlanta, GA 30303 (US).

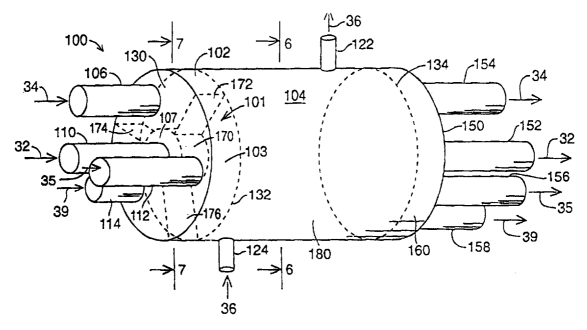
(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: SUPPORTED LIQUID MEMBRANE SEPARATION



(57) Abstract

The present invention comprises a supported liquid membrane carrier liquid with a dielectric constant between approximately (20 and 32), more particularly between (22 and 28), and preferably about (24), and the use of that carrier liquid in a supported liquid membrane device (100) to extract one or more chemical species from a feed liquid (32). The invention further comprises such a device where a temperature gradient is maintained between any of the feed, carrier (36), or strip liquids. In another embodiment of the invention, any of the feed, carrier, or strip liquids are conditioned, for example, by passing the liquids through a purifier, or replacing lost carrier liquid. In another embodiment of the invention, two or more devices (Fig. 5) are connected in series or parallel.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
ΑZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Титкеу
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		
					• .		

1

5

SUPPORTED LIQUID MEMBRANE SEPARATION

10

15

20

FIELD OF THE INVENTION

The present invention is in the field of chemical extraction, and is more particularly in the field of chemical separation by supported liquid membranes.

BACKGROUND OF THE INVENTION

Liquid membrane separation has its origins in liquid-liquid extraction which involves extraction of a solute from a first liquid using a second liquid solvent that is essentially immiscible with the first liquid. A back extraction is then typically done in a separate apparatus to remove the solute from the second liquid. Liquid-liquid extraction can be carried out in a number of devices such as mixer settlers, packed towers, bubble tray columns, asymmetrical membranes, and supported liquid membranes. Of all of these types of liquid-liquid extraction, supported liquid membrane technology belongs to a separate category of extraction that involves the presence of a carrier that forms a reversible complex with the preselected chemical species of interest.

25

30

35

Although other types of liquid-liquid extractions can be carried out using membrane supports, such processes are clearly distinguishable from supported liquid membrane transport in that such methods lack the presence of a carrier that forms a reversible complex to facilitate transport of the preselected chemical species. Such processes do not involve the formation of a reversible chemical complex but rather the physical partitioning of the solute between the two immiscible liquids. As a result, the rate of transport is very slow as the transport involves no reaction to form a chemical complex. Fluxes are low and the devices tend to be large. To facilitate liquid-liquid contact while preserving membrane life, such devices typically use asymmetrical supports such as membranes that are hydrophilic on one side and hydrophobic on the other side or membranes in which a gradient of pore size and density are used.

2

Liquid membrane transport separation is an emerging technology where specific material species are transported selectively and rapidly across a liquid membrane. Though liquid membrane transport was discovered in the early 1970s, the bulk of experimental studies involving this technique has been carried out only in the last few years.

5

10

15

20

25

30

35

In spite of the success of supported liquid membrane techniques in the laboratory, very few pilot scale studies have been undertaken. The primary reason for this low utilization has been the low flux rates and membrane instability. The high surface area per unit volume of hollow fiber microporous support members has increased the flux rate to some extent but flux rate improvements are still necessary to achieve commercial utilization. An even greater problem is membrane instability which generally occurs because of the gradual loss of the liquid membrane to the liquids on each side of the membrane. Such loss can occur because of (1) the solubility of the carrier and its diluent in the feed and strip liquids or (2) capillary displacement as a result of an osmotic pressure differential between the two sides of the membrane due to solution pumping of the feed and strip liquids.

Liquid membrane instability is a prime causes for the slow commercialization of the process. Although considerable research has been conducted to determine the causes for membrane instability, a permanent remedial solution has yet to be found. Studies as to osmotic pressure differences, interfacial membrane-solution tension, and low critical surface tension of the polymer support have been inconclusive.

Generally supported liquid membrane instability is subdivided into (1) instability arising from loss of carrier liquid (extractant) from the membrane phase leading to a loss in permeability, or (2) a complete "break down," resulting in direct contact of the feed and strip liquids. Attempts have been made to explain the complete "break down" effect on the basis of the osmotic pressure gradients across the membrane. Feed and strip liquid transport increases with this gradient, and this transport induces a repulsion of the liquid membrane phase out of the pore of the support which causes the membrane to degrade (Fabani, C., et al., J. Membrane Science, vol. 30, p. 97, 1987). Fabani concluded that osmotic pressure differences play a significant role in determining membrane stability and, as a result of these osmotic pressure differences, the carrier liquid (extractant) and its diluent is washed from the pores of the membrane. In direct contradiction, Danesi, P.R., et al., J. Phys. Chem., vol. 1, p. 1412 (1987) concluded that the "osmotic pressure model" was not responsible for membrane instability. He and Takeuchi, et al., J. Membrane Science, vol. 34, p. 19 (1987) postulated that this form of instability was due to the poor nature of the carrier liquid diluent and low carrier liquid/diluent feed and strip interfacial tensions.

Takeuchi (1987) also found that membrane instability increased with feed velocity and also with increasing hydrostatic pressure gradient across the membrane. Neplenbroeck (1987) concluded that membrane instability depended on the type of feed and strip liquids used and the molecular structure of the carrier. He concluded that there was no direct relation between viscosity of the carrier and diluent and membrane instability but rather that there was a connection with the interfacial tension of the carrier diluent and the feed and strip liquids. However, these relations are ambiguous leaving strong doubt that membrane instability is not caused by osmotic pressure differences.

Other researchers such as Chirazia (1990) and Takeuchi (1987) have suggested that the use of amines as membrane diluents leads to better stability when compared to aliphatic diluents, that the degree of amine solubility in feed and strip solutions contributes to membrane stability, that interfacial tension lowering at the carrier-diluent/feed solution and strip liquid interface increases membrane stability, that the use of polymers which can cross link the carrier to the support can increase membrane stability, that the use of hollow fiber modules rather than flat sheet modules and membrane support pore size has an effect on membrane stability, and that the surface tension of the support in relation to the carrier-diluent/feed and strip interfacial tensions affect membrane stability. Dozol, J.F., et al., J. Memb. Sci., vol. 82, p. 237 (1993) suggested that the solvent solubility of the solvent and the simultaneous drop point of the membrane as proportional to the interfacial tension could effect membrane stability. They felt that the liquid membrane must have surface tension lower than the critical surface tension of the support. They also concluded that the carrier solution viscosity has no effect on membrane stability. In view of these significant differences as to the causes of liquid membrane instability and inconclusive experimental results, there are significant doubts in the minds of researchers regarding the real cause of membrane instability. As such, a viable solution to the membrane stability problem has yet to be found and commercialization of this technique has yet to be achieved.

SUMMARY OF THE INVENTION

5

10

15

20

25

30

35

The present invention comprises a supported liquid membrane carrier liquid with a dielectric constant between approximately 20 and 32, more particularly between 22 and 28, and preferably approximately 24, and the use of that carrier liquid in a supported liquid membrane device to extract one or more chemical species from a feed liquid. The invention further comprises such a device where a temperature gradient is maintained between any of the feed, carrier, or strip liquids. Preferably, any of the feed, carrier, or strip liquids are conditioned, for example, by passing the liquids through a purifier, or replacing lost carrier liquid.

10

15

20

25

30

35

In one embodiment of the invention where three devices are connected in series, the carrier liquid of one device contains an anion exchange carrier, the carrier liquid of another device contains a neutral exchange carrier, and the carrier liquid of the other device contains a cation exchange carrier.

In a further embodiment of the invention, where a second ion is transported from the feed liquid to the strip liquid along with the ion of interest, the second ion is removed from the strip liquid to promote further transport of the ion of interest.

The present invention also includes a device containing the stable supported liquid membrane of the present invention that can be used to rapidly and efficiently remove one or more chemical species from a liquid. Thus, the present invention is therefore useful in removing contaminants from waste fluids from industrial applications, for example, for removing and reusing nickel from aluminum anodizing processes. The present method and device are also useful in removing heavy metals from waste water and in removing organic molecules from a feed stream.

