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Apparatus and method for separating oxygen from air

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United States Patent [191]

Tabatabaie-Raissi

[54) APPARATUS AND METHOD FOR SEPARATING OXYGEN FROM AIR

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- [73) Assignee: University of Central Florida, Orlando, Fla.
- (21) Appl. No.: 919,153
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Related U.S. Application Data

- [63) Continuation-in-part of Ser. No. 637,777, *Jan:* 7, 1991, abandoned.
- [51) Int. *Cl.s* .. C25B 1/02
- [52) U.S. Cl ... 204/129
- [58) Field of Search 204/59 R, 258, 263,
- 204/283, 129; 55/2; 429/30, 31

[56) References Cited

U.S. PATENT DOCUMENTS

OTHER PUBLICATIONS

[111 Patent Number: 5,296,110 [45) Date of Patent: Mar. *22,* 1994

Fee et al.; "Monolithic Fuel Cell Development"; Argonne National Laboratory; pp. 40-43.

Primary Examiner-Kathryn Gorgas *Attorney, Agent, or* Firm-James H. Beusse

[57) ABSTRACT

A process and apparatus for separating and concentrating breathing-grade oxygen, that is therapeutically equivalent to 100% pure oxygen, from ambient air is provided. The oxygen concentrating process employed in the method of the invention is implemented in a housing having a main chamber. A solid anion conducting membrane is situated in the main chamber so as to divide the chamber into separate first and second reaction chambers. Electrocatalytically active cathodic and anodic electrodes are situated on the respective opposed surfaces of the membrane. A direct current source is coupled between the cathodic and anodic electrodes such that when ambient air is provided to the cathodic electrode, therapeutically pure and moist oxygen is produced at the anodic electrode by electrolytic action of hydroxyl ions passing through the solid membrane. The oxygen concentrator advantageously operates at room temperature making it well suited for production of therapeutic oxygen especially for applications where portable oxygen source is required.

2 Claims, 3 Drawing Sheets

Sheet 1of3 **5,296,110**

FIG. 5

APPARATUS AND METHOD FOR SEPARATING OXYGEN FROM AIR

separating a selected gas from a mixture of gases and, 10 more particularly, to an apparatus and method for sepa-

ers are known. Fuel cells typically include cathode and 15 tionally, such alkaline concentrator systems are also anode electrodes situated in, and separated by, an elec-
susceptible to contamination of the output oxygen by anode electrodes situated in, and separated by, an electrolyte. The electrolyte may be either solid or liquid. In trolyte. The electrolyte may be either solid or liquid. In the hazardous alkali vapor emitted from the caustic contrast to batteries which actually store electrical en-
electrolyte. Elaborate components are often required contrast to batteries which actually store electrical en-
electrolyte. Elaborate components are often required to
ergy in a chemical form, fuel cells are structures in provide electrolyte recirculation in these concentrato which the reactants and the reactant products are con- 20 as well as to provide heat exchange and filtration.

for all these reasons, prior electrolytic oxygen con-

for all these reasons, prior electrolytic oxygen continuously supplied and removed to produce electricity. For all these reasons, prior electrolytic oxygen con-
More specifically, a fuel and an oxidant are continu-
centrators have been very complex and generally inap-More specifically, a fuel and an oxidant are continuously supplied to the fuel cell so as to react and thereby directly produce an electrical potential between the cathode and anode electrodes. Thus, a fuel cell is gener- 25 ally regarded as being an energy converter whereas a battery is considered to be an energy source.

Electrolyzers, like fuel cells, also employ cathode and anode electrodes situated in an electrolyte. However, in an electrolyzer a source of DC current is supplied be- 30 and pressures. tween these electrodes such that the electrolyte is separated or decomposed into its component ions via electrolytic action.

ions (anions or cations) are transported. Electrolytic 45 is easily portable. Devices which electrolytically concentrate gases are related in general principle to the fuel cell and the elec- 35 trolyzer technologies discussed above. Fuel cells or electrolyzers are generally classified according to the type of electrolyte which they use to sustain electrolytic conduction between the cathode and anode thereof. Thus, we have either acidic or alkaline fuel cells or 40 electrolyzers depending on the particular electrolyte employed. These acidic or alkaline types of fuel cells or electrolyzers employ negative and/or positive charge ion conducting electrolytes through which charged systems most often used in the past contained electrolytes that were aqueous or molten salt solutions. Examples of such systems include alkaline (KOH or NaOH) electrolyzers and fuel cells, molten carbonate electrolyzers and phosphoric acid fuel cells. Modem electro- 50 lyzers and fuel cells contain solid state electrolytes fabricated from organic compounds such as NAFION (perfluorinated sulfonate ionomer) which is manufactured by DuPont or are fabricated from inorganic compounds such as zirconium oxide.

