Provided herein are metal organic frameworks comprising metal nodes and N-donor organic ligands which have high selectivity and stability in the present of gases and vapors including H₂S, H₂O, and CO₂. Methods include capturing one or more of H₂S, H₂O, and CO₂ from fluid compositions, such as natural gas.
Case study $\text{H}_2\text{S} \gg \text{CO}_2 \gg \text{H}_2\text{O}$

Natural gas → MOF → $\text{H}_2\text{S}$ Removal → MOF → $\text{CO}_2$ Removal → MOF → $\text{H}_2\text{O}$ Removal

FIG 1B
Fluorhydric Acid Solution

Ni$^{2+}$ Source

Second Metal Source

Solvent

Mixture

Reacted Mixture

MOF

FIG 2
**FIG 4A**

**FIG 4B**
FIG 5A

FIG 5B
**FIG 5C**

- Initial optimal Activation
- Adsorption at 25°C
- Re-activation at 50°C

- Adsorption at 25°C
- Re-activation at 25°C
- Adsorption at 50°C

- Optimal Re-activation
- Adsorption at 25°C
- Optimal Re-activation

**FIG 6**

Weight (%) vs. Cycle

- Cycle 1
- Cycle 2
- Cycle 3
- Cycle 4
- Cycle 5
- Cycle 6

Weight (%) range from 0 to 30.
FIG 7

FIG 8
FIG 9

FIG 10
FIG 11

FIG 12
* * * * * * * * * * * HAVEN

**FIG 13**

Flow rate : 10 cc/min @ 25°C

- H$_2$S optimal activation
- CO$_2$ optimal activation
- H$_2$S 25°C activation
- CO$_2$ 25°C activation

**FIG 14**

Flow rate : 10 cc/min @ 50°C

- H$_2$S optimal activation
- CO$_2$ optimal activation
- H$_2$S 50°C activation
- CO$_2$ 50°C activation
Flow rate: 10 cc/min @ 25°C

**FIG 15**
METAL ORGANIC FRAMEWORK ABSORBENT PLATFORMS FOR REMOVAL OF CO2 AND H2S FROM NATURAL GAS

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application claims priority to U.S. Provisional Application No. 62/144,175, filed 7 Apr. 2015, which application is incorporated herein by reference.

BACKGROUND

[0002] Today there is an increasing global desire to reduce greenhouse gas emissions and develop clean alternative vehicle fuels. Methane (CH4), the primary component of natural gas (NG), is of particular interest as it is abundant and has lower carbon dioxide (CO2) emission and more efficient combustion than other hydrocarbons due its high H/C ratio. Biogas, including landfill gas, are also seen as promising renewable energy resources, but, like NG, they contain significant amounts of water, CO2, and hydrogen sulfide (H2S) which must be removed before being transported, stored, and burned as a fuel. For example, NG must contain less than 1-2% CO2 and 4 ppm H2S to meet fuel gas specifications for pipeline transportation. Within many industries, gas dehydration and removal of CO2 and H2S remain some of the most intensive and challenging separations, in part due to the intolerance of many technologies to water.

[0003] Available technologies for refining NG and other biogases are often costly, multi-stage processes. Amine scrubbing is a common liquid phase system used to remove acid gases such as CO2 and H2S from NG. However, stagnant historical operating efficiencies, and the excessive oxidative degradation, evaporation, and the corrosive nature of the amine aqueous solutions create a myriad of performance, safety, and environmental concerns. Solid, porous material systems, such as zeolite and metal organic frameworks (MOFs), offer more environmentally friendly alternatives for CO2 capture, but require cumbersome, multi-stage processes. For example, zeolite has single-species selectivity for CO2 and cyclic adsorption performance in the presence of moisture that require prior dehydration and H2S removal stages. MOFs, similarly, can be designed for CO2 capture but exhibit prohibitively low selectivity for acid gases such as H2S. Further, MOFs exhibit low selectivity and capture at low CO2 partial pressures.

[0004] MOFs generally include porous crystals which are assembled from modular molecular building blocks, and provide a wide array of advantageous material properties, including high surface area, porosity, stability, and sorption potential. While the available building block options, and combinations thereof, are virtually limitless, such potential highlights the statistical difficulty in identifying and assembling MOFs with desired and particularized material properties and multi-faceted functionality. For example, many MOFs exhibit high selectivity towards a particular molecular species, but are highly intolerant to water and H2S.

SUMMARY

[0005] In general, this disclosure describes porous metal organic frameworks (MOFs). In particular, this disclosure describes MOFs suitable for the capture and removal of gases and/or vapors from fluids. It should be noted that although the embodiments of this disclosure are described with respect to examples for gas capture, the embodiments described herein are generally applicable to many fields including gas molecule separation, gas storage, catalysis, sensors, drug delivery, rare gas separation, and proton conductivity.

[0006] In one embodiment, a method of capturing chemical species from a fluid composition comprises contacting a first metal organic framework characterized by the formula [M2M2F(OH2)n]m with a fluid composition comprising one or more of carbon dioxide, water, and hydrogen sulfide and capturing one or more of carbon dioxide, water, and hydrogen sulfide from the fluid composition.

[0007] The details of one or more examples are set forth in the accompanying drawings and the description below. Other features, objects, and advantages will be apparent from the description and drawings, and from the claims.

BRIEF DESCRIPTION OF THE DRAWINGS

[0008] The accompanying drawings illustrate non-limiting example embodiments of the invention.

[0009] FIG. 1A illustrates a method for capturing one or more chemical species from a fluid composition via a metal organic framework, according to one or more embodiments of this disclosure.

[0010] FIG. 1B illustrates a multicolour pressure-temperature swing adsorption

[0011] (PTSA) method for capturing one or more chemical species from a fluid composition via a metal organic framework, according to one or more embodiments of this disclosure.

[0012] FIG. 2 illustrates a method of fabricating a metal organic framework, according to one or more embodiments of the disclosure.

[0013] FIG. 3A illustrates a schematic of an inorganic chain, according to one or more embodiments of this disclosure.

[0014] FIG. 3B illustrates a schematic view of a metal organic framework, according to one or more embodiments of this disclosure.

[0015] FIG. 4A illustrates powder X-ray diffraction data of a metal organic framework, according to one or more embodiments of this disclosure.