One feature of the present invention is the ability to separate a large anion from a mixture of the large anion and smaller anions in a feed stream. The preferred carrier for separation of the large anion is a crown ether. The largest of the anions in the feed solution associates with the crown ether and cation and is carried across the liquid membrane with the carrier and a cation. For example, the present invention provides a means for separating large anions such as, but not limited to, pertechnetate, perrhenate, permanganate, perchlorate, perbromate, periodate, iodide, tetracyanaurate, tetraphenylborate, benzoate, and picrate anions from feed solutions containing smaller anions. The process has been found to be particularly effective in separating radioactive anions such as pertechnetate (TcO-4) from radioactive wastes that often contain substantial amounts of nitrate ion. Technetium (99Tc), a uranium fission product, is a waste material left over after reprocessing fuel rods for plutonium production. The U.S. Department of Energy currently has 1.8 tons of this waste, which has a half-life of over two hundred thousand years.

Chemical treatment involves the use of reactions to transform hazardous waste streams into less hazardous substances. Chemical treatment can be used to recover hazardous substances, which can then be put to good use. Because it can be employed to produce useful byproducts and environmentally acceptable residual effluents, chemical treatment is a far better method of waste management than disposal in a landfill.

Many different forms of chemical treatment are used in hazardous waste management. Chemical reactions are used to reduce wastes in volume, or to convert them to a less hazardous form. These methods include neutralization of acidic and alkaline

waste streams, precipitation of heavy metals to form an insoluble salt, usually with caustic lime, and addition of water-soluble chemicals and polymers to waste waters to promote coagulation and flocculation of heavy metals. These processes are used to separate suspended solids from liquids when their normal sedimentation rates are too slow to be effective. Coagulation is the addition and rapid mixing of a coagulant to neutralize charges and collapse the colloidal particles so they can agglomerate and settle. Flocculation requires gentle agitation to allow bridging of the flocculant chemical between agglomerated colloidal particles, to form large, settleable agglomerates.

5

10

15

20

25

30

35

Another chemical treatment is the use of organic polymers called polyelectrolytes, which react with colloidal material in wastewater by neutralizing the charge, and bridging individual particles to form flocs. Finally, oxidation and reduction can be used to convert toxic pollutants to harmless or less toxic substances. Oxidation is a chemical reaction in which electrons are lost from a chemical compound.

However, all the above-cited methods of chemical treatment of wastes require large physical facilities, and large amounts of capital. Furthermore, one can end up with a huge volume of liquid, even after all treatment steps are carried out.

Another type of waste recovery technology, using what is known as supported liquid membranes (SLM), can be used to separate hazardous chemical species from waste streams. In an SLM device, a porous membrane soaked with a carrier liquid separates a feed stream and a strip liquid. The waste chemical is transported from the feed stream, through the carrier liquid in the membrane, to the strip liquid. This process very selectively removes a chemical from the feed stream, concentrates it in the strip liquid - and does it in one step. The feed stream and the strip liquid do not mix. The key is the thin membrane soaked with carrier liquid, which separates the feed stream and the strip liquid. Since the membrane is very thin, very little of the expensive carrier liquid is necessary, and the whole process can be very economical.

However, no one has figured out a way to make the membrane last very long, that is, stabilize the carrier liquid. Therefore, this technology, though hinting of great promise, has heretofore not been commercialized.

The present invention shows how to stabilize the carrier liquid.

Accordingly, it is an object of the invention to provide a stabilized supported liquid membrane.

It is yet another object of the present invention to provide a method and device that is capable of removing a preselected chemical species from a solution.

10

15

20

25

30

35

Another object of the present invention is to provide a stabilized supported liquid membrane device that can remove a preselected chemical species from a solution over a long period of time without having to replace the supported liquid membrane.

Another object of the present invention is to provide a stabilized supported liquid membrane device that is capable of removing two or more chemical species from a solution.

Accordingly, it is an object of this invention to provide a means for removing large anions such as radioactive pertechnetate from waste streams.

While the forms of the invention herein disclosed constitute currently preferred embodiments, many others are possible. It is not intended herein to mention all of the possible equivalent forms or ramifications of the invention. It is to be understood that the terms used herein are merely descriptive rather than limiting, and that various changes may be made without departing from the spirit and scope of the invention.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 illustrates a supported liquid membrane device including a carrier liquid in a microporous membrane, which separates a feed liquid and a strip liquid.

Figure 2 illustrates a hollow fiber contained liquid membrane module with two hollow fibers, one fiber carrying the feed liquid the second fiber carrying the strip liquid where carrier liquid circulates between the fibers in the module.

Figure 3 is a schematic of the concentration profile across a supported liquid membrane.

Figure 4 is a perspective drawing showing a module featuring the use of hollow fibers and used for parallel processing of multiple strip liquids.

Figure 5 is a cross sectional view of the embodiment shown in Fig. 4 taken along line 6-6.

Figure 6 is a cross sectional view of the embodiment shown in Fig. 4 taken along line 7-7.

Figure 7 is an end view of the embodiment shown in Fig. 4.

DETAILED DESCRIPTION

In a supported, or immobilized, liquid membrane device, a carrier liquid film is immobilized within the pores of a porous membrane. This porous membrane serves only as a framework or supporting layer for the liquid film. These membranes can be prepared by impregnating a porous membrane, typically hydrophobic, with a suitable organic solvent.

Thus, the present invention comprises a supported liquid membrane carrier liquid with a dielectric constant between approximately 20 and 32, and the use of that carrier liquid in a supported liquid membrane device to extract a chemical species from a feed liquid. In other embodiments of the invention, the dielectric constant of the carrier liquid is between approximately 21 and 30 or between approximately 22 and 28, and in a further embodiment of the invention, the dielectric constant is approximately 24.

When the overall dielectric constant of the carrier and the diluent mixture is below about 20 there is very little flux but a stable membrane. The flux continues to improve to a dielectric constant of about 24 with little change in membrane stability. Although the flux continues to improve with increasing dielectric constant of the carrier fluid beyond a dielectric constant of about 28, membrane stability begins to deteriorate appreciably.

In another embodiment of the present invention, mass transfer flux is described by the following equation:

15

10

5

$$J = K(A_F - (K_{d2}/K_{d1})A_S)$$

$$K_{d1} = A_1/A_i$$

20

25

30

35

$$K_{d2} = A_2/A_{si}$$

Where:

J is the mass transfer flux

K is the overall mass transfer coefficient

A is the activity coefficient of the species of interest in the feed solution

AF is the activity coefficient of the bulk feed species of interest

Asi is the strip side interfacial concentration of the species of interest

As is the activity coefficient of the bulk side species of interest

A₁ and A₂ are the activity coefficients of the bound species of interest at the feed and strip interfaces, respectively

K_d is the distribution coefficient.

 K_{d2}/K_{d1} is the ratio of the distribution coefficient on the strip side to that on the feed side.

The ratio K_{d2}/K_{d1} is preferably between approximately 1.0 and 0.0001, where K_{d2}/K_{d1} is the ratio of the distribution coefficient on the strip side to the distribution coefficient on the feed side. In one embodiment of the invention, the ratio K_{d2}/K_{d1} is between approximately 0.1 to 0.0001. In another embodiment of the invention, the ratio

 K_{d2}/K_{d1} is approximately between 0.1 to 0.001. In another embodiment of the invention, the ratio K_{d2}/K_{d1} is between approximately 0.25 and 0.5. In a preferred embodiment of the invention, the ratio K_{d2}/K_{d1} is approximately 0.33.

In another embodiment of the invention, a companion ion that is transported from the feed liquid to the strip liquid along with the ion of interest is removed from the strip liquid, thereby increasing transport of the ion of interest from the feed liquid to the strip liquid. The companion ion can be removed by a variety of means well known to those of ordinary skill in the art. Examples of methods of removing a companion ion include, but are not limited to, ion exchange, solvent extraction, and precipitation.

5

10

15

20

25

30

35

In use, the feed liquid contains a chemical species which is transported from the feed liquid through the carrier liquid in the pores of the membrane to the strip liquid (see Fig. 1). Two types of transport take place. In co-coupled transport, both a metal ion, for example, and its counter ion are transported from the feed solution through the supported liquid membrane (SLM) and into the strip solution. In coupled counter-transport, an acidic carrier, HC, loses a proton and forms a complex MC with the metal ion at the feed solution-SLM interface. This complex diffuses to the SLM-strip solution interface where it liberates the metal cation into the strip solution and simultaneously picks up a proton from the strip solution. The regenerated carrier HC diffuses back to the feed solution-SLM interface, picks up another metal ion and the process continues. The carrier molecule shuttles between the feed solution and the strip solution-SLM interfaces. The driving force in a counter-transport SLM is the pH differential between the feed and strip solutions.