Electrolytic gas concentrator devices, such as oxygen concentrators, are known in the prior art and typically include a cathode and anode electrode situated in an electrolyte, in a manner somewhat similar to the fuel cell and electrolyzer structures discussed above. In such 60 further situated in the first reaction chamber. The appaoxygen concentrators, a source of DC voltage is typically coupled between these electrodes. These oxygen concentrators are known to employ either liquid or solid electrolytes. The nature of the selected electrolyte between the cathode and anode electrodes is such that 65 ing to the first reaction chamber. An oxygen outlet is gases other than oxygen cannot be transferred electrolytically through the electrolyte. Thus, the gas obtained at the anode is substantially pure oxygen whereas the

gas obtained at the cathode is a nitrogen rich sample of air. Electrolytic oxygen concentrators capable of producing medical grade oxygen for patient use are very desirable; however, the processes and structures dis-This is a continuation-in-part of application Ser. No. 5 cussed above have a number of deficiencies which make
07/637,777, filed Jan. 7, 1991, now abandoned. them generally inappropriate for this application. For them generally inappropriate for this application. For example, electrolytic oxygen concentrators which em-BACKGROUND OF THE INVENTION example, electrolytic oxygen concentrators which em-
ploy solid oxide electrolytes typically require very ex-This invention relates in general to techniques for pensive, all solid state fabrication and employ rare earth parating a selected gas from a mixture of gases and, 10 elements. Additionally, these concentrators often operate at relatively high temperatures to assure adequate rating oxygen from other gases by employing electro-
lytic systems.
alkaline oxygen concentrator systems are susceptible to tic systems.
Electrolytic systems such as fuel cells and electrolyz-
carbonation by carbon dioxide in the ambient air. Addicarbonation by carbon dioxide in the ambient air. Additionally, such alkaline concentrator systems are also provide electrolyte recirculation in these concentrators

propriate for low cost home and ambulatory patient oxygen use.

SUMMARY OF THE INVENTION

Accordingly, one object of the present invention is to provide a process for electrolytically concentrating oxygen which operates at or near ambient temperatures

Another object of the present invention is to provide a process for electrolytically concentrating oxygen which substantially eliminates the need to replenish the electrolyte.

Yet another object of the present invention is to provide a process for electrolytically concentrating oxygen which does not require undue maintenance.

Still another object of the present invention is to provide a process for electrolytically concentrating oxygen which is relatively low cost and suitable for concentrating oxygen for therapeutic patient use.

A still further object of the invention is to provide an apparatus for concentrating oxygen which exhibits a sufficiently small weight and volume that the apparatus

Yet another object of the invention is to provide an apparatus for concentrating oxygen which is capable of operating on relatively low DC voltage such as the 12 volts which is available in most vehicles.

In accordance with the present invention, the process is implemented in an apparatus for separating mostly oxygen from air, such apparatus including a housing enclosing a main chamber. The apparatus further includes a solid anion conducting material electrolyte 55 having opposed first and second surfaces. The electrolyte is situated within the main chamber and divides the main chamber into separate first and second reaction chambers. The apparatus also includes a cathode electrode situated on the first surface of the electrolyte and ratus still further includes an anode electrode situated on the second surface of the electrolyte and further situated in the second reaction chamber. An air inlet provides air from outside the housing through the housprovided for exhausting oxygen from the second reaction chamber through the housing to outside the housing. A direct current source is coupled between the

cathode and anode electrodes such that when air is provided to the cathode electrode, oxygen is produced at the anode electrode by electrolytic action of hydroxyl ions or protons passing through the electrolyte.

One important feature of this apparatus is that a small 5 portion of ambient air in the first reaction chamber is allowed to directly pass through, around the rim of the membrane, to the second reaction chamber, diluting the output oxygen in a concentration range varying between no less than 85% (by volume) oxygen and no 10 more than 95% (by volume) pure oxygen.

BRIEF DESCRIPTION OF THE DRAWINGS

The features of the invention believed to be novel are specifically set forth in the appended claims. However, 15 the invention itself, both as to its structure and method of operation, may best be understood by referring to the following description and accompanying drawings:

FIG. I is a block diagram of a planar embodiment of an electrolytic oxygen concentrator using the process of 20 the present invention;

FIG. 2 is a simplified cross-sectional view of the electrolytic oxygen concentrator process of FIG. I taken along section line $2-2$;

FIG. 3 is a more detailed cross-sectional view of the 25 oxygen concentrator process of FIG. I;

FIG. 4 is a schematic view of a portable system embodiment of the electrolytic concentrator of the present invention; and

FIG. 5 is an enlarged view of the electrolysis module 30 of FIG. 4.

