Title of the Invention: **Synthesis of AEI zeolite**

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A composition comprises a synthetic zeolite having an AEI framework structure and an in-situ transition metal dispersed within the cavities and channels of the zeolite. The invention also comprises a method for synthesizing a zeolite comprising: preparing a reaction mixture comprising (a) at least one source of alumina, (b) at least one source of silica, (c) a transition-metal-amine organic templating agent, (d) an alkali metal hydroxide, and (e) distinct second organic templating agent, wherein each of the first and second templating agents is suitable for forming an AEI framework structure and wherein the reaction mixture is essentially free of fluorine; heating the reaction mixture at crystallization conditions for a sufficient time to form zeolite crystals having an AEI framework and optionally an ITE framework and containing the transition metal. The transition metal is preferably copper or iron, and the zeolite can be incorporated into a catalyst for treating exhaust gas containing NOx and/or NH3.
SYNTHESIS OF AEI ZEOLITE

The present invention relates to transition metal zeolite having an AEI framework. The invention also relates to a method for preparing such zeolites and their use as a catalyst.

Zeolites are crystalline or quasi-crystalline aluminosilicates constructed of repeating SiO₄ and AlO₄ tetrahedral units. These units are linked together to form frameworks having regular intra-crystalline cavities and channels of molecular dimensions. Numerous types of synthetic zeolites have been synthesized and each has a unique framework based on the specific arrangement of its tetrahedral units. By convention, each framework type is assigned a unique three-letter code (e.g., “AEI”) by the International Zeolite Association (IZA).

Synthetic zeolites are typically produced using a structure directing agent (SDA), also referred to as a “template” or “templating agent”. SDAs are typically complex organic molecules which guide or direct the molecular shape and pattern of the zeolite’s framework. Generally, the SDA serves to position hydrated silica and alumina and/or as a mould around which the zeolite crystals form. After the crystals are formed, the SDA is removed from the interior structure of the crystals, leaving a molecularly porous aluminosilicate cage.

Zeolites have numerous industrial applications including internal combustion engines, gas turbines, coal-fired power plants, and the like. In one example, nitrogen oxides (NOₓ) in the exhaust gas may be controlled through a so-called selective catalytic reduction (SCR) process whereby NOₓ compounds in the exhaust gas are contacted with a reducing agent in the presence of a zeolite catalyst.

ZSM-5 and Beta zeolites have been studied as SCR catalysts due to their relatively wide temperature activity window. However, the relatively large pore structures of these zeolites have a number of drawbacks. First, they are susceptible to high temperature hydrothermal degradation resulting in a loss of activity. Also, large and medium pore sizes tend to adsorb hydrocarbons which are oxidized as the temperature of the catalyst increases, thus generating a significant exotherm which can thermally damage the catalyst. This problem is particularly acute in lean-burn systems, such as vehicular diesel engines, where significant quantities of hydrocarbon can be
adsorbed during cold-start. Coking by hydrocarbons presents another significant drawback of these relatively large and medium pore molecular sieve catalysts. In contrast, small pore molecular sieve materials, such as those having an AEI framework, offer an improvement in that fewer hydrocarbons are able to permeate into the framework.

To promote the catalytic reaction, transition metals may be included in the zeolite material, either as a substituted framework metal (commonly referred to as “metal-substituted zeolite”) or as a post-synthesis ion exchanged or impregnated metal (commonly referred to as “metal-exchanged zeolite”). As used herein, the term "post-synthesis" means subsequent to zeolite crystallization. The typical process for incorporating a transition metal into a zeolite is by cationic exchange or impregnation of metals or precursors after the molecular sieve is formed. However, these exchange and impregnation processes for incorporating metals frequently lead to poor uniformity of metal distribution, particularly when exchanged into small pore molecular sieve.

Applicants have developed a unique family of transition metal-containing zeolites which are referred to herein as “JMZ-2 zeolite” or “JNZ-2”. This zeolite material contains AEI frameworks and optionally ITE frameworks either as a mixed phase material and/or as an intergrowth, and a non-framework transition metal. Preferably, the transition metal, such as copper, is an ionic species located in the cavities and channels of the zeolite crystal. The transition metal is incorporated into the zeolite material as the zeolite crystals are formed.