A wide variety of materials can be used for the microporous liquid membrane support member, including hydrophobic materials such as microporous polypropylene [Celgard, Hoechst Celanese Corp., Fibers & Film Div., Charlotte NC 28209], Accurel ® (Enka, A.G., Wuppertal), polytetrafluoroethylene [Gore-Tex: W.L. Gore & Associates, Inc., Newark DE 19711], polyvinylidene fluoride, polystyrene, polyethylene, and hydrophilic materials such a microporous regenerated cellulose, cellulose acetate, cellulose acetate-nitrate cellulose triacetate, microporous glass and porcelain. Other materials, such as polybenzimidazoles, polysulfones, polyacrylonitriles, and silicon rubber, may also be used. Typically hydrophobic materials are used with organic liquid carriers, while hydrophilic materials are used with aqueous liquid carriers.

A chiral membrane may also be used to separate enantiomers. For example, a membrane of cellulose tris(3,5-dimethylphenylcarbamate) coated on Teflon may be used with the present invention. (See E. Yashima, *et al.*, <u>J. of Applied Polymer Science</u>, vol. 54, pp. 1087-1091 (1994), incorporated by reference. All literature citations are expressly incorporated herein by reference.

Membranes may be treated to alter their surface properties. For example, hydrophobic films of polyethylene may be treated with chromic acid, sulfuric acid or oxidizing agents to render them less hydrophobic. Typically however, for organic carriers and diluents, untreated polypropylene is used that is symmetrical with respect to both sides and with respect to the density and size of the micropores. Support members can be in the form of flat sheets or desirably hollow fibers which provide a high ratio of membrane surface area to volume of feed solution and strip liquid. To further improve the membrane surface area in contact with the carrier liquid, a fabric of hollow fibers is helically or spirally wound about a central core.

10

5

The microporous liquid membrane support members of the present invention are generally about 20 to about 125 microns in thickness, and desirably about 25 to about 100 microns in thickness. However, it is to be realized that the support members should be as thin as possible consistent with the need to be strong enough to withstand pressure differences across the support member. Some feed solutions, carrier liquids, and strip liquids may cause the membrane support member to swell but this generally does not effect operation of the liquid membrane if the support member remains sufficiently strong. The diameters of the pores of the microporous liquid membrane support members range in size from about 0.01 to about 1 micron with a size of about 0.1 microns typically used. The pores are generally uniform in size and have a uniform density across the membrane.

20

15

The carrier liquid is generally composed of a diluent and a carrier selective for the chemical species of interest. For an aqueous feed solution, the carrier liquid is typically a water immiscible liquid organic solvent. When the feed solution is an organic solution, the carrier liquid is typically an aqueous-based system. It has been found that membrane stability is associated with the overall dielectric constant of the carrier liquid. It is desirable that the dielectric constant be in the range of about 15 to 32, preferably 20 to 28, or 21 to 28, and more particularly, about 24. The diluents listed below, when used within those dielectric parameters, have been found effective with the present invention.

25

To increase the viscosity of the carrier liquid, and so decrease loss of the carrier liquid, and at the same time not decrease diffusivity, one may add a small percentage of a high molecular weight polymer. This technique is described in U.S. Patent No. 5,160,627 to E.L. Cussler, et al., incorporated by reference.

30

35

Appropriate diluents include aromatic hydrocarbons, aliphatic hydrocarbons, aldehydes, ketones, pyridines, ethers, crown ethers, nitriles, phosphoryls, or alcohols which are water immiscible and inert. Such diluents include aliphatic and aromatic hydrocarbons such as kerosenes (including aromatic and aliphatic ones), phenylcyclohexane, diphenylether, benzene, toluene, xylene, decane, dodecane, deconal,

nitrophenyl ethers (nitrophenyl pentyl ether through nitrophenyl decyl ether), diisopropyl benzene, ortho-xylene, dodecane, 2-octanone, 4-(1-butylpentyl)pyridine, anisole, benzonitrile, tributylphosphate, 1-octanol, methyl isobutyl ketone, cyclohexanone, tri-n-butyl phosphate, nonylphenol, di-2-ethylhexyl phthalate, 1-octyl-2-pyrrolidone, Soltrol 220 (paraffin oil from Phillips Petroleum Co.), Isopar M (aliphatic isoparaffinic kerosene: Exxon Chemical Co., Houston, TX), and Isopar V. Typically membrane stability tends to improve when a carrier liquid that is extremely hydrophobic is used. However, such membranes, as typified by a long-chained, hydrophobic diluent such as dodecane, tend to produce very little flux because of the very low diffusion coefficients found in such diluents. On the other hand, polar solvents tend to afford high diffusion coefficients but have low stability. To balance these factors, it has been found desirable to balance the polarity of non-polar solvents with the polarity of a more polar solvent.

5

10

15

20

25

30

35

Most carrier and diluent combinations typically last less than about five days, e.g., nitrophenylpentyl ether in a typical supported liquid membrane configuration. Use of nitrophenyloctyl ether alone as a diluent, improves the lifetime to about 10-20 days. Pure dodecane as a diluent gives a membrane life of about five days. However when a mixture of nitrophenyl ethers is used, such as a 80:20 volume percent mixture of nitrophenyloctyl ether and nitrophenylpentyl ether, the supported liquid membrane lifetime increases unexpectedly to more than 90 days. By adding an alkane such as dodecane, the flux of the preselected chemical species through the membrane structure is further increased through the membrane while reducing diluent cost considerably. In one embodiment of the invention, the ratio of nitrophenyl ether to nitropentyl ether is 3:1 by volume.

Typically a diluent mixture of about 50-80 volume percent nitrophenyloctyl ether, about 5-20 volume percent nitrophenylpentyl ether, and about 5-50 volume percent dodecane can be used with about 60-70 volume percent nitrophenyloctyl ether, about 10-15 volume percent nitrophenylpentyl ether and about 15-30 volume percent dodecane preferred, and about 65 volume percent nitrophenyloctyl ether, about 15 volume percent nitrophenylpentyl ether and about 20 volume percent dodecane most preferred. It is to be realized that other nitrophenyl ethers may be substituted, such as nitrophenylhexyl ether for nitrophenylpentyl ether and other alkanes used, *e.g.*, undecane and dodecane, with the present combination chosen on the basis of its stability, flux increase and low cost.

Suitable carriers include natural receptors, oximes, secondary and tertiary amines, macrocyclic polyethers, cobalt complexes, calixarenes, acidic organophosphorous compounds, carboxylic acids, and fatty acids. Examples of these compounds are valinomycin or beauverin, diethylhexylphosphoric acid, monododecylphosphoric acid, octaphenylphosphoric acid or bis(trimethylpentyl)phosphinic acid, and organophosphorous

