Carbon molecular sieves modified by the incorporation of finely divided elemental copper useful for the selective sorption of oxygen at elevated temperatures. The carbon molecular sieves can be regenerated by reduction with hydrogen. The copper modified carbon molecular sieves are prepared by pyrolysis of a mixture of a copper-containing material and polyfunctional alcohol to form a sorbent precursor. The sorbent precursors are then heated and reduced to produce copper modified carbon molecular sieves. The copper modified carbon molecular sieves are useful for sorption of all concentrations of oxygen at temperatures up to about 200° C. They are also useful for removal of trace amount of oxygen from gases at temperatures up to about 600° C.

21 Claims, 8 Drawing Sheets
FIG. 1

SAMPLE CMS-CU2
SIZE 34.4 mg

FIG. 4

SAMPLE CMS-CU2
SIZE 34.1 mg

WEIGHT (PERCENT)
FIG. 2

SAMPLE CMS-CU2
SIZE 34.4 mg
ADSORBENT n-Butane

Weight (Percent)

Time (Minutes)

99.56%
98.78%
Figure 3: Graph showing the weight change over time for a sample of CMS-CU2 with a size of 34.4 mg adsorbed with isobutane.
FIG. 5

SAMPLE CMS-CU2
REDUCED AT 200°C BY H₂ + Ar
1.0% O₂ – 99% Ar MIXTURE INTRODUCED
AT 1 MINUTE
OXYGEN UPTAKE AT 200°C

FIG. 6

SAMPLE CMS-CU3
REDUCED AT 200°C BY H₂ + Ar
1.0% O₂ – 99% Ar MIXTURE INTRODUCED
AT 16 MINUTES
OXYGEN UPTAKE AT 200°C
SAMPLE CMS-CU4
REDUCED AT 200°C BY H₂ + Ar
OXYGEN INTRODUCED AT 20 MINUTES
OXYGEN UPTAKE AT 200°C

FIG. 7
SAMPLE CMS-CU2
REDUCED AT 200°C BY H₂ + Ar
0.1% O₂ - 99.9% Ar INTRODUCED
OVER SAMPLE AT 300°C

100.30
100.25
100.20
100.15
100.10
100.05
100.00
99.95
99.90
99.93%
100.19%
O₂
RESIDUE 100.04%

FIG. 8

SAMPLE CMS-CU4
REDUCED AT 200°C BY H₂ + Ar
0.1% O₂ - 99.9% Ar INTRODUCED
OVER SAMPLE AT 300°C

100.2
100.1
100.0
99.9
99.8
99.7
99.6
99.5
99.4
99.3
99.2
99.1
99.0
98.9
98.8
98.7
98.6

FIG. 10
SAMPLE CMS-CU2
REDUCED AT 200°C BY H₂ + Ar
0.1% O₂ – 99.9% Ar INTRODUCED
OVER SAMPLE AT 400°C

FIG. 9
FIG. II

SAMPLE IOMCMS 1
SIZE 39.38 mg
ADSORBENT 1.0% O₂ - 99.0% Ar

TEMPERATURE

WEIGHT

96.64%

97.04%

1.0% O₂ IN ARGON

TIME (MINUTES)
COPPER MODIFIED CARBON MOLECULAR SIEVES FOR SELECTIVE OXYGEN REMOVAL

ORIGIN OF INVENTION

The invention described herein was made in the performance of work under a NASA contract, and is subject to the provisions of Public Law 96-517 (35 USC 202) in which the Contractor has elected to retain title.

BACKGROUND OF THE INVENTION

Carbon molecular sieves are amorphous materials having high surface area and pore sizes near the critical dimensions of small molecules. The sieves are prepared by the pyrolysis of natural and synthetic precursors such as coconut shells, pitch, polyfurfuryl alcohol and polycrylonitrile. Carbon molecular sieves are typically made with pore sizes ranging from 3 to 12 Angstroms, pore volumes from 0.2 to 0.5 cc/g, and surface areas from 300 to 1500 m²/g. Properties of the sieves can be controlled by choice of carbon precursor, the pyrolysis conditions, and choice of additives, the latter of which usually influences the pore structure. The adsorptive properties of carbon molecular sieves will therefore vary widely depending on the above-mentioned factors.