[0016] FIG. 4B illustrates powder X-ray diffraction data of a metal organic framework, according to one or more embodiments of this disclosure.

[0017] FIG. 5A illustrates isotherm data for various metal organic frameworks, according to one or more embodiments of this disclosure.

[0018] FIG. 5B illustrates isotherm data for various metal organic frameworks, according to one or more embodiments of this disclosure.

[0019] FIG. 5C illustrates isotherm data for a metal organic framework, according to one or more embodiments of this disclosure.

[0020] FIG. 6 illustrates a block flow diagram of sequence of tests for adsorption column breakthrough studies to evaluate the performances of the materials in temperature swing cyclic (TSC) and vacuum swing regeneration (VSR) modes using CO2/CH4/Ar 5/50/90 mixture, according to one or more embodiments of this disclosure.

[0021] FIG. 7 illustrates a graphical view of column breakthrough test of CO2/H2S/CH4 with 10 cm3/min flow
rate on NiAlF₃O(pyrazine)₂ at 25° C. (1 bar) using TSR mode, according to one or more embodiments of this disclosure.

[0022] FIG. 8 illustrates a graphical view of column breakthrough tests of CO₂/H₂S/CH₄ with 10 cm³/min flow rate on NiAlF₃O(pyrazine)₂ at 25° C. and 50° C. (1 bar) respectively (effect of adsorption temperature on retention time), according to one or more embodiments of this disclosure.

[0023] FIG. 9 illustrates a graphical view of column breakthrough tests of CO₂/H₂S/CH₄ with 10 cm³/min flow rate on NiAlF₃O(pyrazine)₂ at 50° C. (1 bar) after activation at different temperatures (effect of adsorption temperature on retention time of different gases), confirming the recyclability of the material, according to one or more embodiments of this disclosure.

[0024] FIG. 10 illustrates a graphical view of comparison of retention time of CO₂ and H₂S for studied materials during column breakthrough tests at 25° C. (1 bar) after optimal activation of the samples (gas mixture CO₂/H₂S/CH₄ (5/5/90) with 10 cm³/min flow rate was used for the experiments), according to one or more embodiments of this disclosure.

[0025] FIG. 11 illustrates a graphical view of comparison of retention time of CO₂ and H₂S for studied materials during column breakthrough tests at 50° C. (1 bar) after optimal activation of the samples (gas mixture CO₂/H₂S/CH₄ (5/5/90) with 10 cm³/min flow rate was used for the experiments), according to one or more embodiments of this disclosure.

[0026] FIG. 12 illustrates a graphical view of comparison of retention time of CO₂ and H₂S for studied materials during column breakthrough tests at 25° C. and 50° C. (1 bar) respectively after optimal activation of the samples (gas mixture CO₂/H₂S/CH₄ (5/5/90) with 10 cm³/min flow rate was used for the experiments), according to one or more embodiments of this disclosure.

[0027] FIG. 13 illustrates a graphical view of comparison of retention time of CO₂ and H₂S for studied materials during column breakthrough tests at 25° C. after optimal activation and 25° C. activation of the samples respectively (gas mixture CO₂/H₂S/CH₄ (5/5/90) with 10 cm³/min flow rate was used for the experiments) and comparison describes compatibility of the studied materials for Vacuum Swing Regeneration (VSR) at 25° C., according to one or more embodiments of this disclosure.

[0028] FIG. 14 illustrates a graphical view of comparison of retention time of CO₂ and H₂S for studied materials during column breakthrough tests at 50° C. after optimal activation and 50° C. activation of the samples respectively (gas mixture CO₂/H₂S/CH₄ (5/5/90) with 10 cm³/min flow rate was used for the experiments) and comparison describes compatibility of the studied materials for Vacuum Swing Regeneration (VSR) at 50° C., according to one or more embodiments of this disclosure.

[0029] FIG. 15 illustrates a graphical view of comparison of retention time of CO₂ and H₂S for studied materials during column breakthrough tests at 25° C. with fresh samples and samples after five breakthrough cycles respectively after optimal activation (gas mixture CO₂/H₂S/CH₄ (5/5/90) with 10 cm³/min flow rate was used for the experiments) and comparison describes recyclability of the studied materials, according to one or more embodiments of this disclosure.

DETAILED DESCRIPTION

[0030] Provided herein are a series of highly stable and highly tunable MOFs with high affinity and stability to water and H₂S. Such qualities allow for efficient and cost effective methods for dehydration gases, vapors, and solvents capable of replacing many cumbersome and expensive industrial processes. Further, this novel series of MOFs can be designed with a variety of pore sizes and assembled with and without open-metal sites, affording tunable properties for a variety of separation applications. For example, the MOF's provided herein can capture chemical species from fluid compositions under conditions and in the presence of chemical species which render known gas capture technologies inefficient, impractical, or inoperable. In particular, the MOF's provided herein can capture one or more of H₂O, CO₂ and H₂S from a fluid composition, such as natural gas, while maintaining high structural integrity during both adsorption and desorption.

[0031] The present invention is described with reference to the attached figures, wherein like reference numerals are used throughout the figures to designate similar or equivalent elements. The figures are not drawn to scale and they are provided merely to illustrate the invention. Several aspects of the invention are described below with reference to example applications for illustration. It should be understood that numerous specific details, relationships, and methods are set forth to provide an understanding of the invention. One skilled in the relevant art, however, will readily recognize that the invention can be practiced without one or more of the specific details or with other methods in other instances, well-known structures or operations are not shown in detail to avoid obscuring the invention. The present invention is not limited by the illustrated ordering of acts or events, as some acts may occur in different orders and/or concurrently with other acts or events. Furthermore, not all illustrated acts or events are required to implement a methodology in accordance with the present invention.

[0032] As used herein, “fluids” can refer to a gas, liquid, or combination thereof. A gas or liquid can include one or more components. For example, a fluid can include a gas stream comprising CO₂, H₂S and water vapor.

[0033] As used herein, “refining” refers to removing one or more unwanted components or separating one or more components from remaining components of a composition, such as a fluid. For example, refining can include removing a fraction of H₂S from a fluid, such as natural gas.

[0034] As used herein, “poly-functional” refers to the characteristic of having more than one reactive or binding sites. For example, a poly-functional ligand can attach to a metal ion in multiple ways, bridge multiple metal ions, or combinations thereof. Specifically, pyrazine is a poly-functional ligand.