DETAILED DESCRIPTION OF THE INVENTION

FIG. I shows a block diagram of an electrolytic oxy- 35 gen concentrator 10 for implementing the process of the present invention. In this particular embodiment of the invention, concentrator IO is depicted as having a parallelepiped/planar configuration. Those skilled in the art will appreciate that concentrator 10 may exhibit other 40 geometries, such as cylindroidal for example, as long as such other geometries are consistent with the teachings herein.

Concentrator IO includes a housing IS which in one embodiment is fabricated in the form of a two piece 45 between current collecting electrodes SS and 60. More LUCITE (polymethyl methacrylate) shell including housing portions ISA and ISB. LUCITE is a trademark of its respective owner.) These housing portions ISA and ISB are sandwiched together in a planar relationship as depicted in FIG. I. A main chamber (not visible 50 in FIG. I) is situated within housing IS for implementing the later described electrolytic reaction.

Air is supplied to inlet pipe 2S and is withdrawn from outlet pipe 30 as indicated in FIG. I. Substantially pure oxygen in the range of SS to 95 percent by volume is 55 exits oxygen outlet pipe 3S. The operation of concentrator IO both internal and external to housing IS is discussed in detail in the subsequent discussion of FIG. 2.

FIG. 2 shows a cross-sectional view of the electrolytic oxygen concentrator IO of FIG. I taken along 60 section line 2-2. Housing 15 of concentrator 10 includes housing portions ISA and ISB which mate with each other along interface I7 as depicted in FIG. I. Returning to FIG. 2, it is seen that the aforementioned main chamber within housing 15 is actually two sepa- 65 rate reaction chambers, namely a cathodic chamber 20A which is separated from an anodic chamber 20B. As seen in FIG. 2, air inlet pipe 2S leads from the ambi-

ent surroundings of concentrator IO into cathodic chamber 20A and supplies chamber 20A with a fresh supply of air. Air outlet pipe 30 leads from chamber 20A to the ambient surroundings such that already processed air is exhausted from chamber 20A as illustrated. Oxygen outlet pipe 3S leads from chamber 20B to provide separated oxygen to the user.

In this particular embodiment, a solid electrolyte 40 is situated substantially centrally within chamber 20 and is held in position between upper housing walls 15A and ISB at the top 40A of electrolyte 40, and is further held in position between lower housing walls 15C and 15D at the bottom 40B of electrolyte 40. Electrolyte 40 includes opposed major surfaces 40C and 40D wherein surface 40C faces air inlet pipe 2S and outlet pipe 30, and wherein surface 40D faces pure oxygen outlet pipe 3S.

Electrolyte 40 is preferably fabricated from an anion conducting material such as an anion conducting membrane. Anions are defined as those ions which are attracted to the anode of an electrolytic chemical system. Alternatively, electrolyte 40 could be fabricated from a cation conducting material, such as Nation. More particularly, a solid polymer electrolyte (SPE) material such as fluorocarbon polymer may be employed as electrolyte 40. That is, solid electrolyte perfluorosulfunic acid type membranes may be employed as electrolyte 40. One anion conducting membrane material which has been found to be suitable for use as electrolyte 40 is the TOSFLEX (sulfonamide derivativce of perfluorocarbon sulfonic acid ionomer) membrane manufactured by the Tosoh Company of Japan. This membrane material is a solid polymer material.

A catalytic electrode 4S is situated on electrolyte surface 40C. Catalytic electrode 4S is the cathode of concentrator IO. Another catalytic electrode SO is situated on electrolyte surface 40D. Catalytic electrode SO is the anode of concentrator IO.

A current collecting electrode SS is situated on catalytic electrode 4S as shown. Another current collecting electrode 60 which is perforated is situated on catalytic electrode SO.

A source 6S of direct current (DC) power is coupled specifically, a battery is employed as source 6S. The positive terminal of such battery is coupled to current collecting electrode 60 via a switch 70. The negative terminal of the battery is coupled to current collecting electrode SS via a rheostat 7S which can be adjusted to control the amount of electric current supplied to concentrator IO.

It is thus observed in FIG. I that a planar sandwichlike structure is formed by current collecting electrodes SS and 60, catalytic cathode and anode electrodes 4S and SO, and solid polymer electrolyte 40.