The early applications of carbon molecular sieves were in the area of adsorption and separation of gas mixtures. The separation of gas molecules, which sometimes differ by as little as 0.2 Angstroms in their critical dimensions, have been achieved with carbon molecular sieves. Such separations are possible due to kinetic effects caused by the dependence of the intermolecular force field on the molecules size. For example, in a mixture of nitrogen and oxygen, the oxygen molecules will experience an attractive force field while the nitrogen molecules will experience a repulsive force field. Commercial gas separations take advantage of such kinetic effects by using a pressure swing operating procedure to separate oxygen and nitrogen.

Further selectivity has been achieved with metal-containing carbon molecular sieves that also provide certain catalytically properties. For example, the preparation, adsorptive, and catalytic properties of a variety of metal-containing carbon molecular sieves have been reported in The Proceedings of the Second Conference on Industrial Carbon Graphite, Society of the Chemical Industry, London, 1966, The Journal Chemical Society D, 8, 477, 1970, and The Conference on Industrial Carbon Graphite, Third Meeting Date, Society Chemical Industry, London, 1970.

The preparation of a class of carbon molecular sieves which are functionalized with certain inorganic oxides and supported metals were reported in the American Chemical Society Symposium Series, 368, 335, 1988. These carbon molecular sieves were designed to combine the molecular sieving properties of the carbon with the surface chemical and physical properties of the particular inorganic oxides. In particular the preparation and characterization of carbon molecular sieves modified with titania, zirconia and alumina additives was described and molecular sieving effects were reported with relationship to pore size distribution, and inorganic oxide type and content. A carbon molecular sieve modified by the addition of titania prepared according to the procedure described in the last mentioned reference was made for purposes of comparison and designated as prior art sorbent composition CMS1.

SUMMARY OF THE INVENTION

It is an object of this invention to provide a sorbent having both molecular sieving capability and catalytic reacting capability. It is also an object of this invention to provide a sorbent operable for the selective removal of molecular oxygen from a gas. Another object of this invention is to provide a sorbent which will selectively remove molecular oxygen from a gas containing other gaseous species including species having about the same effective diameter as molecular oxygen.

Yet another object of this invention is to provide a sorbent which will selectively remove molecular oxygen from a gas containing other gaseous species including species having about the same effective diameter as molecular oxygen.

It is also an object of this invention to provide a sorbent which will selectively remove molecular oxygen from a gas containing other gaseous species including species having about the same effective diameter as molecular oxygen without the necessity of using a pressure swing type of sorption-desorption operation as is used in commercially available carbon molecular sieves. Still another object of this invention is to provide a sorbent which will not be deactivated by gaseous species other than molecular oxygen, including but not limited to gaseous sulfur, chlorine and similar gaseous species which frequently will permanently deactivate many sorbents.

It is also an object of this invention to provide a sorbent which will remove trace amounts of molecular oxygen from nearly pure gases thereby producing a gas which is completely free of molecular oxygen.

Another object of this invention is to provide a sorbent which will remove molecular oxygen in all concentrations from a gas at temperatures up to about 200°C.

Still another object of this invention is to provide a sorbent which will remove molecular oxygen in concentrations up to 1000 ppm from a gas at temperatures up to about 400°C.

Yet another object of this invention is to provide a sorbent which will remove molecular oxygen in concentrations up to about 100 ppm from a gas at temperatures up to about 600°C.

Another object of this invention is to provide a sorbent which will lower trace amounts of molecular oxygen from a gas to a level less than about 10 ppb, prefera-
bly less than about 1 ppb, and especially preferably less than about one part per trillion or 1 ppt, at temperatures up to about 600° C.

It is also an object of this invention to provide a sorbent comprising finely divided elemental copper dispersed in carbon molecular sieve.

Another object of this invention is to provide a sorbent which can be regenerated.

It is also an object of this invention to provide a process for making the above mentioned sorbents.

This invention is therefore directed towards carbon molecular sieves modified by incorporation of copper into the carbon molecular sieve matrix. Sorbents of this invention are capable of selectively removing molecular oxygen from a gas mixture at temperatures from about 100° C to about 600° C. In this invention copper in elemental form was successfully incorporated in carbon molecular sieves in such a manner that oxidation of the copper is catalyzed while oxidation of the carbon is retarded, and, while the physical separation properties and sorptive capabilities of the carbon molecular sieves are retained and maintained.