[0035] Gas storage and separation using porous materials has experienced significant development in recent years in various industrial applications related to energy, environment, and medicine. Among porous materials, metal organic frameworks (MOFs) are a versatile and promising class of crystalline solid state materials which allow porosity and functionality to be tailored towards various applications. MOF crystal chemistry uses a molecular building block (MBB) approach that offers potential to construct MOFs where desired structural and geometrical information are incorporated into the building blocks prior to the assembly process.
Generally, MOFs comprise a network of nodes and ligands, wherein a node has a connectivity capability at three or more functional sites, and a ligand has a connectivity capability at two functional sites each of which connect to a node. Nodes are typically metal ions or metal containing clusters, and, in some instances, ligands with node connectivity capability at three or more functional sites can also be characterized as nodes. In some instances, ligands can include two functional sites capable of each connecting to a node, and one or more additional functional sites which do not connect to nodes within a particular framework. A MBB can comprise a metal-based node and an organic ligand which extraplate to form a coordination network. Such coordination networks have advantageous crystalline and porous characteristics affecting structural integrity and interaction with foreign species (e.g., gases). The particular combination of nodes and ligands within a framework will dictate the framework topology and functionality. While essentially limitless combinations of nodes and ligands exist, to date, very few MOF materials are H₂S stable which consequently preclude their use in gas separation.

As disclosed in co-owned U.S. Application No.: 62/044,928, a series of isoreticular MOFs with periodically arrayed hexafluorosilicate (SiF₆²⁻) pillars, called SIFSIX-2-Cu-i and SIFSIX-3-Zn, SIFSIX-3-Cu and SIFSIX-3-Ni showed particularly high CO₂ selectivity and capture. These properties in SIFSIX-3-M materials suggest broad applications from ppm level CO₂ removal to bulk CO₂ separation. However, with the exception of SIFSIX-3-Ni, the SIFSIX-3-M materials were not tolerant to H₂S. And although these materials exhibit high structural stabilities in the presence of CO₂, extensive exposure of all SIFSIX-3-M materials to moisture detrimentally induces a phase change and the formation of new 2D stable materials. These 2D materials exhibit relatively unaltered selectivity but diminished CO₂ uptake. This indicates that the SIFSIX-3-M materials series is not sufficiently robust to handle CO₂ and H₂S capture in most critical applications throughout the oil and gas and renewable fuels industries, especially in applications which bring the materials into contact with moisture.

Provided herein are novel functionalized MOFs suitable for the removal of acid gases, particularly CO₂ and H₂S, which additionally exhibit high water vapor tolerance and stability over thousands of cycles. These MOFs eliminate the safety, efficiency, and environmental concerns associated with amine scrubbing techniques while providing high selectivity toward acid gases and negligible uptake of other components, including CH₄. The proposed class of materials will permit NG and biofuels refining approaches based on simultaneous removal of CO₂, H₂S and water vapor. The benefits of this innovative approach include the following: (i) no environmental and safety hazards germane to amine scrubbing (ii) no preliminary separate desulfurization is necessary (iii) no separate gas drying is needed, and (iv) no compression-decompression/cooling of NG is required. Further, the MOFs provided herein offer exceptional thermal and mechanical stability, particularly during adsorption/desorption.

MOFs as provided herein comprise one or more MBBs. Generally, a MBB, or a network of MBBs, can be represented by the formula [(node)ₙ(ligand)ₙ(solvent)ₙ], wherein n represents the number of molecular building blocks. Solvent represents a guest molecule occupying pores within the MOF, for example as a result of MOF synthesis, and can be evaporated after synthesis to provide a MOF with unoccupied pores. In one example, an evacuated MOF can be subsequently enriched with a guest molecule compatible with the MOF framework and/or pores for a particular purpose (e.g., to outfit the MOF for use as a sensor). In other embodiments, guest molecules can include adsorbed gases, such as H₂S. While guest molecules can impart functionality onto a MOF, such are not a permanent fixture of the MOF. Accordingly, the value of c can vary down to zero, without changing the definitional framework of the MOF. Therefore in many instances, MOFs as provided herein will be defined as [(node)ₙ(ligand)ₙ], without reference to a solvent or guest molecule component.

In some embodiments herein, MOFs can be characterized by the formula [(node)ₙ(ligand)ₙ(solvent)ₙ], A non-limiting list of solvents can include one or more of H₂O, DMF, and DEF. In some embodiments, solvent can include a chemical species present after fabrication of the MOF. In some embodiments, solvent can include a functionalizing guest molecule, such as water, DMF, and DEF. Some embodiments herein comprise a porous, uninhabited MOF characterized by the formula [(node)ₙ(ligand)ₙ], wherein node comprises, generally, MₓMₓFₓOₓ(H₂O)ₓ. In some embodiments, Mₓ comprises elements selected from periodic groups IB, IIA, IIIA, IVA, IB, VB, VIIIB, or VIII. In some embodiments, Mₓ comprises elements selected from periodic groups IIIA, IIIB, VB, VIB, or VIIIB, and Mₓ comprises elements selected from periodic groups IIIA, IIIB, VB, VIB, VIIIB, or VIII. In some embodiments, Mₓ can comprise one of the following cations: Cu²⁺, Zn²⁺, Co²⁺, Ni²⁺, Mn²⁺, Zr²⁺, Fe²⁺, Ca²⁺, Ba²⁺, Pb²⁺, Pd²⁺, Ru²⁺, Rh⁺, Cd²⁺, Mg²⁺, Al³⁺, Fe³⁺, Cr³⁺, Zr³⁺, Ru³⁺, and Co³⁺. In some embodiments, Mₓ can be one of the following Al³⁺, Fe³⁺, Cr³⁺, Ti⁴⁺, V⁵⁺, V⁶⁺, Sc³⁺, In³⁺, Nb⁵⁺, Y⁵⁺. In some embodiments, Mₓ can comprise one of the following cations: Cu²⁺, Zn²⁺, Co²⁺, Ni²⁺, Mn²⁺, Zr²⁺, Fe²⁺, Ca²⁺, Ba²⁺, Pb²⁺, Pd²⁺, Ru²⁺, Rh⁺, Cd²⁺, Mg²⁺, Al³⁺, Fe³⁺, Cr³⁺, Ti⁴⁺, V⁵⁺, V⁶⁺, Sc³⁺, In³⁺, Nb⁵⁺, Y⁵⁺. In some embodiments, the ligand can be any bi-functional N-donor linkers based on monocyclic or poly cyclic group (aromatic or not).