According to certain aspects of the invention, JMZ-2 can be prepared via a one-pot synthesis mixture by incorporating a transition metal-amine complex which serves as a first SDA and a second, distinct SDA. As used herein, the terms “first” and “second” with respect to SDA are used to clarify that the two SDAs are distinct compounds, but the terms do not suggest or represent the order or sequent of operation or addition to the synthesis reaction admixture. The combination of two SDAs into a single reaction mixture is referred to herein as a mixed-template and the incorporation of a transition metal into the zeolite during crystallization is referred to as one-pot synthesis. Preferably, Cu-versions of JMZ-2 are synthesized using Cu-tetraethylenepentamine (Cu-TEPA) as the first SDA and N,N-Dimethyl-3,5-dimethylpiperidinium or 1,1-Diethyl-2,6-dimethylpiperidinium as the second SDA.
In a certain embodiment of the invention, provided is a catalyst composition comprising a synthetic zeolite having an AEI framework and optionally an ITE framework and an in-situ transition metal uniformly dispersed within the cavities and channels of the zeolite.

In another embodiment of the invention, provided is a catalyst composition comprising a synthetic zeolite having an AEI framework and optionally an ITE framework, containing about 0.1 to about 7 weight percent of non-framework copper based on the total weight of the zeolite, and containing less than 5 weight percent CuO based on the total weight of the zeolite.

In another embodiments of the invention, provided is a method for synthesizing a zeolite comprising the steps of (1) preparing a reaction mixture comprising (a) at least one source of alumina, (b) at least one source of silica, (c) a transition-metal-amine organic templating agent, (d) seed crystals, and (e) distinct second organic templating agent, wherein each of the first and second templating agents is suitable for forming AEI framework structures; and (2) heating the reaction mixture at crystallization conditions for a sufficient time to form zeolite crystals having an AEI framework and containing the transition metal. In certain embodiments these steps are performed sequentially as described.

In another embodiment of the invention, provided is a catalyst article for treating exhaust gas comprising a catalyst composition described herein, wherein the catalyst composition is disposed on and/or within a honeycomb monolith substrate.

And in yet another embodiment of the invention, provided is a method for treating an exhaust gas comprising contacting a combustion exhaust gas containing NOx and/or NH3 with a catalyst article described herein to selectively reduce at least a portion of the NOx into N2 and H2O and/or oxidize at least a portion of the NH3.

In general, JMZ-2 zeolites are prepared from a one-pot synthesis mixture containing a source of silica, a source of alumina, a first framework organic templating agent in the form of a transition metal-amine, and a second organic templating agent. The transition metal-amine is used to incorporate an ionic species of the transition metal, such as copper, into the channels and/or cavities of the zeolite during crystallization. The non-framework transition metal incorporated into the zeolite during its synthesis is referred to herein as in-situ metal. In certain embodiments, the silica, alumina, templating agents, and seed crystals are mixed to form a reaction mixture, for example a gel, which is then heated to facilitate crystallization. The metal-
containing zeolite crystals precipitate out of the reaction mixture. These crystals are collected, washed, and dried.

As used herein, the term "AEI" refers to an AEI type framework and the term "ITE" refers to an ITE type framework as recognized by the International Zeolite Association (IZA) Structure Commission. The novel synthesis method described herein is capable of producing a zeolite material having about 1 – 99 wt. % aluminosilicates having an AEI framework and about 99 – 1 wt. % aluminosilicates having an ITE framework, provided that the zeolite material comprises at least 70 wt. %, preferably at least about 80 wt. %, at least about 90 wt. %, at least about 95 wt. %, or at least about 99 wt. %, of a combination of AEI and ITE frameworks, based on the total weight of aluminosilicate in the zeolite material. Other secondary phases, such as FAU and/or MOR, may also be present. Preferably, these secondary phases comprise less than about 10 weight percent of the zeolite material, more preferably less than about 5 weight percent, and even more preferably less than about 2 weight percent.

In certain the zeolite material comprises a majority of AEI framework relative to ITE framework. For example, the JMZ-2 zeolite can contain aluminosilicates having an AEI framework and aluminosilicates having an ITE framework in an AEI:ITE ratio of about 1.05:1, about 1.5:1, about 2:1, about 3:1, about 4:1, about 5:1, about 10:1, about 20:1, or about 100:1.