10

15

20

25

30

35

acid esters such as trioctylphosphine oxide, tributyl phosphate, and tripentyl phosphate, fatty acids such as oleic acid, dinonylnaphthalenesulfonic acid, 38% by weight of 2hydroxy-5-nonylacetophenone oxime in kerosene available from Henkel Corp., Tucson, Arizona, as LIX84 (2-hydroxy-5-nonylacetophenone oxime), LIX 64N, LIX 860N, LIX 973N, LIX 984 (LIX 84 and LIX 984 combine an aldoxime and ketoxime in about a 1:1 mole ratio), uranyl salophene, Primene JM-T (a long-chain primary alkylamine from t-C₁₈H₃₇NH₂ to t-C₂₂H₄₅NH₂; Rohm & Haas), tetraoctyl ammonium bromide, trioctylmethyl ammonium chloride, dodecylamine, tridecylamine, tri-n-octylamine, triphenylamine, trioctylmethylammonium chloride, Amberlite-LA-2 (Rohm & Haas Chemical Co., Philadelphia, PA), Alamine 336 (a tertiary amine: General Mills, Inc., Minneapolis MN 55440), Aliquat 336 (methyltricaprylammonium chloride, a quaternary ammonium salt), alkylated cupferrons such as the ammonium salt of N-(alkylphenyl)nitrosohydroxylamine, beta-hydroxyoximes, beta-diketones, alkylated ammonium salts such as tridoddecylammonium chloride, oligoamides such as malonamide and glutaramide, and compounds such as 4-octadecyoxybenzaldehyde, Cyanex 272 (di-(2,2,4-(bis(2,4,4-302 trimethylpentyl)phosphinic acid), Cyanex trimethylpentyl)monothiophosphinic acid), Cyanex 301 (methylpentylphosphinodithioic acid) (American Cyanamid Co., Chemical Products Div., Wayne NJ 07470), tri-noctylphosphine oxide (TOPO) (Hokko Chemical Ind. Co. Ltd., Chuo-ku Tokyo, Japan), di-n-octyl sulfide, diethylcarbamoyl methyl phosphonate, n-octylphenyl N,Ndiisobutylcarbamoyl methylphosphine oxide, dinonylnaphthalene sulfonic acid, Nmethyl, N-butyl decanamide, N, N-di(n-octyl) hexanamide. Other suitable carriers are disclosed Wienck, M.M., et al., J. Am. Chem. Soc., 1990, vol. 112, pp. 797-801; C.Hill, et al., J. of Inclusion Phenomena and Molecular Recognition in Chemistry, vol. 19, pp. 399-408, 1994; W.F. van Straaten-Nijenhuis, et al., Recl. Trav. Chim. Pays-Bas, vol. 112, pp. 317-324 (6 June 1993); Nijenhuis, W.F., et al., J. Am. Chem. Soc., vol. 113, p. 3607 (1991); W.F. van Straaten-Nijenhuis, et al., J. Membr. Sci., vol. 82, p. 277 (1993); U.S. Patent No. 4,879,316 to Alexandratos, Spiro D., et al.; U.S. Patent No. 4,661,257 to Kreevoy, et al., U.S. Patent No. 5,443,731 to Moyer, B.A., et al., all incorporated by reference.

Crown Ethers

Crown ethers may also be used as carriers. Useful crown ethers include: 12-crown-4, 13-crown-4, 14-crown-4, 15-crown-5, 16-crown-4,16-crown-5, 18-crown-6, 19-crown-6, 20-crown-6, 21-crown-7, or 24-crown-8, which may have attached to them including but not limited to benzene rings (benzocrowns), cyclohexyl rings

10

15

20

25

30

35

(cyclohexylcrowns), alkyl groups (e.g., t-butyl or t-octyl groups), or hydroxyl, ketone, ether, phosphoryl, sulfoxide, amide, or pyridine substituents. Other crown ethers include dicyclohexano-18-crown-6, 4-tert-butylcyclohexano-15-crown-5 or bis-4,4'(5')[tert-butyl)cyclohexano-18-crown-6 from Parish Chemical Company (Vineyard, Utah), calix[4]crown-6, lariat ether amides, such as sym-dibenzo-16-crown-5-oxacetamide compounds. Typically a crown ether with lipophilic character is used as the carrier, since this property, namely, the tendency to remain in a solvent, is directly related to the stability of the liquid membrane. Such a property counteracts the slight but steady distribution of the carrier molecules into the strip and feed solutions over time, which eventually depletes the carrier from the liquid membrane. To improve the lipophilic character of a crown ether such as dicyclohexano-18-crown-6, hydrophobic groups such as aliphatic groups are added to the basic carbon framework of the crown ether. Typical aliphatic groups include t-butyl, t-octyl, n-octyl, 2-ethylhexyl, isooctyl, isodecyl, isotridecyl, decyl, and dodecyl.

Carriers such as dithizone may be used in separating metal ions such as cadmium, copper, lead, mercury, or zinc, while carriers such as thioxine may be used in separating metal ions such as antimony, arsenic, bismuth, cadmium, copper, cobalt, gallium, gold, indium, iridium, iron, lead, manganese, mercury, molybdenum, nickel, osmium, palladium, platinum, rhenium, rhodium, ruthenium, selenium, silver, tantalum, tellurium, thallium, tin, tungsten, vanadium, or zinc. Where it is desired to separate a neutral species such as an organic molecule, e.g., an aromatic hydrocarbon from a hydrocarbon feed solution, the carrier for such an aromatic hydrocarbon may be, e.g., polyethylene glycol and benzene.

To separate enantiomers from the feed stream, a chiral carrier may be used. Such carriers, such as (S)-N-(1-naphthyl)leucine octadecyl ester, are described in U.S. Patent No. 5,080,795 to Pirkle, *et al.*, incorporated herein by reference.

Although a carrier in liquid form may be used as the carrier liquid, preferably the carrier liquid used in the device and processes of the present invention will contain from about 2 to about 70 percent by weight of the carrier, more preferably from about 10 to 50 percent by weight of the carrier, and most preferably from about 30 to 45 percent by weight of the carrier.

The preferred carrier will depend upon the preselected chemical species. For example, for copper and neodymium ions or zinc and copper ions, the preferred carrier liquid includes the acidic organophosphorous compounds such as diethylhexylphosphoric acid or trilauryl ammonium chloride.

Typical cation exchangers include carriers such as long chain alkyl, sulfonic, carboxylic, phosphoric, phosphoric, and phosphinic acids, beta-diketones, and

5

10

15

20

25

30

35

hydroxyoximes. Typical anion exchangers include carriers such as long chain alkylamines and their salts, phosphoric acid esters, and phosphine oxides while neutral carriers include crown ethers, phosphonates, and phosphine oxides. The crown ethers are typically dissolved in the diluent at concentrations ranging from about 1.0 x 10⁻⁵ molar to about 5 molar, with about 0.1 to 1 molar being a typical concentration.

The carrier liquid may contain a surfactant. Suitable surfactants can assist in stabilizing the mixtures, *i.e.*, reducing any tendency for the mixture to separate. The suitable surfactants are generally non-ionic surfactants with hydrophilicity-lipophilicity balance (HLB) numbers from about 8 to about 15, preferably from about 9 to about 10. Suitable surfactants include polyoxyalkylene, alkyl ethers, *e.g.*, polyoxyethylene lauryl ether, polyoxyalkylene alkyl phenols, polyoxyalkylene esters, sorbitan esters such as polyoxyalkylene sorbitan esters, polyoxyalkylene sorbitol esters, and polyols. Such surfactants can generally be added in amounts of about 1 to about 40 percent by weight based on the total amount of carrier, diluent, and surfactants.

The carrier liquid can contain additional materials which improve its overall operation. These materials include pyridines, amines, amides, ethers, nitriles, sulfoxides, phosphoryl compounds including, for example, phosphine oxides and the alkyl esters of phosphoric, phosphonic, and phosphinic acids, fluorocarbons, and alcohols. Such added materials are typically present in the diluent at a concentration ranging from about 0.01 molar to about 2 molar, with about 1.0 molar being preferred. The typical volume percent of the added materials is about 0.5 to about 50 percent by volume.

In the operation of the device 10, a feed liquid 20 containing the chemical species of interest 15 flows past a microporous membrane 50 impregnated with the carrier liquid 30, and the species of interest is transported to the strip liquid 40 on the other side of the membrane (See Fig. 1). When the device 10 is a hollow fiber contained module (see Fig. 2), the feed liquid 20 flows through the feed liquid tube 60, the strip liquid 40 flows through the strip liquid tube 70, and the carrier liquid 30 circulates in the module 80 on the outside of tubes 60 and 70.

Several supported liquid membrane devices may be connected in series, where the strip stream of one device becomes the feed stream of the following device. To maintain the driving force for chemical separation, if, for example, three devices were in series, they would transport the chemical species of interest across the membrane as a neutral species, a cation, and an anion. Of course, any of those could be first, second, or third. Therefore, the carriers would be anionic, cationic, or neutral carriers. Examples of these carriers are listed below.

Separate strip cells, each with a different strip liquid, can be used to separate two or more chemical species from the feed liquid. When using a hollow fiber configuration, it is not necessary to use a barrier to form separate cells. Rather, when hollow fibers are employed, the feed solution flows through the interior of one set of fibers while separate strip liquids flow through individual bundles of strip fibers with the carrier liquid circulating on the outside of each bundle of fibers.

Furthermore, when a chemical ion of interest is transported along with along with a second ion to the strip side of the microporous membrane, removal of the second ion will actually increase transport of the ion of interest to the strip liquid. For example, Na⁺ may be transported to the strip liquid along with TcO_4^- . Removal of the Na⁺ and replacement with Mg^{2+} to preserve ionic neutrality will unexpectedly increase transport of TcO_4^- .