In some embodiments, a ligand can comprise a polydentate, or polyfunctional ligand, such as a bi-functional ligand, a tri-functional ligand, or ligands with four or more functional sites. In some embodiments, a ligand can comprise an N-donor linker. In some embodiments a ligand can comprise a poly-functional ligand. In some embodiments, a ligand can comprise a plurality of N-donor functional groups. In some embodiments, a ligand can comprise a monocyclic or poly cyclic group structure, wherein the cyclic groups can be aromatic or non-aromatic. In some embodiments, a ligand can comprise a nitrogen-containing monocyclic or poly cyclic group structure. In some embodiments, a ligand can comprise a nitrogen-containing heterocyclic ligand, including pyridine, pyrazine, pyrimidine, pyridazine, triazine, thiazole, oxazole, pyrrole, imidazole, pyrazole, triazole, oxadiazole, thiaziazole, quinoline, benzoxazole, benzimidazole, and thiazole.
cations: Cu$^{2+}$, Zn$^{2+}$, Co$^{2+}$, Ni$^{2+}$, Mn$^{2+}$, Zr$^{2+}$, Fe$^{3+}$, Cr$^{3+}$, Ba$^{2+}$, Pb$^{2+}$, Pt$^{2+}$, Ru$^{2+}$, Rh$^{2+}$, Cd$^{2+}$, Mg$^{2+}$, Al$^{3+}$, Fe$^{2+}$, Fe$^{3+}$, Cr$^{3+}$, Cr$^{4+}$, Ru$^{3+}$, Ru$^{4+}$, and Co$^{2+}$. $M_2$ can be one of the following: Al$^{3+}$, Fe$^{2+}$, Fe$^{3+}$, Cr$^{3+}$, Ti$^{2+}$, V$^{3+}$, V$^{5+}$, Sc$^{3+}$, In$^{3+}$, Nb$^{5+}$, Y$^{3+}$, and the ligand can be any bi-functional N-donor linkers based on monocyclic or polycyclic group (aromatic or not).

[0043] The utility of MOFs such as those provided herein are highly dependent upon the framework’s structural features such as structural strength, density, functionality, pore aperture dimensions, pore dimensions, the ratio of pore aperture dimensions to pore dimensions, pore accessibility, and the presence of a plurality of pore dimensions and/or pore aperture dimensions (e.g., a poly-porous MOF). The originality of this new class of crystalline porous materials is based, in part, on the fact that the shape of cavities, (i.e. square or rectangle based channels), is controlled from a structural point of view using appropriate cations and organic linkers. The novel MOF architectures disclosed herein offer a novel improvement on some MOF architectures by replacing silicon components with other metals, such as Al$^{3+}$, Fe$^{2+}$, Fe$^{3+}$, V$^{3+}$, V$^{4+}$, V$^{5+}$, Nb$^{5+}$, to afford highly stable materials with or without open metal sites. In some embodiments, the use of specific cations, such as Al$^{3+}$, Fe$^{2+}$, Fe$^{3+}$, Cr$^{3+}$, Cr$^{4+}$, Ti$^{3+}$, Ti$^{4+}$, V$^{3+}$, V$^{5+}$, Sc$^{3+}$, In$^{3+}$, V$^{5+}$, in $M_2$ site positions can introduce open-metal sites within the channels that enhance properties of gas capture.

[0044] These, and other features, collaborate to achieve MOFs with high affinity and stability to water and H$_2$S. Additionally, the novel series of MOFs structures disclosed herein can be designed with a variety of pore sizes and/or open-metal sites which afford tunable properties for a variety of gas/vapor/solvent separation applications. Tuning, in some embodiments, can include modification of the organic and/or inorganic components of the MOF. For example, lighter metal-based clusters can be used to lower the framework density and increase the relative wt. % of captured CO$_2$ and/or H$_2$S. Further, the MOF platforms as provided herein allow for an unprecedented high degree of tuning control at the molecular level, allowing the size and shape of channels within a MOF architecture to be rigorously controlled and adapted to specific separation of numerous gases, beyond CO$_2$ and H$_2$S.

[0045] In some embodiments, a representative $M_nM_{2}F_{x}O_{y}$(H$_2$O)$_{z}$([Ligand])$_{x}$ MOF structure can include a Ni$_x$M$_y$ constituent, an M$_y$ constituent group selected from one of Al, Fe, V, or Nb, and a Ligand comprising a pyrazine constituent group, wherein x can vary from 0 to 10. All such embodiments offer high affinity and stability to water vapor and H$_2$S, unlike the Cu and Zn-based analogues of SiF$_3$SiX$_3$-M materials made with Si. In some embodiments a MOF characterized by the formula $M_nM_{2}F_{x}O_{y}$(H$_2$O)$_{z}$([Ligand])$_{x}$ wherein M$_x$ equals Ni, M$_y$ equals Al, Fe, V or Nb, and ligand equals pyrazine, the pore size (channel size) of the resulting MOF can be about 3.3 Å to about 3.8 Å. In some embodiments, the channels are square/rectangular. In the same or in an alternative embodiment, a MOF can have a specific surface area of about 250 m$^{2}$/g to about 500 m$^{2}$/g. In either of the same MOFs or in an alternative embodiment, a MOF can have a pore volume of about 0.1 cm$^{3}$/g to about 0.25 cm$^{3}$/g. In a different embodiment, a more elongated ligand can provide an analogous MOF with much higher porosity.

[0046] FIG. 1A illustrates a method 100 for capturing 120 one or more chemical species from a fluid composition 110 via a first MOF 105. A method 100 for capturing 120 one or more chemical species from a fluid composition 110 can comprise contacting 115 a metal organic framework 105 characterized by the formula $[M_nM_{2}F_{x}O_y$(H$_2$O)$_{z}$([Ligand])$_{x}$ with a fluid composition 110. Fluid composition 310 can comprise two or more chemical species. Fluid composition 310 can comprise natural gas. In one embodiment, fluid composition 110 comprises one or more of carbon dioxide, water, and hydrogen sulfide. In such an embodiment, fluid composition 110 can further comprise methane.