In other embodiments, the zeolite material comprises a majority of ITE framework relative to AEI framework. For example, the JMZ-2 zeolite can contain aluminosilicates having an ITE framework and aluminosilicates having an AEI framework in an ITE:AEI ratio of about 1.05:1, about 1.5:1, about 2:1, about 3:1, about 4:1, about 5:1, about 10:1, about 20:1, or about 100:1.

As used herein the term "zeolite" means a synthetic aluminosilicate molecular sieve having a framework constructed of alumina and silica (i.e., repeating SiO_{4} and AlO_{4} tetrahedral units), and preferably having a molar silica-to-alumina ratio (SAR) of at least 10, for example about 20 to about 50.

The zeolites of the present invention are not silica-aluminophosphates (SAPOs) and thus do not have an appreciable amount of phosphorous in their framework. That is, the zeolite frameworks do not have phosphorous as a regular repeating unit and/or do not have an amount of phosphorous that would affect the basic physical and/or chemical properties of the material, particularly with respect to the material’s capacity to selectively reduce NO_{x} over a broad
temperature range. In certain embodiments, the amount of framework phosphorous is less than 0.1 weight percent, preferably less than 0.01 or less than 0.001 weight percent, based on the total weight of the zeolite.

Zeolites, as used herein, are free or substantially free of framework metals, other than aluminium. Thus, a "zeolite" is distinct from a "metal-substituted zeolite" (also referred to as "isomorphous substituted zeolite"), wherein the latter comprises a framework that contains one or more non-aluminium metals substituted into the zeolite's framework.

Suitable silica sources include, without limitation, fumed silica, silicates, precipitated silica, colloidal silica, silica gels, dealuminated zeolites such as dealuminated zeolite Y, and silicon hydroxides and alkoxides. Silica sources resulting in a high relative yield are preferred. Typical alumina sources also are generally known and include aluminates, alumina, other zeolites, aluminium colloids, boehmites, pseudo-boehmites, aluminium hydroxides, aluminium salts such as aluminium sulphate and alumina chloride, aluminium hydroxides and alkoxides, alumina gels.

As a first SDA, a transition metal-amine complex is utilized. Suitable transition metals include those which are known for use in promoting SCR of NO\textsubscript{x} compounds in exhaust gases, with Cu and Fe being preferred, and Cu being particularly preferred. Suitable amine components for the metal-amine complex include organic amines and polyamines which are capable of directing AEI framework formation. A preferred amine component is tetraethylenepentamine (TEPA). The metal-amine complex (i.e., Cu-TEPA) may be pre-formed or formed in-situ in the synthesis mixture from individual metal and amine components.

A second framework templating agent, other than the above-noted copper-amine complex, is selected for directing AEI synthesis. Examples of suitable second templating agents include N,N-Diethyl-2,6-dimethylpiperidinium cation; N,N-Dimethyl-9-azoniabicyclo 3.3.1nonane; N,N-Dimethyl-2,6-dimethylpiperidinium cation; N-Ethyl-N-methyl-2,6-dimethylpiperidinium cation; N,N-Diethyl-2-ethylpiperidinium cation; N,N-Dimethyl-2-(2-hydroxyethyl)piperidinium cation; N,N-Dimethyl-2-ethylpiperidinium cation; N,N-Dimethyl-3,5-dimethylpiperidinium cation; N-Ethyl-N-methyl-2-ethylpiperidinium cation; 2,6-Dimethyl-1-Azoniabicyclo 5.4 decane cation; N-Ethyl-N-propyl-2,6-dimethylpiperidinium cation; 2,2,4,6,6-Pentamethyl-2-azoniabicyclo 3.2.1 octane cation; and N,N-Diethyl-2,5-dimethyl-2,5-dihydropyrrolo cation, with N,N-Dimethyl-3,5-dimethylpiperidinium or 1,1-Diethyl-2,6-dimethylpiperidinium being particularly preferred. The anion associated with the
cation may be any anion which is not detrimental to the formation of the zeolite. Representative anions include halogen, e.g., fluoride, chloride, bromide and iodide, hydroxide, acetate, sulphate, tetrafluoroborate, carboxylate, and the like. Hydroxide is the most preferred anion. In certain embodiments, the reaction mixture and subsequent zeolite is free or essentially free of fluorine.