A carrier liquid can be conditioned to improve the stability and operation of the supported liquid membrane device. This is accomplished by: (1) passing through a conditioner such as a purifier to remove impurities; (2) providing a supply of carrier liquid to replace lost carrier or diluent, or both; and (3) removing pre-selected ions when more than one pre-selected ion is chosen for separation using a device such as an ion exchange column. Purification can be accomplished by filters, sorption devices employing materials such as activated carbon, distillation apparatus, liquid-liquid extraction devices, ion-exchangers and other means. One or more preselected species can be removed from the carrier liquid prior to passage to the strip liquid. Devices such as an ion exchanger, liquid-liquid extraction unit, sorption devices, distillation apparatus, and others can accomplish this.

Similarly, the feed solution and strip liquid may also have a conditioning step in which a conditioner is used to remove membrane fouling impurities, or add and remove various species from these liquids to improve the operation of the overall device and process including improvement of the flux of the pre-selected chemical species. Thus, removal of various species from the feed solution or strip liquid using such techniques as ion exchange, activated carbon processing, or reverse osmosis can improve significantly the flux of the preselected chemical species through the supported liquid membrane. Similarly addition of various chemical species including non-preselected species to either the feed solution or strip liquid can also improve preselected chemical transport across the supported liquid membrane. Such additions and removals are made so as to maintain an overall ionic strength differential across the support liquid membrane structure that results in improved distribution coefficients of the preselected chemical species at both the feed and

strip side of the membrane structure and a resulting increase of preselected chemical species complex flux across the membrane.

The present invention contemplates the use of a wide variety of devices and methods for preparing the feed liquid so as to accommodate a wide variety of commercial applications, including, but not limited to, the cleanup of radioactive materials, contaminated soils, contaminated ground water, processing waste streams and other hazardous waste sites. Such preparation devices and methods include the use of supercritical solvents such as carbon dioxide or more conventional aqueous and organic solvent methods and devices, separation methods and devices including vacuum distillation devices to remove volatile substances including, but not limited to, organic vapors, perstraction devices and methods to separate on the basis of partial pressures, gas separators, concentrators, filters, and the like. Such techniques are especially attractive to the clean up of metals and organics at manufactured gas plants. As such the present invention contemplates multiple and simultaneous operation of a wide variety of pretreatment processing to prepare one or more feed solutions including both organic and aqueous feed streams of metals, aromatics such as benzene, toluene, ethyl benzene and xylene, and hydrocarbons.

The pH of an aqueous feed liquid can be varied depending upon the choice of the carrier. With carriers such as acidic organophosphorous compounds, organophosphorous acid esters, or beta-hydroxyoximes, the pH of the feedstream is generally within the range of about 3.0 to 6.0 and more preferably from about 4.0 to about 5.5. The pH of the strip liquid is generally within the range about 0.1 to about 2.0, more preferably from about 0.5 to about 1.0. The strip liquid can be an aqueous acidic solution, e.g., a nitric acid solution or other suitable acidic solution such as sulfuric acid. Some applications for metal transport may require extraction from basic solution, and complexing agents such as beta-diketones can be used with pH ranges greater than about 8.

Although the process and related device can be used at ambient temperatures or at higher or lower temperatures, a temperature gradient across the supported liquid membrane structure 20 (see Figure 1) has been found often to be effective in increasing the flux of the preselected chemical species across the liquid membrane structure. This may be achieved by heating the feed liquid, the carrier liquid, the strip liquid, or a combination of two of the solutions. Electrical heating coils, boilers, heat exchangers, heating jackets, water/steam baths or other heating devices can be used. Cooling of the feed, carrier, or strip liquids, or a combination of any two, can be done with a chiller, a heat exchanger, dry ice/solvent bath or other cooling device. Alternatively, heating and cooling can be accomplished by heating or cooling the feed carrier or strip compartments directly, as for

16

5

10

15

20

25

30

35

example by placing a heating element or heat exchanger directly into the feed, carrier or strip liquid compartments of the supported liquid membrane device. In certain situations it can be advantageous to heat or cool only the carrier liquid to establish the proper temperature gradient for the system.

Of course, various other process parameters must be considered in establishing the temperature gradient including the stability of materials in the feed solution including the preselected chemical species, the boiling points and volatility of the carrier and its diluents, and the cost of maintaining such gradients versus the decreased throughput times at ambient temperatures. Flux increases of about an order of magnitude are obtained when a 30-50 degree Celsius temperature gradient is maintained.

When using a hollow fiber contained liquid membrane device, the feed solution, carrier liquid, and strip solution may be maintained at flow rates of, for example, about 1 cm/sec to about 5 m/sec. The flow rate of a more viscous fluid may be maintained at a flow rate of 3-5 times that of a less viscous fluid. However, it is to be realized that such flows rates are not necessary to the practice of the invention and that a batch-type processing without feed, strip and carrier flows are contemplated as part of the present invention.

One feature of the present invention is the use of a crown ether carrier to separate a large anion from a mixture of the large anion and smaller anions in the feed solution. It has been found that when a crown ether is used as a carrier to carry a small cation across the membrane, the largest of the anions in the feed solution associates with the crown ether and cation and is carried across the liquid membrane with the carrier and cation. As a result it is possible to remove the largest of the anions from the feed, which gives the invention particular utility in separating large anions such as pertechnetate, perrhenate, permanganate, perchlorate, perbromate, periodate, iodide, tetracyanaurate, tetraphenylborate, benzoate, and picrate anions from feed solutions containing smaller anions. The process has been found to be particularly effective in separating radioactive anions such as pertechnetate from radioactive wastes that often contain substantial amounts of nitrate ion. For example, a crown ether carrier can form a cation complex with a sodium ion at the interface of the feed solution and the liquid membrane. To preserve charge neutrality with respect to the sodium cation and crown ether complex, and between the feed solution and the strip liquid, an anion also associates itself with the crown ether cation complex and is also transported to the interface of the strip solution and liquid membrane, where the complex dissociates with the anion and the cation, e.g., Na⁺, entering the feed liquid. The crown ether then returns to the interface of the feed solution and the liquid membrane. Because the pertechnetate anion, TcO₄-, is the largest anion in the feed

17

solution, it is selectively carried across the liquid membrane, while the large nitrate anion, $N0_3$ -, because it is smaller than the pertechnetate anion, remains in the feed solution.

While not wanting to be bound by the following hypothesis, the mass transfer flux (J) is given by the equation of Dr. Srinivas Kilambi:

5

$$J = K(A_F - (K_{d2}/K_{d1})A_S)$$

$$K_{d1} = A_1/A_i$$

10

20

25

30

$$K_{d2} = A_2/A_{si}$$

Where:

J is the mass transfer flux

K is the overall mass transfer coefficient

A is the activity coefficient of the species of interest in the feed solution

AF is the activity coefficient of the bulk feed species of interest

 A_{Si} is the activity coefficient of the strip side interfacial species of interest

A_S is the activity coefficient of the bulk side species of interest

A₁ and A₂ are the activity coefficients of the bound species of interest at the feed and strip interfaces, respectively

K_d is the distribution coefficient.

 K_{d2}/K_{d1} is the ratio of the distribution coefficient on the strip side to that on the feed side (See Figure 3).

The ratio K_{d2}/K_{d1} is preferably between approximately 1.0 and 0.0001, where K_{d2}/K_{d1} is the ratio of the distribution coefficient on the strip side to the distribution coefficient on the feed side. In a further embodiment of the invention, the ratio K_{d2}/K_{d1} is between approximately 0.1 to 0.0001. In another preferred embodiment of the invention, the ratio K_{d2}/K_{d1} is approximately between 0.1 to 0.001. In another embodiment of the invention, the ratio K_{d2}/K_{d1} is between approximately 0.25 and 0.5. In a preferred embodiment of the invention, the ratio K_{d2}/K_{d1} is approximately 0.33.