[0047] Method 100 can further comprise capturing 120 one or more chemical species from the fluid composition 110. Capturing 120 can comprise capturing one or more of carbon dioxide, water, and hydrogen sulfide. Capturing 120 can comprise capturing two or more of carbon dioxide, water, and hydrogen sulfide. In some embodiments, capturing 120 comprises physical adsorption of the one or more captured chemical species by the first MOF 105. In some embodiments, capturing 120 comprises chemisorption of the one or more captured chemical species by the first MOF 105. Chemisorption can occur by one or more captured chemical species chemically interacting with one or more open metal sites of the first MOF 105. In some embodiments, capturing 120 comprises physical adsorption and chemisorption of the one or more captured chemical species by the metal organic framework. Capturing 120 can comprise wholly or partially containing a chemical species within a pore of a MOF. In some embodiments, capturing 120 consists of chemisorption. In some embodiments, capturing 120 consists of physical adsorption. Capturing 120 can occur in a capturing environment. A capturing environment can comprise one or more of ambient temperature or pressure. A capturing environment can comprise a pressurize-controlled vessel. A capturing environment can comprise a temperature-controlled vessel. A capturing environment can comprise a pressurize-controlled and temperature-controlled vessel. A capturing environment can comprise the first MOF 105 in the form of a fixed bed, a packed column, or combinations thereof.

[0048] Capturing 120 can further comprise changing the temperature of the capture environment to alter the affinity of one or more chemical species for the MOF. Similarly, capturing 120 can further comprise changing the pressure of the capture environment to alter the affinity of one or more chemical species for the MOF. Additionally, capturing 120 can further comprise changing the temperature and pressure of the capture environment to alter the affinity of one or more chemical species for the MOF. In such embodiments, the chemical species can particularly include carbon dioxide, water, and hydrogen sulfide.

[0049] Method 100 can further comprise desorbing one or more of carbon dioxide, water, and hydrogen sulfide from the MOF. Method 100 can further comprise desorbing two or more of carbon dioxide, water, and hydrogen sulfide from the MOF. Desorbing can occur in a desorbing environment. A desorbing environment can comprise one or more of ambient temperature or pressure. A desorbing environment can comprise a temperature-controlled and pressure-controlled vessel. A desorbing environment can comprise a temperature-controlled and pressure-controlled vessel. While a MOF can capably capture a number of gaseous or vapor phase species, a
relative affinity hierarchy will, in principle, exist among those species. In embodiments where method 100 comprises desorbing two or more of carbon dioxide, water, and hydrogen sulfide from the MOF, carbon dioxide, water, and hydrogen sulfide can be sequentially desorbed in descending order of affinity. In a particular embodiment, H\textsubscript{2}S is desorbed first, CO\textsubscript{2} is desorbed second, and H\textsubscript{2}O is desorbed last.

[0050] In some embodiments, the relative affinity hierarchy of chemical species can be manipulated. Accordingly, desorbing can further comprise changing the temperature of the desorbing environment to alter the affinity of one or more chemical species for the MOF. Similarly, desorbing can further comprise changing the pressure of the desorbing environment to alter the affinity of one or more chemical species for the MOF. Additionally, desorbing can further comprise changing the temperature and pressure of the desorbing environment to alter the affinity of one or more chemical species for the MOF. In such embodiments, the chemical species can particularly include carbon dioxide, water, and hydrogen sulfide. In some embodiments, one or more of capturing and desorbing is effected by multilocation pressure-temperature swing adsorption. Multilocation pressure-temperature swing adsorption is known in the art.

[0051] Method 100 can further comprise contacting 115 the fluid composition 110 with a second MOF characterized by the formula \( [\text{M}_x \text{M}_y \text{F}_{z}(\text{OH},\text{O})_n,\text(Ligand)}(\text{solvent})_m ] \). In some embodiments, the second MOF is different from the first MOF 105. In some embodiments, contacting 115 the fluid composition with the first MOF 105 and the second MOF occurs separately. In some embodiments, contacting 115 the fluid composition with the first MOF 105 and the second MOF occurs simultaneously. Method 100 can further comprise contacting 115 the fluid composition 110 with three or more MOFs.

[0052] In some embodiments, the first MOF comprises one or more MOFs as provided herein. For example, the first MOF can comprise an MOF characterized by the formula \( [\text{NiNbF}_{x}(\text{OH},\text{O})_n,\text(Ligand)}(\text{solvent})_m ] \). In some embodiments, the second MOF comprises one or more MOFs as provided herein. For example, the second MOF can comprise an MOF characterized by the formula \( [\text{NiF}_{x}(\text{OH},\text{O})_n,\text{(Ligand)}(\text{solvent})_m ] \), wherein \( M \) comprises Al\textsuperscript{3+}, Fe\textsuperscript{3+}, or Fe\textsuperscript{2+}. In some embodiments, the MOF is characterized by the formula \( [\text{NiNbF}_{x}(\text{OH},\text{O})_n,\text{(Ligand)}(\text{solvent})_m ] \), wherein \( M \) comprises Al\textsuperscript{3+}, Fe\textsuperscript{3+}, or Fe\textsuperscript{2+}. FIG. 1B illustrates a method 101 for removing one or more of carbon dioxide, water, and hydrogen sulfide from natural gas via a MOF. Removing can one or more of carbon dioxide, water, and hydrogen sulfide from natural gas can occur simultaneously. Removing can one or more of carbon dioxide, water, and hydrogen sulfide from natural gas can occur separately. Removing can occur in a multilocation pressure-temperature swing adsorption (PTSA) system. In one embodiment, the MOF comprises one of Ni(Al/Fe/V/Nb)\textsubscript{2}(OH,\text{O})\textsubscript{3}(pyrazine)(solvent). In another embodiment, the MOF comprises one or more of Ni(Al/Fe/V/Nb)\textsubscript{2}(OH,\text{O})\textsubscript{3}(pyrazine)(solvent). MOFs as provided herein can be fabricated using a solvo(hydro)thermal synthetic procedure. As shown in FIG. 2, a method for fabricating 200 a MOF 230 can include combining 205 reactants. Reactants can include one or more of a fluorohydric acid solution 206 with a Ni\textsuperscript{2+} source 207, a second metal source 208, and a solvent 209 to form a mixture 210. A Ni\textsuperscript{2+} source 207 can include one or more of nickel nitrate, hydrated nickel nitrate, nickel chloride, hydrated nickel chloride, nickel fluoride, hydrated nickel chloride, nickel oxide, or hydrated nickel oxide. The second metal source 208 can include an Al\textsuperscript{3+} source, an Fe\textsuperscript{2+} source, an Fe\textsuperscript{3+} source, a Cr\textsuperscript{3+} source, a Cr\textsuperscript{2+} source, a Ti\textsuperscript{4+} source, a V\textsuperscript{3+} source, a V\textsuperscript{5+} source, a Sc\textsuperscript{2+} source, an In\textsuperscript{3+} source, a Nb\textsuperscript{5+} source, or a Y\textsuperscript{3+} source, for example. These metals can be in the form of nitrates, hydrated nitrates, chlorides, hydrated chlorides, fluorides, hydrated fluorides, oxides, hydrated oxides, and combinations thereof. The solvent 209 can include one or more of H\textsubscript{2}O, dimethylformamide (DMF), and diethylformamide (DEF).