One-pot synthesis is conducted by combining predetermined relative amounts of the silica source, aluminium source, transition metal-amine complex, the second organic templating agent, and optionally a source of hydroxide ions such as NaOH, and seed crystals, such as AEI zeolite, under various mixing and heating regimens as will be readily apparent to those skilled in the art. JMZ-2 can be prepared from a reaction mixture having the composition shown in Table 1 (shown as weight ratios). The reaction mixture can be in the form of a solution, gel, or paste, with a gel being preferred. Silicon- and aluminium-containing reactants are expressed as SiO₂ and Al₂O₃, respectively.

<table>
<thead>
<tr>
<th></th>
<th>Typical</th>
<th>Preferred</th>
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</thead>
<tbody>
<tr>
<td>SiO₂ / Al₂O₃</td>
<td>10 – 100</td>
<td>15 – 65</td>
</tr>
<tr>
<td>OH-SiO₂</td>
<td>0.5 – 2</td>
<td>0.7 – 1.5</td>
</tr>
<tr>
<td>Template 1 / Template 2</td>
<td>0.01 – 0.3</td>
<td>0.05 – 0.15</td>
</tr>
<tr>
<td>Templates / SiO₂</td>
<td>0.05 – 0.5</td>
<td>0.10 – 0.25</td>
</tr>
<tr>
<td>Transition metal / Template 1</td>
<td>0.02 – 5</td>
<td>0.1 – 2</td>
</tr>
<tr>
<td>H₂O / SiO₂</td>
<td>20 – 80</td>
<td>30 – 45</td>
</tr>
</tbody>
</table>

Reaction temperatures, mixing times and speeds, and other process parameters that are suitable for conventional AEI synthesis techniques are also generally suitable for the present invention. Without limitation, the following synthesis steps may be followed to synthesize JMZ-2. An aluminium source (e.g., dealuminated zeolite Y) is mixed in water with sodium hydroxide to promote dissolution of alumina. The mixture is combined with an organic templating agent (e.g., 1,1-Diethyl-2,6-dimethylpiperidinium) and mixed by stirring or agitation for several minutes (e.g., about 5 – 30). A silica source (e.g., TEOS) is added and mixed for several minutes (e.g., about 30 – 120 minutes) until a homogeneous mixture is formed. Then, a copper source (e.g. copper
sulphate) and TEPA are added to the mixture and mixed by stirring or agitation for several minutes (e.g., about 15 – 60 minutes). Hydrothermal crystallization is usually conducted under autogenous pressure, at a temperature of about 100 to 200 °C for a duration of several days, such about 1 – 20 days, a preferably about 1 – 3 days.

At the conclusion the crystallization period, the resulting solids are separated from the remaining reaction liquid by standard mechanical separation techniques, such as vacuum filtration. The recovered solids are then rinsed with deionized water, and dried at an elevated temperature (e.g., 75 – 150 °C) for several hours (e.g., about 4 to 24 hours). The drying step can be performed under vacuum or at atmospheric pressure.

The dried JMZ-2 crystals are preferably calcined, but can also be used without calcination. It will be appreciated that the foregoing sequence of steps, as well as each of the above-mentioned periods of time and temperature values are merely exemplary and may be varied.

In certain embodiments, a source of alkali metal, such as sodium, is not added to the synthesis mixture. The phrase "essentially alkali-free" or "alkali-free" as it is used herein means that alkali metal is not added to the synthesis mixture as an intentional ingredient. An "essentially alkali-free" or "alkali-free" catalyst as referred to herein means generally that the catalyst material contains an inconsequential level of alkali metal with regard to the intended catalytic activity. In certain embodiments, the JMZ-2 zeolite contains less than about 0.1 weight percent, and preferably less than about 0.01 weight percent, alkali metal such as sodium or potassium.