For a neutral preselected chemical species x, the flux equation for species x

is:

$$J_X = k_X(A_{X,S}/K_{X,F})A_{X,S})$$

10

15

20

25

30

35

where J_X is the flux of the preselected ion through the liquid membrane; k_X is the mass transfer coefficient of the preselected species and is a direct function of the diffusivity of the preselected species in the carrier liquid and the membrane porosity and an inverse function of the membrane thickness and tortuosity; $A_{X,F}$ is the activity coefficient of species on the feed side of the liquid membrane structure; $A_{X,S}$ is the activity coefficient of species x on the strip side of the liquid membrane structure; $K_{X,S}$ is the activity coefficient of species x in the strip liquid interface divided by its activity coefficient on the strip side of the liquid membrane structure; and $K_{X,F}$ is the activity coefficient of species x in the feed solution interface divided by its activity coefficient on the feed side of the liquid membrane structure. From this equation it is readily apparent that one can control the flux of the preselected species by increasing the concentration of species x in the liquid membrane on the feed side or at the bulk interface on the strip side of the membrane. Alternatively one can decrease the concentration of species x at the bulk interface on the feed side of the liquid membrane or in the liquid membrane on the strip side.

For ionic co-transfer, the flux equation can be written as:

$$J_X = k_X(A_{X,F}A_{CX,F} - (K_{X,S}/K_{X,F})A_{X,S}A_{CX,S})$$

where x is the preselected ionic species and cx is the oppositely charged ion that is cotransported to maintain charge neutrality. One may obtain an uphill transfer of species x against its concentration gradient by decreasing the concentration of the oppositely charged ion on the strip side to such a point that the concentration difference between the feed and strip side for the oppositely charged ion is greater than for the preselected species even though there is a greater concentration of preselected species x on the strip side than on the feed side of the membrane. That is, species x will transport against its concentration gradient so long as the concentration difference between the oppositely charged ion is greater than the concentration difference between the selected species going in the uphill direction. The concentration of the oppositely charged ion must be greater on the feed side than on the strip side and that concentration difference must be greater than the concentration difference between the preselected species on the strip side to that on the feed side.

As yet a further possibility and as noted previously, one can control further the flux of the preselected species by increasing the concentration of species x in the liquid membrane on the feed side or at the bulk interface on the strip side of the membrane. Alternatively or in combination, one can decrease the concentration of species x at the bulk

interface on the feed side of the liquid membrane or in the liquid membrane on the strip side.

For ionic counter-transfer, the flux equation can be written as:

$$J_X = K_X(A_{X,F}A_{CX,S} - (K_{X,S}/K_{X,F})A_{X,S}A_{CX,F})$$

5

10

15

20

25

30

35

where x is the preselected ionic species and cx is the oppositely charged ion that is countertransported to maintain charge neutrality. One may obtain an uphill transfer of species x against its concentration gradient by decreasing the concentration of the oppositely charged ion on the strip side to such a point that the concentration difference between the strip and feed side for the oppositely charged ion is greater than for the preselected species even though there is a greater concentration of preselected species x on the strip side than on the feed side of the membrane. That is, species x will transport against its concentration gradient so long as the concentration difference between the oppositely charged ion is greater than the concentration difference between the selected species going in the uphill direction. The concentration of the oppositely charged ion must be greater on the strip side than on the feed side and that concentration difference must be greater than the concentration difference between the preselected species on the strip side to that on the feed side.

As yet a further possibility and as noted previously for neutral and cotransport, one can control further the flux of the preselected species by increasing the concentration of species x in the liquid membrane on the feed side or at the bulk interface on the strip side of the membrane. Alternatively or in combination, one can decrease the concentration of species x at the bulk interface on the feed side of the liquid membrane or in the liquid membrane on the strip side.

Figs. 4-7 illustrate a module for carrying out supported liquid membrane separation using hollow fibers with multiple and parallel strip liquid flows. The device generally designated as 100 comprises cylinder 104 with a first end 130 and a second end 150. Inlets 110, 106, 112, and 114 and outlets 152, 154, 156, and 158 are provided for feed solution 32 and strip solutions 34, 35, and 39, respectively. Inlet manifold 101 and outlet manifold 160 are provided for maintaining the feed solution 32 and strip liquids 34, 35, and 39 as separate flows. The inlet manifold 101 is formed from plate 132, a circular divider 170, and wall dividers 172, 174, and 176. The dividers 170, 172, 174, 176, plate 132 and end 130 form impermeable cells 102, 103, 105, 107 that maintain and separate each of the fluids 32, 34, 35 and 39. A similar arrangement of dividers (not

10

15

20

25

30

35

shown) is used in conjunction with plate 134 and end 150 to provide cells that maintain separate outlet flows of feed solution 32 and strip liquids 34, 35, and 39.

As shown in Figure 4, bundles of hollow fibers are provided to each of the cells 102, 103, 105 and 107 typically by forming plates 132 and 134 with epoxy cement and imbedding the hollow fibers so that the interior of each fiber is open to the appropriate cell. As shown in Figures 5 and 6, the interior of fibers 140, 142, 144, and 146 are open to and convey feed solution 32, and strip liquids, 34, 35, and 39, respectively through chamber 180 after which the hollow fibers are sealed to a set of similarly situated outlet cells at the opposite end of chamber 180.

One of more open screens such as screen 190 (Fig. 5) may be used to provide support to the various bundles of hollow fibers. To improve the flux of the carrier complex, the groups of fibers can be formed into fabric and spirally or helically wound about a central core in chamber 180. Baffles and other flow diverters can be provided to increase the flow path of the carrier liquid 36 through chamber 180.

In operation, a cross flow of carrier liquid 36 enters at the bottom of chamber 180 through inlet 124 and flows over and around the exterior of the hollow fiber tubes 140, 142, 144, and 146 in chamber 180 and leaves though outlet 122. The carrier liquid 36 fills the micropores of all of these tubes. The feed solution 32 enters into the circular chamber defined by circular partition 170 through inlet 110 and then flows through the interior of microporous hollow fibers 146 where it contacts the carrier liquid 36 in the micropores of hollow fibers 146. The preselected chemical species form complexes with the carrier and are transported to the pores of the hollow fibers 140, 142, and 144 carrying the strip liquids 34, 35 and 39. The strip liquids 34, 35, and 39 flowing on the inside of hollow fibers 140, 142, and 144 contact the carrier liquid in the micropores of the hollow fibers and each removes (decomplexes) one or more different preselected chemical species from the complexes in the carrier liquid 36.

This invention is further illustrated by the following examples, which are not to be construed in any way as imposing limitations upon the scope thereof. On the contrary, it is to be clearly understood that resort may be had to various other embodiments, modifications, and equivalents thereof, which, after reading the description herein, may suggest themselves to those skilled in the art without departing from the spirit of the present invention.

Example 1

An initial feed of 300 parts per million (ppm) chromium (VI) was extracted using a carrier liquid having a 33 vol% Amberlite L-A2 carrier (N-

dodecyltrialkylmethylamine), i.e., an ion exchange resin, and a diluent of pure dodecane (67 vol%). This carrier liquid of Amberlite carrier and dodecane diluent had a dielectric constant of approximately 15 and afforded a flux of less than 1 g m²/hr. The stability of the membrane was relatively high, lasting for approximately 16 days. When 1 vol% nitrophenyl octyl ether was added to the carrier liquid (33% Amberlite carrier, 66% dodecane diluent and 1% nitrophenyl octyl ether), the resulting dielectric constant was approximately 20 and the flux had increased by a factor of 1.6. The general stability was about the same as the pure dodecane diluent. When a 5% solution of nitrophenyl octyl ether was used (33% Amberlite, 62% dodecane, and 5% nitrophenyl octyl ether), the overall dielectric constant of the carrier liquid increased to approximately 24 while the flux increased by a factor of 2.6 times over that of the pure dodecane diluent. The membrane was stable for approximately 18 days. When a 10 vol% nitrophenyl octyl ether solution with 57 vol% dodecane, and 33 vol% Amberlite was used, the flux increased by a factor of 4 times with the dielectric constant increasing to about 28. The membrane was stable for approximately 3 weeks. Finally, when pure nitrophenyl octyl ether was used in place of the dodecane, the resulting Amberlite and nitrophenyl octyl ether solution had a dielectric constant of more than 30 and provided the greatest flux; however the membrane was stable for less than a day.

20

25

30

35

15

5

10

Example 2

Similar results were obtained for nickel or zinc at an initial feed concentration of 300 ppm and a carrier liquid having a 30 vol% bis(2-ethylhexyl) hydrogen phosphate carrier and a diluent mixture of 60-70 vol% dodecane and 0-10 vol% of nitrophenyl octyl ether. The resulting carrier liquid had a dielectric constant of about 10 to about 15 and afforded a flux of less than about 1 gm²/hr. The membrane was stable for about 2 to about 3 days. With the carrier concentration held constant, the nitrophenyl octyl ether concentration was increased to about 40 vol% to about 70 vol% and the dodecane lowered to about 0 vol% to about 30 vol%, to afford a dielectric constant of about 30 to about 40. Such carrier liquid afforded an increase of flux of about two times that of the flux with a 10% nitrophenyl octyl ether diluent, and the membrane was stable for about 15 days.