[0055] The method for fabricating 200 can further comprise reacting 215 the mixture 210 until sufficient to form a reacted mixture 220. Reacting 215 can include contacting the fluoroacidic solution 206, the Ni\textsuperscript{2+} source 207, the second metal source 208, and the solvent 209. Reacting 215 can further comprise stirring or agitating the mixture 210, or heating the mixture 210. Heating the mixture 210 can comprise heating to a temperature between about 80°C to about 200°C. The reacted mixture 220 can be further processed 225 to provide a fabricated MOF 230. Processing 220 can include one or more of filtering the reacted mixture 220, rinsing the reacted mixture 220 with water, removing excess reactants from the reacted mixture 220. In some embodiments, guest molecules are optionally evacuated from a fabricated MOF 230. Guest molecules can include solvent guest molecules, or derivatives thereof.

[0056] One MOF synthesis strategy provided herein comprises linking inorganic chains using appropriate N-donor based linkers to deliberately generate channels along one crystallographic direction. The inorganic chains are built up from the trans-connection between \( M_2N_2F_6 \) and \( M_2F_4 \) \((\text{H}_2\text{O})_2 \) octahedra or between \( M_2N_2F_6 \) and \( M_2F_4 \) \((\text{H}_2\text{O})_2 \) octahedra or between \( M_2N_2F_6 \) octahedra and \( M_2F_4 \) \((\text{OH},\text{O}) \) octahedra. FIG. 3A illustrates an example of an inorganic chain, built up from \( M_2N_2F_6 \) and \( M_2F_4 \) \((\text{H}_2\text{O})_2 \) octahedra. The resulted inorganic chains are linked to each other using bi-functional N-donor organic ligands, thereby generating channels with different sizes and shapes depending on the nature of the organic linker. FIG. 3B illustrates a schematic view of one embodiment of a MOF comprising a NiNbF\textsubscript{6}O\textsubscript{6} (pyrazine), structure, viewed along the c-axis.

[0057] In one embodiment, a representative \( M_2N_2F_6 \) \((\text{Ligand})_2 \) MOF structure can include a Ni M\textsubscript{6} constituent, a Nb M\textsubscript{6} constituent group, and a Ligand comprising a pyrazine constituent group. FIG. 4A illustrates powder X-ray diffraction data of this MOF, characterized by the formula \( \text{NiNbF}_6\text{O}(\text{pyrazine})_2(\text{solvent})_m \), confirming the high stability of the MOF in the presence of water. FIG. 4B illustrates powder X-ray diffraction data of this MOF, confirming the high stability of the MOF in the presence of H\textsubscript{2}S.

[0058] FIGS. 5A-C illustrates the broad potential for MOFs characterized by the formula \( \text{NiAlFeV/NbF}_6\text{O}(\text{pyrazine})_2(\text{solvent})_m \), and variants thereof. FIG. 5A illustrates CO\textsubscript{2} isotherms at 298 K for each of these four MOFs, which show extremely high sorption of CO\textsubscript{2} across all test pressures (0-1 bar). FIG. 5B illustrates H\textsubscript{2}S isotherms at 298K for three MOFs characterized by the formula
Ni(Al/Fe/V)F₂(OH/H₂O)₅(pyrazine)₂(solvent)₂, which shows across all test pressures (0–1, wherein 1 corresponds to 0.1 H₂S partial pressure).

[0059] FIG. 5C illustrates H₂O sorption isotherms at 298K for Ni(Al/Fe/V/Nb)F₂(OH/H₂O)₅(pyrazine)₂(solvent)₂, indicating the high suitability of this MOF for gas dehydrogenation applications, among others. These results demonstrate the value of the MOF embodiment characterized by the formula Ni(Al/Fe/V/Nb)F₂(OH/H₂O)₅(pyrazine)₂(solvent)₂ as a platform for refining a number of valuable hydrocarbon gases and fluids, including methane, natural gas, and biogas. The exemplary performance and properties of the Nb²⁺ based MOFs disclosed herein are notably achievable in spite of having no open metal sites. These and other results can be expected in similar other embodiments, with or without open metal sites, such as MOF structure characterized by the formula Ni₅M₃F₆O₁₆(pyrazine)₂, wherein M₃ can be one of the following: Al³⁺, Fe²⁺, Fe³⁺, Cr²⁺, Cr³⁺, Ti³⁺, V³⁺, V⁵⁺, Sc³⁺, In³⁺, Nb⁵⁺, Y³⁺. These and other results can be expected in similar other embodiments, with or without open metal sites, such as MOF structure characterized by the formula M₃Ni₅F₆O₁₆(pyrazine)₂, wherein M₃ can be one of the following cations: Cu²⁺, Zn²⁺, Mn²⁺, Ni²⁺, Mn²⁺, Zr²⁺, Fe²⁺, Ca²⁺, Ba²⁺, Pb²⁺, Pt²⁺, Pd²⁺, Ru²⁺, Rh²⁺, Cd²⁺, Mg²⁺, Al³⁺, Fe²⁺, Fe³⁺, Cr²⁺, Cr³⁺, Ru²⁺, Ru³⁺. These and other results can also be expected in similar other embodiments, with or without open metal sites, such as MOF structure characterized by the formula M₅Ni₅F₆O₁₆(pyrazine)₂, wherein M₅ can be one of the following cations: Cu²⁺, Zn²⁺, Mn²⁺, Ni²⁺, Mn²⁺, Zr²⁺, Fe²⁺, Ca²⁺, Ba²⁺, Pb²⁺, Pt²⁺, Pd²⁺, Ru²⁺, Rh²⁺, Cd²⁺, Mg²⁺, Al³⁺, Fe²⁺, Fe³⁺, Cr²⁺, Cr³⁺, Ru²⁺, Ru³⁺.