Oxidation of the carbon of the sorbent of this invention was successfully retarded at temperatures up to about 200° C. While the sorbent removed molecular oxygen from a gas which was 100% oxygen. At still higher temperatures, and after a period of time, oxidation of the carbon of the sorbent occurred. The period of time varied depending on the percent oxygen in the treated gas, and the temperature and composition of the particular sorbent. However, usage of the sorbents of this invention is nevertheless useful at such higher elevated temperatures for selectively removing molecular oxygen from gases containing small concentrations or trace amounts of oxygen up to the point of the onset of carbon oxidation.

Accordingly there is provided by the principles of this invention a sorbent comprising a carbon molecular sieve and finely divided elemental copper dispersed in the carbon molecular sieve. The sorbent has both molecular sieving capability and catalytic activity. The sorbent is operable for selectively removing molecular oxygen from a gas containing molecular oxygen. The sorbent removes molecular oxygen by converting the molecular oxygen to copper oxide by catalytic oxidation of the finely divided elemental copper. In one embodiment the sorbent is operable for selectively removing molecular oxygen from a gas containing any concentration of molecular oxygen at temperatures up to about 200° C.

In another embodiment of this invention the sorbent is operable for selectively removing molecular oxygen from a gas containing a concentration of molecular oxygen less than about 1000 ppm at temperatures up to about 400° C. In still another embodiment the sorbent is operable for selectively removing molecular oxygen from a gas containing a concentration of molecular oxygen less than about 100 ppm at temperatures up to about 600° C. In a further embodiment the sorbent is operable for selectively removing molecular oxygen from a gas containing a concentration of molecular oxygen less than about 100 ppt at temperatures up to about 600° C.

In one embodiment of this invention the finely divided elemental copper is from about 1 to about 40% of the weight of the sorbent. In a further embodiment the finely divided elemental copper is from about 2 to about 30% of the weight of the sorbent. In a still further embodiment the finely divided elemental copper is from about 3 to about 25% of the weight of the sorbent.

In one embodiment of this invention at least about 10% of the finely divided elemental copper of the sorbent will selectively react with molecular oxygen in a gas containing molecular oxygen. In a still further embodiment at least about 20% of the finely divided elemental copper of the sorbent will selectively react with the molecular oxygen. In yet another embodiment of this invention at least about 30% of the finely divided elemental copper of the sorbent will selectively react with the molecular oxygen.

The sorbent of this invention, after sorbing molecular oxygen, can be regenerated by heating and reducing the sorbent at an elevated temperature for a predetermined period of time. In one embodiment of this invention the sorbent is regenerated by heating the sorbent to an elevated temperature of at least as high as about 150° C. in a reducing environment for a predetermined period of time. In still another embodiment of this invention the sorbent can be regenerated by heating the sorbent to an elevated temperature of at least as high as about 150° C. in a dilute hydrogen-inert gas mixture for a predetermined period of time. In yet another embodiment of this invention the sorbent is regenerated by heating the sorbent to an elevated temperature of at least as high as about 200° C. in a reducing environment for a predetermined period of time.

There is also provided by the principles of this invention a sorbent containing finely divided copper oxide dispersed in a carbon molecular sieve, the sorbent being formed by pyrolyzing a mixture of a carbon-containing substance and a copper-containing material into a copper oxide containing a carbon molecular sieve and, after pyrolyzing, reducing the copper oxide containing carbon molecular sieve to form a reduced sorbent having finely divided elemental copper dispersed in a carbon molecular sieve matrix. In one embodiment of this invention the copper-containing material is selected from the group consisting of cupric acetate, cupric acetyl acetate and mixtures thereof. In another embodiment of this invention the carbon-containing substance is selected from the group consisting of polyfurfuryl alcohol, polyacrylonitrile, pitch, coconut shells, and mixtures thereof.

There is also provided by the principles of this invention a process for making a sorbent precursor having finely divided copper oxide dispersed in a carbon molecular sieve comprising forming a mixture by dispersing or dissolving a copper-containing material, a carbon-containing substance, and a distributing agent selected from the group consisting of (i) dispersing agents operable for dispersing the copper-containing material in the mixture, (ii) dispersing agents operable for dissolving the copper-containing material, and (iii) mixtures thereof. In this invention the copper-containing substance is transformable into a carbon molecular sieve by pyrolysis, and the copper-containing material is simultaneously transformable into copper oxide by the pyrolysis. The process further comprises pyrolyzing the thusly formed mixture at an elevated temperature in a non-deleteriously reacting environment for a predetermined period of time operable for transforming the mixture into a sorbent precursor having finely divided copper oxide dispersed in a carbon molecular sieve matrix.