Concentrator IO employs the following chemical reactions to separate oxygen from air. In this particular embodiment of the invention, the process is a 4 electron process. More particularly, the reaction at cathode 4S is summarized as:

EQUATION 1

O₂ (from air in 20A) + 2 H₂O + 4e (from 55) - 40H -

Upon the occurrence of the reaction described in Equation 1, the OH^- anion species permeates and migrates through electrolyte 40 to anode 50 where the reverse of the above reaction occurs as described by Equation 2.

Oxygen is thus produced in reaction chamber 20.B and is output pipe 35. A mixture of air gases, now less oxygen exhausted out output pipe 35 for therapeutic patient use. It should be noted that a portion of the air in the first 10° haust pipe 30. Air is continually circulated through inlet reaction chamber 20A is allowed to flow through the sine 25 and across electrods 55 and out out reaction champer 20A is allowed to flow through the pipe 25 and across electrode 55 and out outlet pipe 30 to current collecting electrode 55 and catalytic electrode 45, around the rim of the electrolyte 40 and through the $\frac{100}{100}$ current collecting electrode 60 and catalytic electrode 50, to the second reaction chamber 20B, diluting the 15 cally equivalent to 100% pure oxygen from air has been
output oxygen to a level between no less than 85% (by
determined about air has a mercanistal that a mathed output oxygen to a level between no less than 85% (by described above, it will be appreciated that a method volume) and no more than 95% (by volume) pure αx .

FIGS. 1 and 2 is shown in FIG. 3. Like numbers are 20 solid anion conducting material electrolyte including a cath-
used to indicate like elements in FIGS. 1, 2 and 3, rectable providing a cath-surfaces and providing a ca ferred to jointly. The electrolytic reaction generates method further includes the steps of providing an anode heat at cathode 45 which must be dissipated. For this heat at cathode 4S which must be dissipated. For this electrode on the second surface of the electrolyte and purpose, current collecting electrode 55 which is adjapurpose, current collecting electrode SS which is adja-
cent cathode 4S includes a plurality of coolant carrying 25 providing a direct current source coupled between the
channels or duets 57 . Channels 57 are intersecon channels or ducts 57. Channels 57 are interconnected electrode such that collecting electrode SS such that collecting cathode electrode such that covagen is produced at the

such that collectrode is such that collected at the same and electrode by electrolytic action of hydr flows throughout electrode 55 to remove heat there-
from Water may be englanded as the angles for each passing through the electrolytic. This method is advanfrom. Water may be employed as the coolant for con-
contrator 10, As seen in FIG, 3, the unnermost obannel 20, tageously performed at room temperature although centrator 10. As seen in FIG. 3, the uppermost channel 30 tageously performed at room temperature although
57 is coupled to a heat exchanger 90 by appropriate satisfactory performance may also be obtained within 57 is coupled to a heat exchanger 90 by appropriate provided by the coolant carried by the coolant is the temperature range of approximately 20 degrees
removed from the coolant hefore the coolant reaches a Celsius to approximately 100 degrees Celsius. The optiremoved from the coolant before the coolant reaches a Celsius to approximately 100 degrees Celsius. The opti-
storage tapk 95 connected to best exchanger 90. Heat mum electrolyte temperature is expected to be within storage tank 95 connected to heat exchanger 90. Heat exchanger 90 is connected to compartment 95 to permit 35 the range of approximately 80 degrees Celsius to ap-
the storage of cooled coolant therein A nump 100 proximately 85 degrees Celsius. the storage of cooled coolant therein. A pump 100 proximately 85 degrees Celsius.
(FIG. 1) is connected to storage tank 95 and to the It is difficult to maintain electrolytic membrane 40 at (FIG. 1) is connected to storage tank 95 and to the It is difficult to maintain electrolytic membrane 40 at lowermost channel 57 in concentrator 10 to numn cool. 100% relative humidity at temperatures above 100 delowermost channel 57 in concentrator 10 to pump cool-
ant back into concentrator 10. In this manner, a continuuting rees Celsius. For the membrane electrolyte 40 to operant back into concentrator 10. In this manner, a continu-
ous flow of coolant is provided through channels $57 \text{ to } 40$ ate properly, it is necessary to keep electrolyte 40 fully ous flow of coolant is provided through channels 57 to 40 remove heat from current collecting electrode SS and saturated so that undesired membrane dryout and the cathode 4S. resulting loss of ionic conductivity therethrough do not