EXAMPLE 1

[0060] In order to study the performance of NiAlF₃O(pyrazine)₂ for adsorption of CO₂ and H₂S, a series of cyclabes testing were carried out in a flow mode system using the sequence of tests described in FIG. 6.

[0061] NiAlF₃O(pyrazine)₂, a material from a related SiF₆ platform which is known to be very selective to CO₂ was tested. Interestingly, the gas adsorption test with CO₂, H₂/CH₄ 5/5/90 at 25°C. In TSR mode showed a retention time in the column of 27 min/g for H₂S and 31 min/g for CO₂, respectively (FIG. 7), indicative of similar affinity of the NiAlF₃O(pyrazine)₂ framework to both CO₂ and H₂S. These results showed that H₂S and CO₂ could be removed simultaneously with the same affinity.

[0062] Most importantly, heating the adsorption column in a second TSR (temperature swing regeneration mode) cycle to 50°C for NiAlF₃O(pyrazine)₂ did not significantly affect the retention time (25 vs. 27 min/g) for both H₂S and CO₂, respectively (FIG. 8).

[0063] FIG. 9 shows that TSR mode could be practiced for NiAlF₃O(pyrazine)₂ using reactivation at 50°C and even VSR mode at 25°C with a minimum loss in the H₂S retention time while keeping the same affinity for H₂S and CO₂. FIG. 9 shows that for NiAlF₃O(pyrazine)₂, the H₂S and CO₂ uptakes were reduced by 20% between TSR and VSR (vacuum swing regeneration) mode when adsorption is carried at 50°C.

[0064] Finally, the reproducibly test (6th cycle) after 5 adsorption-desorption tests showed that NiAlF₃O(pyrazine)₂ exhibit a reproducible H₂S and CO₂ adsorption performance. In this case, NiAlF₃O(pyrazine)₂ can be utilized as a potential material for the simultaneous removal of H₂S and CO₂.

EXAMPLE 2

[0065] With the exception of NiAlF₃O(pyrazine)₂, it was shown that other MOF materials studied exhibit higher selectivity and higher retention time for H₂S than CO₂ at 25°C. (FIG. 10). Y-14,NDC-fcu-MOF is the best material from the studied materials for H₂S removal with retention time of 70 min/g(5% H₂S, 10 cm³/min flow rate). Retention time for H₂S decreases for almost 40% when temperature is increased to 50°C. (FIGS. 11, 12), with highest retention time of 41 min/g (5% H₂S, 10 cc/min flow rate) for Y-14,NDC-fcu-MOF. Retention time of H₂S and CO₂ for Y-14,NDC-fcu-MOF and Y-fum-fcu-MOF does not change during VSR at 25°C. (FIG. 13), while retention time for all the samples does not change during VSR at 50°C. (FIG. 14).

[0066] Unlike all other studied materials here, NiAlF₃O(pyrazine)₂, has the same retention time of 28 min/g (5% H₂S, 5% CO₂, 10 cm³/min flow rate) for both H₂S and CO₂ at 25°C. (FIG. 10). By increasing temperature to 50°C, the retention time for CO₂ and H₂S only decrease slightly to 26 min/g (5% H₂S, 5% CO₂, 10 cm³/min flow rate) unlike other materials (FIGS. 11, 12). During VSR at 25°C for NiAlF₃O(pyrazine)₂, the retention time for H₂S decreases slightly (FIG. 13), however the retention time for H₂S and CO₂ remains unchanged during VSR at 50°C. (FIG. 14). NiAlF₃O(pyrazine)₂, is a good candidate for simultaneous removal of H₂S and CO₂ over the wide temperature range, while other studied materials can be good candidates for removal of H₂S first and then CO₂ in the presence of CH₄ containing gas streams at 25°C.

[0067] As shown in FIG. 15, uptake for all the materials remains practically unchanged between 1st cycle and 6th cycle under the same condition showing that all these materials selected for mixed gas testing are very stable.

[0068] The CO₂ and H₂S uptakes for all the MOFs studied in the course of this project, using TSR and VSR mode, are reported in Tables 1 and 2.

<table>
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<tr>
<th></th>
<th>CO₂ uptake</th>
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<tbody>
<tr>
<td></td>
<td>wt %</td>
<td>wt %</td>
</tr>
<tr>
<td><strong>TABLE 1</strong></td>
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</tr>
<tr>
<td></td>
<td>298K</td>
<td>323K</td>
</tr>
<tr>
<td></td>
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<td>323K</td>
</tr>
<tr>
<td><strong>Materials</strong></td>
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<tr>
<td>Ga₆-MOF</td>
<td>0.58</td>
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<td>Y-FTZB-fcu-MOF</td>
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<td>0.78</td>
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<td>Y-14,NDC-fcu-MOF</td>
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<td>Y-fum-fcu-MOF</td>
<td>0.78</td>
<td>0.40</td>
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<td>NiAlF₃O(pyrazine)₂(solvent)</td>
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<td>2.55</td>
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<td>NiSiF₄ DPA i</td>
<td>1.57</td>
<td>1.32</td>
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*CH₄ uptake not detected by the flow mode system and CO₂/CH₄, H₂/CH₄ selectivity is calculated assuming 0.1 mg as the lowest detectable uptake.
TABLE 2

<table>
<thead>
<tr>
<th>Materials</th>
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<td>Y-fum-fcu-MOF</td>
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<td>0.59</td>
<td>3.87</td>
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<td>2.28</td>
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TABLE 3

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<td>1800</td>
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<tr>
<td>NiSiF6-DPA</td>
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<td>3342</td>
<td>1928</td>
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</table>

TABLE 4

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<td>2185</td>
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<td>NiAlF5(H2O)(pyrazine)(solvent)</td>
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<td>1671</td>
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<tr>
<td>NiSiF6-DPA</td>
<td>1028</td>
<td>867</td>
<td>3342</td>
<td>1928</td>
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</tbody>
</table>

*CH4 uptake not detected by the flow mode system and CO2/CH4, H2S/CH4 selectivities are calculated assuming the uptake of CH4 is equal to 0.1 mg, which is the lowest detectable uptake by the setup.