In a further embodiment the process also comprises reducing the finely divided copper oxide in the sorbent.
precursor to form a reduced sorbent having finely divided elemental copper dispersed in a molecular sieve. In one embodiment the reduction of the sorbent precursor is by treating the sorbent precursor with molecular hydrogen at an elevated temperature to form the reduced sorbent.

In another embodiment the copper-containing material is cupric acetate, the carbon-containing substance is polyfurfuryl alcohol, and the distributing agent is methanol. In still another embodiment the copper-containing material is cupric acetyl acetonate and the carbon-containing substance and the distributing agent are polyfurfuryl alcohol.

There is also provided by the principles of this invention a process for removing molecular oxygen from gases; and, a process for producing gases which are free of molecular oxygen. Accordingly there is provided a process for reducing the concentration of molecular oxygen in a gaseous mixture comprising contacting the gaseous mixture with a carbon molecular sieve having finely divided elemental copper homogeneously distributed in the carbon molecular sieve at an elevated temperature for a period of time effective for transforming the molecular oxygen into copper oxide by catalytic oxidation of the finely divided elemental copper thereby reducing the concentration of the molecular oxygen in the gaseous mixture; and, after the period of time, recovering the gaseous mixture having a reduced concentration of molecular oxygen. In one embodiment the concentration of molecular oxygen in the gaseous mixture before contacting with the carbon molecular sieve is no greater than about 100 ppm, and the elevated temperature and the period of time of the contacting are effective for reducing the concentration of the molecular oxygen in the gaseous mixture to no greater than about 10 ppb and preferably to no greater than about 1 ppb. In another embodiment the elevated temperature is from about 100° C. to about 600° C. In still another embodiment the process further comprises, prior to the contacting the gaseous mixture with the sorbent, conditioning the carbon molecular sieve at an elevated temperature of at least about 300° C. under a vacuum for a predetermined period of time operable for degassing the sorbent.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is an adsorption curve for CO2 on a first sorbent precursor composition of this invention conducted at 35° C.

FIG. 2 is an adsorption curve for n-butane on the first sorbent precursor composition of this invention conducted at 35° C.

FIG. 3 is an adsorption curve for isobutane on the first sorbent precursor composition of this invention conducted at 35° C.

FIG. 4 is an adsorption curve for 100% oxygen on the first sorbent precursor composition of this invention conducted at 35° C.

FIG. 5 is an oxygen uptake curve for oxygen from a 1% O2/99% Ar gas mixture on the reduced first sorbent composition of this invention conducted at 200° C.

FIG. 6 is an oxygen uptake curve for oxygen from a 1% O2/99% Ar gas mixture on a reduced second sorbent composition of this invention conducted at 200° C.

FIG. 7 is an oxygen uptake curve for oxygen from 100% oxygen on a reduced third sorbent composition of this invention conducted at 200° C.

FIG. 8 is an oxygen uptake curve for oxygen from a 0.1% O2/99.9% Ar gas mixture on the reduced first sorbent composition of this invention conducted at 300° C.

FIG. 9 is an oxygen uptake curve for oxygen from a 0.1% O2/99.9% Ar gas mixture on the reduced first sorbent composition of this invention conducted at 400° C.

FIG. 10 is an oxygen uptake curve for oxygen from a 0.1% O2/99.9% Ar gas mixture on the reduced third sorbent composition of this invention conducted at 300° C.

FIG. 11 is an adsorption curve for a 1.2% O2/98.8% Ar gas mixture on a titania modified carbon molecular sieve, which is not part of this invention, conducted at 200° C.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The preparation of copper-containing carbon molecular sieves of this invention can be carried out by mixing a carbon-containing substance operable for pyrolyzing into a carbon molecular sieve, with effective amount of a copper-containing material operable for simultaneous pyrolyzing into finely divided copper oxide and, pyrolyzing the mixture under inert conditions to produce an oxidized sorbent or sorbent precursor. In one embodiment the carbon-containing substance is polyfurfuryl alcohol which is mixed with an effective amount of a copper-containing compound, and then pyrolyzed to produce a homogeneously mixed copper oxide in a carbon molecular sieve matrix or sorbent precursor. The thusly produced sorbent precursor was then heated in a hydrogen-containing environment to reduce the copper oxide to elemental copper while maintaining the copper homogeneously dispersed in the matrix of the carbon molecular sieve thereby producing a reduced sorbent.