-

Example 3

When a nitrophenyl octyl ether concentration of about 10 vol% to about 30 vol% and a 40 vol% to about 60 vol% dodecane solution was used, the dielectric constant of the carrier liquid was approximately 20-28, and a flux of 6 to 8 times that of the flux

using a 10% nitrophenyl octyl ether solution was achieved. The membrane was stable for approximately 10 to 15 days.

Example 4

A 200 ppm silver feed solution was extracted with a carrier of diethylsulfide. The diethylsulfide carrier was maintained at a concentration of 10% in the carrier liquid. When nitrophenyl octyl ether was used alone with the carrier, the dielectric constant of the resulting carrier liquid was approximately 24, and afforded a flux of more than 3 gm²/hr. The membrane was stable for more than 35 days. When dodecane (90%) was substituted for the nitrophenyl octyl ether, the dielectric constant was approximately 10 and the flux was less than 1 gm²/hr. Furthermore, the membrane was stable for less than 2 days.

Example 5

In this Example, the separation of hexavalent chromium from an acid feed solution using a carrier liquid employing an amine carrier is shown. In the first step of the reaction, the feed chromium reacts with the amine to form an amine complex, *i.e.*,

20

15

5

10

On the strip side of the process, the chromium reacts with a base in another exothermic reaction, i.e.,

$$2H_2CrO_4 + 2NaOH Na_2Cr_2O_7 + 2H_2O + Heat.$$

25

30

35

Since both the first and second reactions liberate heat and the greatest concentration of chromium ion is in the carrier liquid, cooling of the carrier liquid is a more efficient process than cooling either the feed liquid or strip liquid or both, since only a small volume of liquid needs to be cooled. The feed and strip liquid volumes are much larger than that of the carrier liquid.

Example 6

A supported liquid membrane device was established with a feed solution containing 50 ppm Na⁺ TcO₄⁻, and a strip solution containing 100 ppm Na⁺ TcO₄⁻. An ion exchange column was used to remove Na⁺ from the strip solution, and replace it with

23

Mg²⁺. During the course of the experiment, the concentration of TcO₄⁻ increased to 110 ppm, then 120 ppm, and continued to increase.

Those skilled in the art will now see that certain modifications can be made to the invention herein disclosed with respect to the illustrated embodiments, without departing from the spirit of the instant invention. And while the invention has been described above with respect to the preferred embodiments, it will be understood that the invention is adapted to numerous rearrangements, modifications, and alterations, all such arrangements, modifications, and alterations are intended to be within the scope of the appended claims.

5

20

25

30

Claims

- 1. A supported liquid membrane device for extracting a chemical species from a feed liquid comprising the feed liquid in a feed compartment and a strip liquid in a strip compartment, wherein the feed and strip compartments are separated by a porous membrane containing a carrier liquid with a dielectric constant approximately between 20 and 32.
- The device of Claim 1, wherein the dielectric constant is between approximately 22 and 28.
 - 3. The device of Claim 1, wherein the dielectric constant is approximately 24.
- The device of Claim 1, wherein the carrier liquid contains a polar solvent.
 - 5. The device of Claim 4, wherein the polar solvent is a nitrophenyl alkyl ether.
 - 6. The device Claim 5, wherein the nitrophenyl alkyl ether is nitrophenyloctyl ether or nitrophenylpentyl ether.
 - 7. The device of Claim 1, wherein the carrier liquid is a non-polar solvent.
 - 8. The device of Claim 7, wherein the nonpolar solvent is kerosene, benzene, toluene, xylene, decane, dodecane, or deconal, or mixtures of the nonpolar solvents.
 - 9. The device of Claim 7, wherein the carrier liquid is approximately 50 to 20 volume percent nitrophenyloctyl ether, approximately 5-20 volume percent nitrophenylpentyl ether, and approximately 5-50 volume percent.

PCT/US97/16386 WO 98/10852

25

The device of Claim 7, wherein the carrier liquid is 65 volume 10. percent nitrophenyloctyl ether, about 15 volume percent nitrophenylpentyl ether, and about 20 volume percent dodecane.

5

The device of Claim 1, wherein a temperature gradient is 11. maintained between any of the feed, carrier, or strip liquids.

10

The device of Claim 1, further comprising one or more supported 12. liquid membrane devices connected in series or parallel.

The device of Claim 1, wherein three supported liquid membrane 13. devices are connected in series and the carrier liquid of one device contains an anion exchange carrier, the carrier liquid of another device contains a neutral exchange carrier, and the carrier liquid of the other device contains a cation exchange carrier.

15

A method of extracting one or more chemical species from a feed 14. liquid comprising introducing the feed liquid through a supported liquid membrane device, the supported liquid membrane device comprising a feed compartment and a strip liquid in a strip compartment, wherein the feed and strip compartments are separated by a porous membrane containing a carrier liquid with a dielectric constant approximately between 20 and 32.

20

The method of Claim 14, wherein the dielectric constant is between 15. approximately 22 and 28.

25

The method of Claim 14, wherein the dielectric constant is 16. approximately 24.

polar solvent. 30

The method of Claim 14, wherein the carrier liquid contains a 17.

The method of Claim 17, wherein the polar solvent is a nitrophenyl 18. alkyl ether.

26

- 19. The method of Claim 18, wherein the nitrophenyl alkyl ether is nitrophenyldecyl ether, nitrophenyl nonyl ether, nitrophenyloctyl ether, nitrophenyl hexyl ether, or nitrophenylpentyl ether.
- 20. The method of Claim 14, wherein the carrier liquid is a non-polar solvent.

5

10

15

20

25

30

35

- 21. The method of Claim 20, wherein the nonpolar solvent is kerosene, benzene, toluene, xylene, decane, dodecane, or deconal, or mixtures of the nonpolar solvents.
- 22. The method of Claim 21, wherein the carrier liquid is approximately 50 to 20 volume percent nitrophenyloctyl ether, approximately 5-20 volume percent nitrophenylpentyl ether, and approximately 5-50 volume percent.

23. The method of Claim 21, wherein the carrier liquid is 65 volume percent nitrophenyloctyl ether, about 15 volume percent nitrophenylpentyl ether and about 20 volume percent dodecane.

- 24. The method of Claim 14, further comprising the step of maintaining a temperature gradient between any of the feed, carrier, or strip liquids.
- 25. The method of Claim 14, further comprising connecting one or more supported liquid membrane devices in series or parallel.
- 26. The method of Claim 14, wherein the supported liquid membrane device includes a feed liquid in a feed compartment, a carrier liquid compartment, and a strip liquid in a strip liquid compartment, further comprising the step of conditioning any of the feed, carrier, or strip liquids.
 - 27. A method of extracting an ion from a feed liquid comprising:
- (1) using a supported liquid membrane device including a feed liquid and a strip liquid to transport the ion and a companion ion from the feed liquid to the strip liquid; and
 - (2) removing the companion ion from the strip liquid.

27

28. A method of extracting one or more chemical species from a feed liquid comprising the step of using a supported liquid membrane device where the ratio K_{d2}/K_{d1} is preferably between approximately 1.0 and 0.0001.