between the uppermost portion of electrolyte 40 and the concentrator 10 depends on several factors such as the lowermost portion of electrolyte 40 by appropriate 45 gas pressure at cathode reaction chamber 20A formed lowermost portion of electrolyte 40 by appropriate 45 piping. Storage tank 95B and the connecting piping to by the interface of channels 59 with cathode 45 and the electrolyte 40 are filled with water to keep the solid anode reaction chamber 20B formed by the interface of electrolyte 40 are filled with water to keep the solid anode reaction chamber 20B formed by the interface of polymer electrolyte 40 fully saturated as discussed later channels 62 with anode 50, as well as the ionic conducpolymer electrolyte 40 fully saturated as discussed later

are situated extending throughout current collecting ment system such as structures SS, S7, 90, 9SA, and 100, electrode SS. Channels S9 open upon cathode 4S such and further to provide an electrolyte moisture managethat fresh air is provided to cathode 45 at which the ment system such as structures 40 and 95B.
reaction of Equation 1 takes place. Inlet pipe 25 is con-
The minimum voltage required to elecnected to the uppermost air passage 59 to provide fresh 55 separate oxygen from air is determined from the Nemst air thereto and to the remaining channels 59 connected equation (Equation 3). thereto. Outlet pipe 30 is coupled to the lowermost air passage 59 to allow spent air to escape from concentrator 10. Air channels 59 are interconnected to permit fresh air to flow among channels 59 and reach cathode 60 4S.

A plurality of oxygen exhaust passages or channels 62 are situated extending throughout current collecting electrode 60. Channels 62 are interconnected and open upon anode 50 such that oxygen, produced at anode 50 65 as per the reaction of Equation 2, is provided an exit route from concentrator 10. More specifically, channels 62 are connected to oxygen outlet pipe 3S through

which oxygen is exhausted for collection or other use as desired.

EQUATION 2 **In accordance with the invention, only hydroxyl ions**
 (QU_n) minima from antihiis algebra of f (the set) $(OH⁻)$ migrate from catalytic electrode 45 (the cath-*5* ode) through solid anion conducting polymer electro-40H-.....02 (liberated to 20B)+2 H20+4e (from 60) lyte 40 to catalytic electrode 50 (the anode). Thus, only oxygen is thus produced in reaction chamber 20B and is output pipe 35. A mixture of air gases now less oxygen rich, remains at electrode 55 for exhausting out of exprovide the oxygen source required for concentrator

While a process for producing oxygen therapeutivolume) and no more than 95% (by volume) pure oxy-
gen.
closed. This mathed includes the state of me, it'll $\frac{1}{2}$ more detailed cross-section of concentrator 10 of closed. This method includes the steps of providing a closed. This method includes the steps of providing a closed. ode electrode on the first surface of the electrolyte. The

Also as seen in FIG. 3, storage tank 95 is connected occur. It is noted that the best operating temperature for
tween the uppermost portion of electrolyte 40 and the concentrator 10 depends on several factors such as the in more detail. tivity of membrane electrolyte 40. For these reasons, it
A plurality of channels or cross flow air passages 59 50 is essential to provide an electrolyte thermal manageis essential to provide an electrolyte thermal manage-

The minimum voltage required to electrolytically

EQUATION 3

$E=(RT/nF)1n(P₀₂[anode]/P₀₂[catnode])$

wherein:

R is the gas constant $(8.317 \text{ JK}^{-1}\text{mol}^{-1})$

T is the temperature (in degrees K)

n is the number of electrons exchanged (four for each oxygen molecule)

 F is Faraday's constant (96,485 coulombs g^{-1} or $JV^{-1}g^{-1}$ equivalent)

Po2[cathode] is the oxygen partial pressure (in atm.) at the cathode 45

The calculated theoretical open circuit voltage be- 5 tween current collecting electrodes 55 and 60 necessary to drive the reactions of Equations 1 and 2 above is 60 help to distribute current evenly over catalytic cath-
determined to be 0.01 volts. This value is determined ode electrode 45 and catalytic anode electrode 50. determined to be 0.01 volts. This value is determined ode electrode 45 and catalytic anode electrode 50.
according to Equation 3 assuming that breathing air is Moreover, current collectors 55 and 60 provide conveused as the feed gas supplied to inlet pipe 25. It is further 10 nient attachment points for connection that the oxygen partial source 65 . assumed for this determination that the oxygen partial source 65.
pressure at anode 50 ($P_{02[anode]}$) is 1 atmosphere and that In one embodiment of the oxygen concentrator, coolpressure at anode 50 ($P_{02[anode]}$) is 1 atmosphere and that In one embodiment of the oxygen concentrator, cool-
the oxygen partial pressure at cathode 45 ($P_{02[canbed]}$) is ant carrying channels 57 which extend throughout cur the oxygen partial pressure at cathode 45 *(Po_{2[cathode]}*) is 0.21 atmospheres. The operating temperature, T, is