The copper compounds or copper-containing materials added to the polyfurfuryl alcohol were cupric aceta- tate and cupric acetyl acetone. Since cupric acetate did not dissolve in polyfurfuryl alcohol, the cupric acetate was first dissolved in methanol which served as a distributing agent for the cupric acetate. The cupric acetate-methanol solution was then added to the polyfurfuryl alcohol. As the polyfurfuryl alcohol dissolved in the methanol the cupric acetate was seen to precipitate. However, since the precipitate consisted of very small particles which were observed to mix well and become uniformly distributed in the bulk of the mixture, the precipitation did not present a problem. After mixing, the remaining methanol was removed from the mixture by room temperature evaporation. In particular, a solution of 1.0 g cupric acetate in 200 ml methanol was added to 2 ml of polyfurfuryl alcohol. The mixture was stirred and allowed to evaporate at room temperature for 16 hours. The resulting mixture, which was a paste, was stirred and stored for subsequent pyrolysis.

In another embodiment of this invention the addition of cupric acetyl acetone to polyfurfuryl alcohol was performed by physical mixing alone since this copper-containing material appeared to distribute by dissolving in the polyfurfuryl alcohol and produce a homogeneous paste. The resulting paste was stirred and stored for subsequent pyrolysis.

Both of the thusly prepared mixtures or pastes were pyrolyzed to form a carbon molecular sieve having finely divided copper oxide uniformly distributed...
of bent precursors was carried out by using carbon dioxide and the sample was contacted with the first probe gas argon flow, and continuing heating in vacuum to 0.5 hr. The characterization of the sorbent. The vacuum condition was then terminated by flushing in argon to flowing at a rate of about 100 cc/min for a period of one hour. The increase in the sample weight with time was monitored using an automatic data acquisition accessory to the TGA. The characterization of the sorbent precursors was carried out by using carbon dioxide, n-butane and isobutane as probe gases. The reported critical dimensions or characteristic diameters of these probe gases are 3.3, 4.3, and 5.0 Angstroms, respectively. The sorbent precursor samples were preconditioned by flushing in argon to 300° C., terminating the argon flow, and continuing heating in vacuum to 300° C. for one hour, and then cooling under vacuum to 35° C. to ready the samples for subsequent adsorptive study. This procedure is referred to herein as preconditioning the sorbent. The vacuum condition was then terminated and the sample was contacted with the first probe gas flowing at a rate of about 100 cc/min for a period of one hour. The increase in the sample weight with time was monitored using an automatic data acquisition accessory to the TGA. Subsequently the sample was reconditioned by heating under vacuum using the preconditioning procedure described above, and brought in contact with the next probe gas.

For oxygen chemisorptive tests, the reduction of copper oxide modified carbon molecular sieves of this invention was carried out by heating the sample in a flow of 2-5% hydrogen/95-98% argon gas mixture for 12-16 hours at a temperature of 220° C. to form a carbon molecular sieve having finely divided elemental copper uniformly distributed therein or reduced sorbent. The characterization of the copper modified carbon molecular sieves or sorbents of this invention and investigation of their oxygen uptake were conducted using a Du Pont 951 Thermogravimetric Analyzer hereinafter referred to as "TGA". The characterization of the sorbent precursors was carried out by using carbon dioxide, n-butane and isobutane as probe gases. The reported critical dimensions or characteristic diameters of these probe gases are 3.3, 4.3, and 5.0 Angstroms, respectively. The sorbent precursor samples were preconditioned by flushing in argon to 300° C., terminating the argon flow, and continuing heating in vacuum to 300° C. for one hour, and then cooling under vacuum to 35° C. to ready the samples for subsequent adsorptive study.

This procedure is referred to herein as preconditioning the sorbent. The vacuum condition was then terminated and the sample was contacted with the first probe gas flowing at a rate of about 100 cc/min for a period of one hour. The increase in the sample weight with time was monitored using an automatic data acquisition accessory to the TGA. Subsequently the sample was reconditioned by heating under vacuum using the preconditioning procedure described above, and brought in contact with the next probe gas.

For oxygen chemisorptive tests, the reduction of copper oxide modified carbon molecular sieves of this invention was carried out by heating the sample in a flow of 2-5% hydrogen—95-98% argon gas mixture at a temperature of about 200° C. The sample size was typically 30 to 50 mg and the gas mixture flow rate was about 200 cc/min. The total reduction time was about 12-16 hours. At the end of each reduction, hydrogen flow was terminated and the sample brought to the desired test temperature in a flow of argon alone. Flow of 100% oxygen, or an oxygen-argon gas mixture, over the sample was initiated and the change in sample weight with time was monitored.