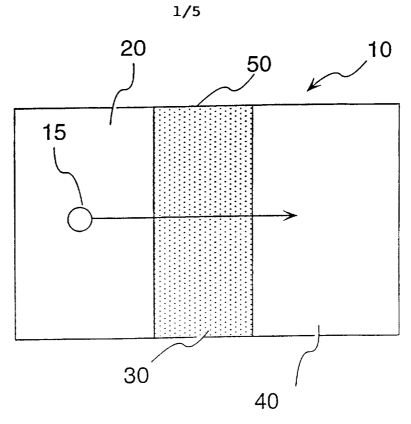
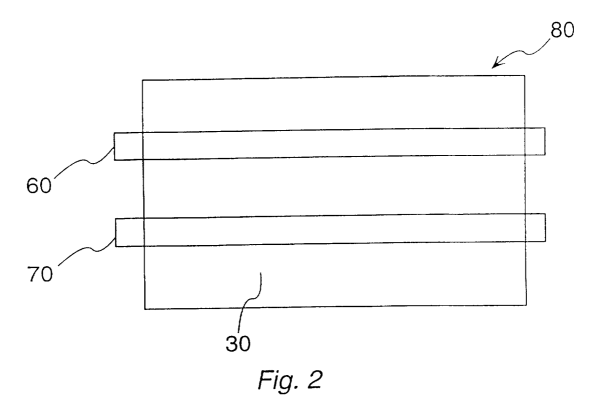


Fig. 1



SUBSTITUTE SHEET (RULE 26)

2/5

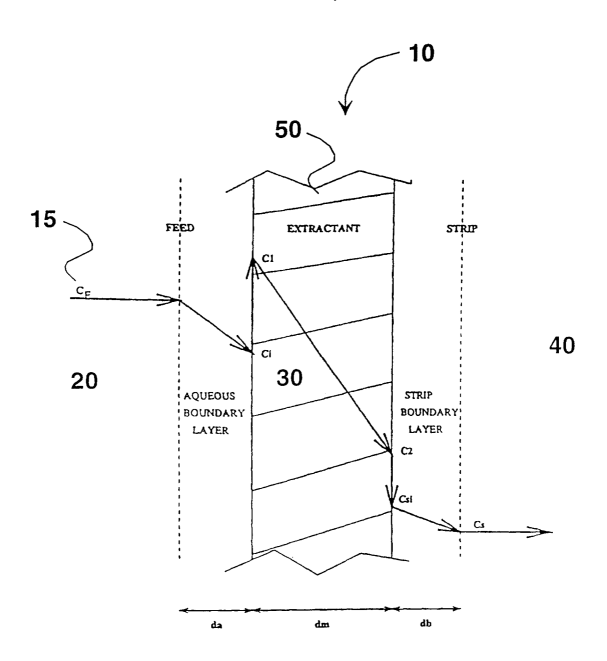
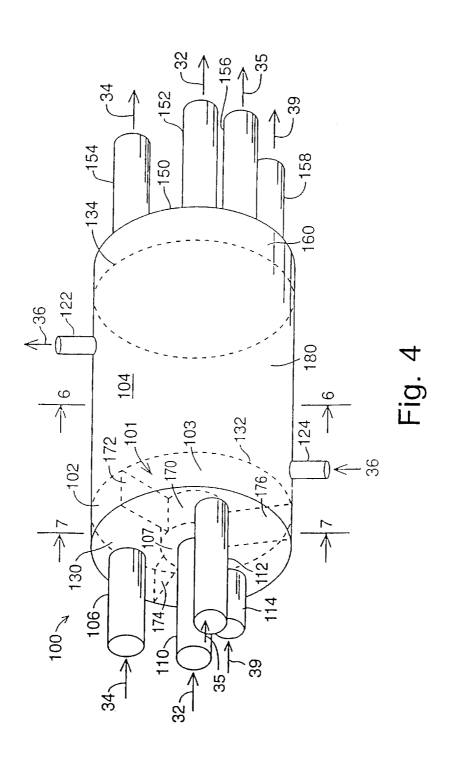
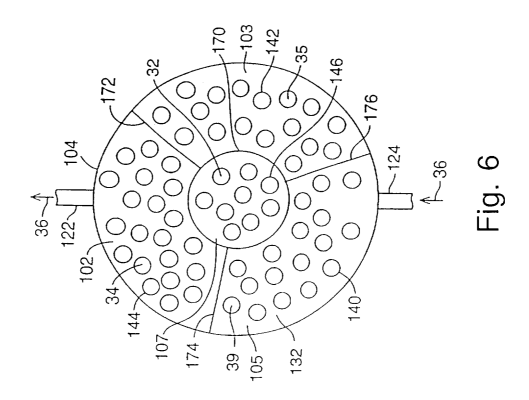
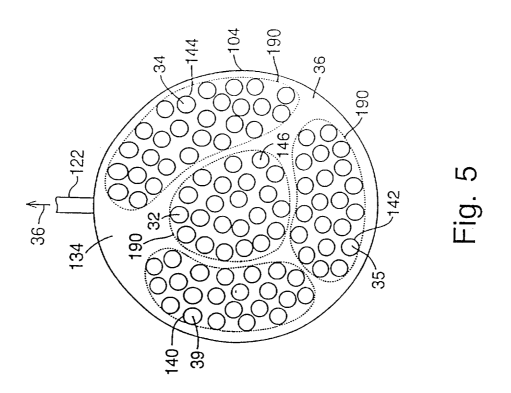


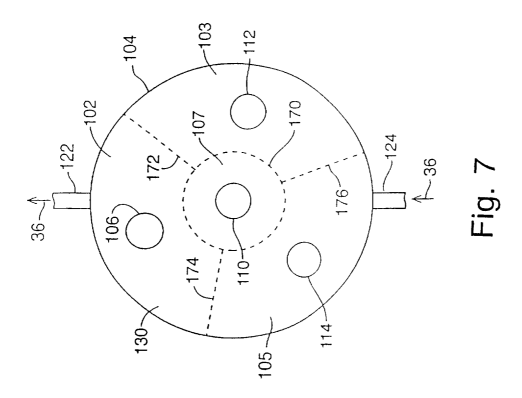
Fig. 3



4/5







INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/16386

A. CLASSIFICATION OF SUBJECT MATTER								
IPC(6) :BO1D 17/00 US CL :210/643, 640, 638, 654, 321.72; 96/5, 11								
According to	International Patent Classification (IPC) or to both n	ational classification and IPC						
	DS SEARCHED							
	ocumentation searched (classification system followed	by classification symbols)						
U.S. : 210/643, 640, 638, 654, 321.72; 96/5, 11								
Documentat	ion searched other than minimum documentation to the	extent that such documents are included	in the fields searched					
Electronic d	ata base consulted during the international search (nar	ne of data base and, where practicable,	search terms used)					
C. DOC	UMENTS CONSIDERED TO BE RELEVANT							
Category*	Citation of document, with indication, where app	ropriate, of the relevant passages	Relevant to claim No.					
X 	US 5,114,579 A (TAKIGAWA) 19 Ma	ay 1992, entire disclosure.	1-3, 7, 8, 12, 14- 16, 20-21, 25, 27					
Y			11, 28					
X	US 5,592,973 A (PEMSLER et al) 03	1-3, 7, 8, 16-16, 20, 21						
Y	US 4,617,125 A (DANESI) 14 October 1986, abstract, column 3, lines 26-68, and column 4, lines 1-68.							
Y	US 5,189,189 A (MISAWA et al) 2 column 4, lines 31-48, c olumn 5, line	1-3, 7,, 8, 14-16, 20						
X Furt	her documents are listed in the continuation of Box C.	See patent family annex.						
Special extensions of cited documents: "T" later document published after the international filing date or priority								
'A' de	date and not in contract with the application out cited to understand							
E earlier document published on or after the international filing date *X* document of particular relevance; the claimed invention cannot considered novel or cannot be considered to involve an inventive a								
ci	secument which may throw doubts on priority claim(s) or which is ted to establish the publication date of another citation or other	when the document is taken alone "Y" document of particular relevance; the						
'O' de	ecial reason (as specified) cument referring to an oral disclosure, use, exhibition or other eans	considered to involve an inventive combined with one or more other suc being obvious to a person skilled in	ch documents, such combination					
•P• d	ocument published prior to the international filing date but later than the priority date claumed	*&* document member of the same pater	nt family					
	Date of the actual completion of the international search Date of mailing of the international search report							
04 JANUARY 1998 0 3 FEB 1998								
Commissi	Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Authorized officer School Sc							
Box PCT Washingto	Box PCT Washington, D.C. 20231 ANA FORTUNA							
Facsimile No. (703) 305-3230 Telephone No. (703) 308-3857								

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US97/16386

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Category	Chanon of document, while indication, where appropriate, of the reservant passages	THE PARTY OF THE P
Y	US 5,332,531 A (HORWITZ et al) 26 July 1994, abstract, column 5, lines 44-47.	1-3, 14-16
Y	US 5,080,795 A (PIRKLE et al) 14 January 1992, abstract, and column 8, lines 9-28.	1-4, 7-8,14-16, 20, 26-28
E	US 5,607,591 A (DOZOL et al) 04 March 1997, entire disclosure.	4-6, 9, 10, 11, 14 17 -19, 22 and 2