age needed to bias concentrator 10 is approximately two orders of magnitude greater than the above detertwo orders of magnitude greater than the above deter-
mined 0.01 volt theoretical value. This variance is due
carrying channels 57 are formed as an integral part of to the internal resistance of the electrolytic cell formed 20 by concentrator 10 and further due to diffusion and by concentrator 10 and further due to diffusion and When fabricating cathode electrode 45 and anode activation limitations under current flow.

electrode 50, it is desirable that all reactants have ready

the anodic and cathodic overpotentials (concentration, ohmic and activation polarizations) must be properly 25 centrator 10, gaseous reactants are present which must
removed from the cell of concentrator 10 to avoid cell by physically come into contact with the surface of cath overheating. The amount of such excess heat resulting from the exothermic reaction described in Equation 1 is determined to be -330.5 kJ/gmol of oxygen consumed facilitated by the presence of a large area of material
by the cell. Cathode 45 operates at a higher temperature 30 which exhibits very high electrocatalytic activity on by the cell. Cathode 45 operates at a higher temperature 30 which exhibits very high electrocatalytic activity on than anode 50 due to the generation of this excess heat. surfaces of cathode and anode electrodes 45 and 50. than anode 50 due to the generation of this excess heat. surfaces of cathode and anode electrodes 45 and 50.
Cathodic cooling to remove this excess heat from con-
At cathode electrode 45, the electrocatalyst must be Cathodic cooling to remove this excess heat from concentrator 10 is provided by the thermal management centrator 10 is provided by the thermal management selected to be able to efficiently promote oxygen activa-
system of structures 55, 57, 90, 95A, and 100.

electrolyte moisture (humidity) management system anode electrode 50 must permit efficient oxygen regen-
(structures 40, 95B and connective piping therebe- eration (evolution). Once charge transfer is accom-(structures 40, 95B and connective piping therebe-
tween) for preventing dryout of the membrane forming electrolyte 40. In this arrangement, electrolyte 40 is reaction sites through the electronically conducting provided with a continuous supply of water in closed 40 electrode material and adjacent current collector. provided with a continuous supply of water in closed 40 electrode material and adjacent current collector.
loop fashion as seen in FIG. 3 to keep electrolyte 40 The oxygen evolution electrode, anode 50, must oploop fashion as seen in FIG. 3 to keep electrolyte 40 saturated or as saturated as possible under given operating conditions. Since a closed loop formed by electro-
lyte 40, storage tank 95B and interconnective piping is cated from PTFE (polytetrafluoroethylene) bonded employed in concentrator 10, ions from electrolyte 40 45 are recirculated through storage tank 95B and are not inickel screen which exhibits a lower redox potential lost. This moisture management system solves the prob-
than the platinum black anodes which are commonly lem of moisture balance control at the solid polymer employed in fuel cell/electrolyzer applications. In a electrolyte matrix 40 and in the immediate vicinity of preferred embodiment of the invention, the hydrophilic cathode electrode 45 and anode electrode 50. In this 50 anode 50 is intimately bonded to the solid polymer manner, electrolyte 40 is maintained under acceptable electrolyte 40 in the form of a very thin nickel cobalt manner, electrolyte 40 is maintained under acceptable electrolyte 40 in the form of a very thin nickel cobalt operating conditions with respect to its jonic conductiv-
spinel oxide deposit onto which the gold-plated curren ity. Without such a moisture management system, the collector is pressed.

ionic conductivity of solid polymer electrolyte 40 In concentrator 10 of the present invention, it is necionic conductivity of solid polymer electrolyte 40 In concentrator 10 of the present invention, it is nec-
would progressively decrease as concentrator 10 is 55 essary for cathode electrode 45 to reduce oxygen elecwould progressively decrease as concentrator 10 is 55 operated, while the cell resistance of concentrator 10 trochemically in a manner which results in a high activwould simultaneously increase as a result of moisture ity for sole production of hydroxyl ions $(OH⁻)$. It is loss, by entrapment and evaporation, from electrolyte contemplated that many different materials and cathode 40 through porous cathode and anode electrodes, 45 designs may be employed as cathode 45 in concentrator