Applicants also discovered that the foregoing one-pot synthesis procedure permits adjusting the transition metal content of the crystals based on the composition of the starting synthesis mixture. For example, a desired Cu or Fe content can be directed by providing a predetermined relative amount of Cu or Fe source in the synthesis mixture, without requiring post synthesis impregnation or exchange to increase or decrease the copper loading on the material.

In certain embodiments, the synthesized zeolite contains about 0.1 to about 5 weight percent copper, iron, or combination thereof, for example about 0.5 wt.% to about 5 wt.%, from about 1 to about 3 wt.%, from about 0.5 to about 1.5 wt.%, and about 3.5 wt.% to about 5 wt.%. For example, a controlled Cu loading of 0.5-5% by weight, 0.1-1.5% by weight or 2.5-3.5% by weight, for example, can be achieved without additional post-synthesis processing. In certain embodiments, the zeolite is free of post-synthesis exchanged metal, including copper and iron.
The transition metal is catalytically active and substantially uniformly dispersed within the AEI and optionally ITE frameworks. Here, a substantially uniformly dispersed transition metal means that the zeolite substance contains not more than about 5 weight percent transition metal in the form of a transition metal oxide (e.g., CuO, FeO, Fe₂O₃, Fe₃O₄), also referred to herein as a free transition metal oxide, or a soluble transition metal oxide, relative to the total amount of that transition metal in the JMZ-2 zeolite. For example, the JMZ-2 zeolite contains not more than about 5 weight percent, not more than about 3 weight percent, not more than about 1 weight percent, and not more than about 0.1 weight percent, from example about 0.01 to about 5 weight percent, about 0.01 to about 1 weight percent, or about 0.01 to 3 weight percent CuO based on the total weight of copper in the zeolite material. Applicants have found that minimizing the concentration of CuO improves the hydrothermal durability and exhaust gas treatment performance of the JMZ-2 zeolite.

Preferably, the JMZ-2 zeolite contains a majority of in-situ transition metal compared to free transition metal oxides. In certain embodiments, the JMZ-2 zeolite contains a weight ratio of free transition metal oxides (e.g., CuO) to in-situ transition metal (e.g. ionic Cu) of less than about 1, less than about 0.5, less than about 0.1, or less than about 0.01, for example about 1 to about 0.001, about 0.5 to about 0.001, about 0.1 to about 0.001, or about 0.01 to about 0.001.

Preferably, the JMZ-2 zeolite does not contain framework transition metals in an appreciable amount. Instead, the copper or iron is present as an ionic species within the interior channels and cavities of the zeolite framework. Accordingly, the metal-containing JMZ-2 zeolite is not a metal-substituted zeolite (e.g., a zeolite having a metal substituted into its framework structure) and not necessarily a metal-exchanged zeolite (e.g., a zeolite that underwent a post synthesis ion exchange). In certain embodiments, the JMZ-2 zeolite is free or essentially free of metals other than copper and aluminium or is essentially free of metals other than iron and aluminium. For example, in certain embodiments, the JMZ-2 zeolite is free or essentially free of nickel, zinc, tin, tungsten, molybdenum, cobalt, bismuth, titanium, zirconium, antimony, manganese, magnesium, chromium, vanadium, niobium, ruthenium, rhodium, palladium, gold, silver, indium, platinum, iridium, and/or rhenium. In certain embodiments, the JMZ-2 zeolite is free or essentially free of iron. In certain embodiments, the JMZ-2 zeolite is free or essentially free of calcium. In certain embodiments, the JMZ-2 zeolite is free or essentially free of cerium.
The JMZ-2 zeolite is useful as a catalyst in certain applications. The JMZ-2 catalyst can be used without a post-synthesis metal exchange. However, in certain embodiments, the JMZ-2 can undergo a post-synthesis metal exchange. Thus, in certain embodiments, provided is a catalyst comprising a JMZ-2 zeolite containing one or more catalytic metals exchanged into the channels and/or cavities of the zeolite post zeolite-synthesis in addition to in-situ copper or in-situ iron. Examples of metals that can be post-zeolite synthesis exchanged or impregnated include transition metals, including copper, nickel, zinc, iron, tungsten, molybdenum, cobalt, titanium, zirconium, manganese, chromium, vanadium, niobium, as well as tin, bismuth, and antimony; noble metals including platinum group metals (PGMs), such as ruthenium, rhodium, palladium, indium, platinum, and precious metals such as gold and silver; alkaline earth metals such as beryllium, magnesium, calcium, strontium, and barium; and rare earth metals such as lanthanum, cerium, praseodymium, neodymium, europium, terbium, erbium, ytterbium, and yttrium. Preferred transition metals for post-synthesis exchange are base metals, and preferred base metals include those selected from the group consisting of manganese, iron, cobalt, nickel, and mixtures thereof.