EXAMPLES

The copper modified carbon molecular sieves or sorbents of this invention that were characterized and investigated are listed in Table 1. Table 1 shows the particular copper additive or copper-containing material used in the preparation of the sorbent and the copper and carbon weight percents in the sorbent. The copper weight percent was obtained by spectroscopic analysis. The carbon percent was obtained by mass balance on the sorbent precursor taking into account that the oxidized sorbent consists of copper oxide as CuO and carbon.

All of the copper modified carbon molecular sieves of this invention exhibited molecular sieving properties in addition to catalytic oxidation properties attributable to the finely divided elemental copper. FIGS. 1 to 3 show adsorption results obtained with a preconditioned sample of the sorbent of this invention having sorbent composition CMS-CU2, at a temperature of 35° C., using carbon dioxide, n-butane, and isobutane, respectively, as the probe gases. FIG. 1 shows that the carbon dioxide adsorption is about 5.25% of the sample sorbent weight. FIG. 2 shows that the n-butane adsorption is about 0.7% of the sample sorbent weight. FIG. 3 shows that the isobutane adsorption is about 0.10% of the sample sorbent weight. It can therefore be concluded that most of the pores of the sorbent of this composition are smaller than 4.3 Angstroms and that very few are larger than 5.0 Angstroms.

After preconditioning at 300° C. under vacuum, the oxygen uptake at 35° C. from a stream of pure oxygen on a sorbent precursor sample, also having composition CMS-CU2, is shown in FIG. 4. From FIG. 4 it is seen that the total oxygen uptake is about 0.3% of the sorbent weight. Since this sample was preconditioned in vacuum the 0.3 wt % increase represents oxygen uptake. The uptake of argon in a similarly conditioned precursor sample at 35° C. was about 0.6% of the sorbent weight. These results are consistent with the somewhat larger molecule size and lower molecular weight of oxygen compared to argon.

At temperatures significantly above room temperature, the physical absorption for most gases on all the sorbents of this invention is expected to be negligible. The chemisorption sites provided by the sorbents after reduction, however, will provide for substantial chemisorption of oxygen and thereby result in the selective removal of oxygen from a gaseous mixture.

The oxygen uptake by reduced sorbents at a temperature of about 200° C. is shown in FIGS. 5 to 7 for three sorbent compositions of this invention, designated as sorbent compositions CMS-CU2, CMS-CU3 and CMS-CU4, respectively. The gas used to contact the reduced sorbents consisted of 1.0% oxygen and 99.0% argon for CMS-CU2 and CMS-CU3, and pure oxygen for CMS-CU4. The maximum oxygen uptake and corresponding copper utilization values are shown in Table 1. The data show that at 200° C. the reduced first and third sorbent compositions, i.e. compositions CMS-CU2 and CMS-CU4, the total oxygen uptake was about 1.8% and 2.0%, respectively, of the sorbent sample weight. The copper utilization, i.e. the percent by weight of the copper oxidized in the sorbent sample, was about 30 to about 34%. The reduced second sorbent composition, CMS-CU3, had an oxygen uptake of about 0.7% of the sorbent sample weight and a copper utilization of about 70%. FIGS. 5 to 7 also show that the rate of oxygen uptake was faster for the first sorbent composition than for the other two compositions.

Oxygen uptake at temperatures of 300° C. and 400° C. was investigated with reduced samples using a 0.1% oxygen and 99.9% argon gas mixture. FIGS. 8 and 9 show the oxygen uptake curve for first sorbent composition at 300° C. and 400° C., respectively. At 300° C. FIG. 8, the reduced sorbent's oxygen uptake was about 0.25% of the sample weight, however, with continued exposure to the oxygen/argon gas stream the sample
weight began to decrease which was attributed to carbon ignition due to hot spots in the sample. A similar result is seen in FIG. 9 for the reduced sorbent sample tested at 400°C wherein initial oxygen uptake occurred for a few minutes followed by a decrease in sorbent weight. FIG. 10 shows the oxygen uptake curve for the reduced third sorbent composition, i.e. sorbent composition CMS-CU4, at 300°C when subjected also to a 0.1% oxygen in argon gas stream. As seen in FIG. 10 there was a very short period of oxygen uptake followed by a continuous decline in sample weight indicating carbon ignition.