Current collecting electrodes 55 and 60 are fabricated sis Company, Inc. The hydrophobic cathode electrode from perforated expanded metal screens in one embodi- 65 45 of the present invention is fabricated from 0.3 to 0.4 ment of the invention. One material which was found to mg/cm^2 platinum catalyzed on a carbon cloth support be acceptable to fabricate electrodes 55 and 60 is ex- which is 12 to 14 mils thick and made by Electrosynthe-

Po_{2[anode]} is the oxygen partial pressure (in atm.) at the sheet having a sheet thickness of 10 mils and a strand anode 50 width of 58 mils. Alternatively, perforated titanium and width of 58 mils. Alternatively, perforated titanium and
stainless steel metals may also be employed to fabricate current collectors 55 and 60. The anodic current collector 60 is plated with gold to prevent corrosion and ohmic losses within the cell. Current collectors 55 and 60 help to distribute current evenly over catalytic cathaccording to Equation 3 assuming that breathing air is Moreover, current collectors 55 and 60 provide conve-
used as the feed gas supplied to inlet pipe 25. It is further 10 mient attachment points for connective leads to

rent collector 55 may be fabricated from metallic tubing assumed to be room temperature, 20 degrees Celsius. 15 or other heat conductive sealed pipework so as to pre-
It is noted, however, that in actual practice the volt-
vent the coolant from leaking into expanded metal cur-It is noted, however, that in actual practice the volt-

is not the coolant from leaking into expanded metal cur-

is approximately
 $\frac{1}{2}$ rent collector 55 and adjacent regions of the concentracarrying channels 57 are formed as an integral part of the material used to fabricate collector 55.

tivation limitations under current flow.
The waste heat generated within the cell as a result of access to a large catalytically active interface between access to a large catalytically active interface between electrolyte 40 and current collectors 55 and 60. In conphysically come into contact with the surface of cath-
ode and anode electrodes 45 and 50 so that charge transfer (oxidation and reduction) can occur. This process is facilitated by the presence of a large area of material

tion (reduction) through the reaction described in Equa-As described earlier, concentrator 10 also includes an 35 tion 1 above. Similarly, the electrocatalyst selected for ectrolyte moisture (humidity) management system anode electrode 50 must permit efficient oxygen regenplished, the electrons flow to or from the respective

> erate at relatively low oxygen overvoltage. It has been cated from PTFE (polytetrafluoroethylene) bonded nickel cobalt oxide (NiCO₂O₄) on gold plated 100 mesh than the platinum black anodes which are commonly preferred embodiment of the invention, the hydrophilic spinel oxide deposit onto which the gold-plated current

and 50, to reaction chambers 20A and 20B. 60 10 consistent with the cathode requirements set forth
The gas pressure at cathode 45 (the input pressure at earlier. One particularly effective gas diffusion elecearlier. One particularly effective gas diffusion elecreaction chamber 20A) can be as low as approximately trode that may be employed as cathode 45 is the "Proto-1 atm. with an optimum value of approximately 5 atm. tech Fuel Cell Electrode" marketed by Electrosynthe-Current collecting electrodes 55 and 60 are fabricated sis Company, Inc. The hydrophobic cathode electrode 45 of the present invention is fabricated from 0.3 to 0.4 panded metal from Exmet, Inc. which employs a nickel sis Company, Inc. It should again be understood that contemplated consistent with the guidelines and teachings set forth above.

Based on the above considerations, FIG. 4 illustrates a corrugated configuration to create oxygen and air
a device or system 90 having at least 10 stacked cells 92 5 passages between cells. The oxygen is extracted from a device or system 90 having at least 10 stacked cells 92 *S* passages between cells. The oxygen is extracted from providing at least 95% (by volume) oxygen enriched one side of each cell using manifolds (not shown) at one air, at a flow rate of 0.5 liter per minute for patient use. edge of the stack of cells. Similarly, air is introdu air, at a flow rate of 0.5 liter per minute for patient use. edge of the stack of cells. Similarly, air is introduced to Based on requiring four Faradays of charge, the current the opposite side of the cells from another m Based on requiring four Faradays of charge, the current the opposite side of the cells from another manifold
pull of the device is 131 Amps. To operate an individual coupled to another edge of the cells. While various pull of the device is 131 Amps. To operate an individual coupled to another edge of the cells. While various unit from a 12 VDC car battery, a total of 10 cells, 1.2 10 numbers of cells can be stacked in this manner, it is unit from a 12 VDC car battery, a total of 10 cells, 1.2 10 numbers of cells can be stacked in this manner, it is
VDC each, is required. Assuming a current density of convenient to stack ten galls since the voltage drop approximately 300 A/ft², the electrolytic unit 94 will require a total of approximately 64 in² of memrequire a total of approximately 64 in² of mem-
brane/electrode surface area. For the 10 cell unit, each and brane/electrode surface area. For the 10 cell unit, each age.