[0069] Because if low uptake of CH4 vs H2S and CO2 during the mixed gas tests, it is impossible to detect and quantify the CH4 uptake for the other studied materials. Accordingly, in order to determine the CO2/CH4 and CO2/CH4 H2S selectivity (Tables 3 and 4), it was assumed that the non-detectable uptake of CH4 is equal to the accuracy of the setup (0.1 mg).

[0070] Overall, all the materials selected exhibit very high selectivity for H2S vs CH4. As expected, the NiAlF5(O)(pyrazine) exhibited the highest selectivity for CO2. It is noted that adsorption selectivity values higher than 700-1000 are often attached with elevated degree of inaccuracy, hence, this values should be considered as infinite. Accordingly, the high values for CO2/CH4 and H2S/CH4 selectivities reported in tables 3 and 4 are highly qualitative.

[0071] In summary, highly stable recyclable MOF materials for H2S and CO2 removal from natural gas with a wide range of H2S/CO2 selectivity are described that can be used for TSR and VSR at room and higher temperatures. Highly selective materials are shown for CO2 having open metal sites (such as NiAlF5(H2O)(pyrazine)(solvent)) that have the potential to remove CO2 and H2S simultaneously with high selectivity.

1. A method of capturing chemical species from a fluid composition, the method comprising contacting a first metal organic framework characterized by the formula [MnF6(H2O)(Ligand)(solvent)] with a fluid composition comprising one or more of carbon dioxide, water, and hydrogen sulfide; and capturing one or more of carbon dioxide, water, and hydrogen sulfide from the fluid composition.

2. The method of claim 1, wherein M comprises elements selected from periodic groups IB, II, IIIA, IIIA, IVA, IVB, VIB, VIIA, and VIIIB, and M6 comprises elements selected from periodic groups IIIA, IIIA, IVB, VIB, VIIA, and VIIIB. Ligand comprises an organic, poly-functional, N-donor ligand.

3. The method of claim 1, wherein M comprises Cu2+, Zn2+, CO2-, Ni2+, Mn2+, Zr4+, Fe2+, Ca2+, Ba2+, Pb2+, Pt4+, Ru4+, Rh2+, Cd2+, Mg2+, Al3+, Fe3+, Fe3+, Cr3+, Cr2+, Ru4+, Rh4+, or Co3+, and M6 comprises Al3+, Fe3+, Fe3+, Cr3+, Cr2+, Ti3+, V3+, V3+, Sc3+, In3+, N5+, or O3+.

4. The method of claim 1, wherein M comprises Ni2+ and M6 comprises Al3+, Fe3+, Fe3+, V3+, V3+, or Nb5+.

5. The method of claim 1, wherein the ligand comprises pyridine, pyrazine, pyrimidine, pyridazine, triazine, thiazole, oxazole, pyrrole, imidazole, pyrazole, triazole, oxadiazole, thiadiazole, quinoline, benzoazole, or benzimidazole.

6. The method of claim 1, wherein capturing comprises physical absorption of the one or more captured chemical species by the metal organic framework.

7. The method of claim 1, wherein capturing comprises chemisorption of the one or more captured chemical species by the metal organic framework.

8. The method of claim 7, wherein chemisorption occurs by one or more captured chemical species chemically interacting with one or more open metal sites of the metal organic framework.

9. The method of claim 1, wherein the fluid composition further comprises methane.

10. The method of claim 1, further comprising capturing two or more of carbon dioxide, water, and hydrogen sulfide from the fluid composition.

11. The method of claim 10, further comprising desorbing two or more of carbon dioxide, water, and hydrogen sulfide.

12. The method of claim 11, wherein two or more of carbon dioxide, water, and hydrogen sulfide are sequentially desorbed in descending order of affinity.

13. The method of claim 1, further comprising changing the pressure of the capture environment to alter the affinity of one or more of carbon dioxide, water, and hydrogen sulfide for the metal organic framework adsorbent.

14. The method of claim 1, further comprising changing the temperature of the capture environment to alter the
affinity of one or more of carbon dioxide, water, and hydrogen sulfide for the metal organic framework adsorbent.

15. The method of claim 14 further comprising changing the temperature of the capture environment to alter the affinity of one or more of carbon dioxide, water, and hydrogen sulfide for the metal organic framework adsorbent.

16. The method of claim 1, simultaneously or separately contacting the fluid composition with a second metal organic framework, characterized by the formula \[ [\text{M}_\text{c}\text{M}_\text{d} \text{F}_{\text{c} \text{d}}(\text{O}/\text{H}_2\text{O})_{\text{c}}(\text{Ligand})_{\text{c}}(\text{solvent})_{\text{c}}]_{\text{c}} \], wherein the second metal organic framework is different from the first metal organic framework.

17. The method of claim 16, wherein the first metal organic framework is characterized by the formula \[ [\text{NiNbF}_{\text{c} \text{d}}(\text{O}/\text{H}_2\text{O})_{\text{c}}(\text{Ligand})_{\text{c}}(\text{solvent})_{\text{c}}]_{\text{c}} \], and the second metal organic framework is characterized by the formula \[ [\text{M}_\text{c}\text{M}_\text{d} \text{F}_{\text{c} \text{d}}(\text{O}/\text{H}_2\text{O})_{\text{c}}(\text{Ligand})_{\text{c}}(\text{solvent})_{\text{c}}]_{\text{c}} \], wherein \( \text{M}_\text{c} \) comprises Ni+ and \( \text{M}_\text{d} \) comprises Al\(^{3+}\), Fe\(^{2+}\), or Fe\(^{3+}\).

18. The method of claim 1, wherein the metal organic framework adsorbent is in the form of a fixed bed, a packed column, or combinations thereof.

19. The method of claim 1, further comprising desorbing one or more of carbon dioxide, water, and hydrogen sulfide.

20. The method of claim 19, wherein one or more of capturing and desorbing is effected by multicolonm pressure-temperature swing adsorption.