The data demonstrates that all of the reduced sorbents of this invention can be used for selective oxygen removal from a gas at temperature up to about 200°C. For any concentration of oxygen in the gas, temperatures of 300°C or higher, after a period of time, carbon ignition occurred when the oxygen concentration in the gas was 0.10% or higher. However, up until ignition occurs, or, for lower concentrations of oxygen in the treated gas, the reduced sorbents removed oxygen even at temperatures of 300°C and 400°C. It is believed that for trace amounts of oxygen, i.e., about 100 ppm, and especially about 20 ppm or lower, the reduced sorbents of this invention will remove oxygen up to temperatures of about 600°C. In particular, since the capabilities of the sorbents of this invention to lower oxygen levels to less than 1 ppm are supported by thermodynamic calculations for the oxidation of copper to CuO and resulting equilibrium values of oxygen partial pressures, the sorbents of this invention are capable of reducing oxygen concentrations in the treated gas to below 1 ppm at temperature up to about 600°C.

The experimental data herein show that copper oxide can be incorporated in carbon molecular sieves, which after reduction, produce copper modified carbon molecular sieves that are effective sorbents for selective removal of oxygen while retaining their molecular sieving properties.

For purpose of comparison to FIGS. 5 to 7, reference is now made to FIG. 11 which shows a sorption test with a known titania modified carbon molecular sieve composition, briefly referred to earlier as prior art sorbent composition CMS1, conducted at 200°C. It resulted in a combined argon and oxygen uptake of about 0.35% on a sorbent sample weight basis. It is to be noted that this sorbent took up both oxygen and argon. As mentioned before, FIGS. 5 to 7 show the oxygen uptake curve for the copper modified carbon molecular sieves of this invention at 200°C. It was about 1.8, 0.7 and 2.0%, respectively, all of which are substantially higher than that achieved with sorbent CMS1.

While the preferred embodiments of the present invention have been described, it should be understood that various changes, adaptations and modifications may be made thereto without departing from the spirit of the invention and the scope of the appended claims. It should be understood, therefore, that the invention is not to be limited to minor details of the illustrated invention shown in preferred embodiment and the figures and that variations in such minor details will be apparent to one skilled in the art.

Therefore it is to be understood that the present disclosure and embodiments of this invention described herein are for purposes of illustration and example and that modifications and improvements may be made thereto without departing from the spirit of the invention or from the scope of the claims. The claims, therefore, are to be accorded a range of equivalents commensurate in scope with the advances made over the art.

### TABLE 1

<table>
<thead>
<tr>
<th>Sorbent</th>
<th>Copper Additive in Precursor</th>
<th>Copper Uptake at 200°C (Wt %)</th>
<th>Percent Copper Utilization</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMS-CU2</td>
<td>Cupric Acetate</td>
<td>1.80</td>
<td>30.0</td>
</tr>
<tr>
<td>CMS-CU3</td>
<td>Cupric Acetylacetate</td>
<td>0.70</td>
<td>73.0</td>
</tr>
<tr>
<td>CMS-CU4</td>
<td>Cupric Acetylacetate</td>
<td>2.00</td>
<td>34.0</td>
</tr>
</tbody>
</table>

What is claimed is:

1. A copper modified carbon molecular sieve-sorbent operable for selectively removing molecular oxygen from a gas containing molecular oxygen, the copper modified carbon molecular sieve having an effective pore size no greater than about 4.3 Angstroms and containing finely divided copper dispersed therein, the copper modified carbon molecular sieve-sorbent formed by pyrolyzing a mixture of a carbon-containing substance selected from the group consisting of polyfur-furyl alcohol, polyacrylonitrile and mixtures thereof, and a copper-containing material selected from the group consisting of cupric acetate, cupric acetylacetate and mixtures thereof, into a copper oxide containing carbon molecular sieve and, after pyrolyzing, reducing the copper oxide therein to form a copper modified carbon molecular sieve-sorbent having an effective pore size no greater than about 4.3 Angstroms and having finely divided elemental copper dispersed therein operable for selectively removing molecular oxygen from a gas containing molecular oxygen.

2. The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent has catalytic activity.

3. The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent is operable for selectively removing molecular oxygen from a gas at temperatures up to about 200°C.

4. The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent is operable for selectively removing molecular oxygen from a gas by converting the molecular oxygen to copper oxide by oxidation of the finely divided elemental copper contained in the copper modified carbon molecular sieve.