cell would be $2\frac{1}{2}'' \times 2\frac{1}{2}''$ in size, and would require 15 While only certain preferred features of the invention

a total of approximately 157 Watts a total of approximately 157 watts of power. From have been shown by way of illustration, many modifica-
these considerations, a conservative estimate of the tions and abanges will occur to those skilled in the art these considerations, a conservative estimate of the tions and changes will occur to those skilled in the art.

actual weight of the device is 22 lbs. This figure in-

It is therefore to be understood that the negati claim actual weight of the device is 22 lbs. This figure in-
cludes the weight of the cooling unit, support structure,
 $\frac{1}{2}$ is therefore, to be understood that the present claims

few atmospheres and 60° -70° C. Waste heat within the What is claimed is:
SPEOC (Solid Polymer Electrolytic Oxygen Concen-
trategy unit can be dissipated in the number of a can be a separated in the number of a capaci SPEOC (Solid Polymer Electrolytic Oxygen Concen-
trator) unit can be dissipated in two ways. In one ap- $\frac{1}{1}$. A process for separating and enriching air with proach, ambient air may be passed continuously 25 $O(X)$ comprising the steps of: through the feed (cathode) side of the cells. This would providing a solid polymer anion conducting material
require annoymetally 71 SCEM (standard cubic feet
rectrolyte including first and second opposed surrequire approximately $7\frac{1}{2}$ SCFM (standard cubic feet electronic material surper minute) of air flow, total, to cool the cells down to
about 70° C. Alternatively, it is possible to recirculate providing a cathode electrode on the first surface of
the cell electrolyte water (at a rate of approximate the cell electrolyte water (at a rate of approximately 50 30 the electrolyte;
ml/min, total) through a simple heat exchanger, at-
tached to the casing It annears that the later approach of the electrolyte; tached to the casing. It appears that the later approach of the electrolyte;
should result in a less costly low weight waste heat connecting a direct current source between the cathshould result in a less costly, low weight waste heat connecting a direct current s
handling system Based on the considerations above it ode and anode electrodes; handling system. Based on the considerations above, it ode and anode electrodes;
is estimated the size and weight of the first generation 35 flowing unprocessed ambient air through the electrois estimated the size and weight of the first generation 35 flowing unprocessed ambient air through the electro-
SPEOC units are less than $2' \times 3' \times 1'$ and 22 lbs. respec. lyte; producing medical grade output oxygen a SPEOC units are less than $2' \times 3' \times 1'$ and 22 lbs., respectively.

The foregoing describes a process and system for ions passing through the electrolyte; and
ectrolytically separating oxygen from air which oper-
bypassing a portion of the air flow around the rim of electrolytically separating oxygen from air which oper-
ates at or near ambient temperatures and pressures. The 40 the solid polymer electrolyte for diluting the outates at or near ambient temperatures and pressures. The 40 the solid polymer electrolyte for diluting the out-
apparatus and method substantially eliminate the need put oxygen to a level therapeutically equivalent to apparatus and method substantially eliminate the need to replenish the electrolyte and requires little mainte-
nance. Moreover, the apparatus and method are rela-
2. The process of claim 1 wherein said steps are pernance. Moreover, the apparatus and method are rela-
tively low cost and are suitable for concentrating oxy-
formed at a temperature within the range of approxitively low cost and are suitable for concentrating oxy-

FIG. 5 illustrates one arrangement of stacked cells for Celsius. use in the oxygen concentrator of FIG. 4. Each cell

other suitable material for fabrication of cathode 45 are comprises an anode platen 101, a cathode plate 102 and contemplated consistent with the guidelines and teach-
a solid polymer membrane 104. The cells are separated gs set forth above.
Based on the above considerations, FIG. 4 illustrates a corrugated configuration to create oxygen and air convenient to stack ten cells since the voltage drop across a cell is preferably 1.2 volts and ten cells provide

are intended to cover all such modifications and
air pump, and control module.
The cell operating pressure and temperature are a
The cell operating pressure and temperature are a
the invention.

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- anode electrode by electrolytic action of hydroxyl ions passing through the electrolyte; and
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gen for therapeutic patient use. 45 mately 20 degrees Celsius to approximately 100 degrees

50

SS

60

65