5. The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent is operable for selectively removing molecular oxygen from a gas containing a concentration of molecular oxygen less than about 1000 ppm at temperatures up to about 400°C.

6. The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent is operable for selectively removing molecular oxygen from a gas containing a concentration of molecular oxygen less than about 100 ppm at temperatures up to about 600°C.

7. The molecular sieve-sorbent of claim 1, wherein the finely divided elemental copper is from about 1 to about 40% of the weight of the molecular sieve-sorbent.

8. The molecular sieve-sorbent of claim 1, wherein the finely divided elemental copper is from about 2 to about 30% of the weight of the molecular sieve-sorbent.
The molecular sieve-sorbent of claim 1, wherein the finely divided elemental copper is from about 3 to about 25% of the weight of the molecular sieve-sorbent.

The molecular sieve-sorbent of claim 1, wherein at least about 10% of the finely divided elemental copper of the molecular sieve-sorbent will react with the molecular oxygen in a gas containing molecular oxygen.

The molecular sieve-sorbent of claim 1, wherein at least about 20% of the finely divided elemental copper of the molecular sieve-sorbent will react with the molecular oxygen in a gas containing molecular oxygen.

The molecular sieve-sorbent of claim 1, wherein at least about 30% of the finely divided elemental copper of the molecular sieve-sorbent will react with the molecular oxygen in a gas containing molecular oxygen.

The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent after sorbing molecular oxygen, can be regenerated by heating and reducing the molecular sieve-sorbent at an elevated temperature for a predetermined period of time.

The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent after sorbing molecular oxygen, can be regenerated by heating and reducing the molecular sieve-sorbent at an elevated temperature of at least as high as about 150° C. in a reducing environment for a predetermined period of time.

The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent after sorbing molecular oxygen, can be regenerated by heating the molecular sieve-sorbent at an elevated temperature of at least as high as about 200° C. in a reducing environment for a predetermined period of time.

The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent after sorbing molecular oxygen, can be regenerated by heating the molecular sieve-sorbent at an elevated temperature of at least as high as about 150° C. in a dilute hydrogen-inert gas mixture for a predetermined period of time.

The molecular sieve-sorbent of claim 1, wherein the molecular sieve-sorbent after sorbing molecular oxygen, can be regenerated by heating and reducing the molecular sieve-sorbent at an elevated temperature of at least as high as about 200° C. in a dilute hydrogen-inert gas mixture for a predetermined period of time.

The process of claim 17, wherein said reducing of the finely divided copper oxide in the copper modified carbon molecular sieve-sorbent precursor is by treating with molecular hydrogen at an effective elevated temperature.

A process for making a copper modified carbon molecular sieve-sorbent operable for selectively removing molecular oxygen from a gas containing molecular oxygen, the copper modified carbon molecular sieve having an effective pore size no greater than about 4.3 Angstroms and having finely divided copper oxide dispersed therein, the process comprising:

(a) forming a mixture by dispersing or dissolving a copper-containing material in a carbon-containing substance, wherein the copper-containing material comprises cupric acetylacetonate and the carbon-containing substance being simultaneously transformable into copper oxide by pyrolysis, the copper-containing material being simultaneously transformable into copper oxide by pyrolysis;

(b) pyrolyzing the mixture from step (a) at an elevated temperature in a non-deleteriously reacting environment for a predetermined period of time operable for transforming the mixture into a copper modified carbon molecular sieve-sorbent precursor having finely divided copper oxide dispersed therein and having an effective pore size no greater than about 4.3 Angstroms and having finely divided copper oxide dispersed therein.

(b) pyrolyzing the mixture from step (a) at an elevated temperature in a non-deleteriously reacting environment for a predetermined period of time operable for transforming the mixture into a copper modified carbon molecular sieve-sorbent precursor by pyrolysis;
having finely divided copper oxide dispersed therein and having an effective pore size no greater than about 4.3 Angstroms, the pyrolyzing being conducted first at a temperature of about 100° C. for about 1 hour, then at a temperature of about 200° C. for about 0.5 hour, then at a temperature of about 300° C. for about 0.5 hour, then at a temperature of about 400° C. for about 0.5 hour, then at a temperature of about 500° C. for about 0.5 hour, and then at a temperature of about 600° C. for about 3 hours; and,

(c) reducing the finely divided copper oxide in the copper modified carbon molecular sieve-sorbent precursor to form a copper modified carbon molecular sieve-sorbent having finely divided elemental copper dispersed therein.