Metals incorporated post-synthesis can be added to the molecular sieve via any known technique such as ion exchange, impregnation, isomorphous substitution, etc. The amount of metal post-synthesis exchanged can be from about 0.1 to about 3 weight percent, for example about 0.1 to about 1 weight percent, based on the total weight of the zeolite.

In certain embodiments, the metal-containing zeolite contains post-synthesis exchanged alkaline earth metal, particularly calcium and/or magnesium, disposed within the channels and/or cavities of the zeolite framework. Thus, the metal-containing zeolite of the present invention can have transition metals ($T_M$), such as copper or iron, incorporated into the zeolite channels and/or cavities during synthesis and have one or more exchanged alkaline earth metals ($A_M$), such as calcium or potassium, incorporated post-synthesis. The alkaline earth metal can be present in an amount relative to the transition metal that is present. For example, in certain embodiments, $T_M$ and $A_M$ are present, respectively, in a molar ratio of about 15:1 to about 1:1, for example about 10:1 to about 2:1, about 10:1 to about 3:1, or about 6:1 to about 4:1, particularly were $T_M$ is copper and $A_M$ is calcium. In certain embodiments, the relative cumulative amount of transition metal ($T_M$) and alkali and/or alkaline earth metal ($A_M$) is present in the zeolite material in an amount relative to the amount of aluminium in the zeolite, namely the framework aluminium. As used herein, the
(T_{M} + A_{M}):Al ratio is based on the relative molar amounts of T_{M} + A_{M} to molar framework Al in the corresponding zeolite. In certain embodiments, the catalyst material has a (T_{M} + A_{M}):Al ratio of not more than about 0.6. In certain embodiments, the (T_{M} + A_{M}):Al ratio is not more than 0.5, for example about 0.05 to about 0.5, about 0.1 to about 0.4, or about 0.1 to about 0.2.

In certain embodiments, Ce is post-synthesis impregnated into the JMZ-2, for example by adding Ce nitrate to a copper promoted zeolite via a conventional incipient wetness technique. Preferably, the cerium concentration in the catalyst material is present in a concentration of at least about 1 weight percent, based on the total weight of the zeolite. Examples of preferred concentrations include at least about 2.5 weight percent, at least about 5 weight percent, at least about 8 weight percent, at least about 10 weight percent, about 1.35 to about 13.5 weight percent, about 2.7 to about 13.5 weight percent, about 2.7 to about 8.1 weight percent, about 2 to about 4 weight percent, about 2 to about 9.5 weight percent, and about 5 to about 9.5 weight percent, based on the total weight of the zeolite. In certain embodiments, the cerium concentration in the catalyst material is about 50 to about 550 g /ft^3 (about 1.77 to about 19.42 g/l). Other ranges of Ce include: above 100 g /ft^3 (3.53 g/l), above 200 g /ft^3 (7.06 g/l), above 300 g /ft^3 (10.59 g/l), above 400 g /ft^3 (14.13 g/l), above 500 g /ft^3 (17.66 g/l), from about 75 to about 350 g /ft^3 (about 2.65 to about 12.36 g/l), from about 100 to about 300 g /ft^3 (about 3.53 to about 10.59 g/l), and from about 100 to about 250 g /ft^3 (about 3.53 to about 8.83 g/l).

For embodiments in which the catalyst is part of a washcoat composition, the washcoat may further comprise binder containing Ce or ceria. For such embodiments, the Ce containing particles in the binder are significantly larger than the Ce containing particles in the catalyst.

Applicants further discovered that the foregoing one-pot synthesis procedure permits adjusting the SAR of the catalyst based on the composition of the starting synthesis mixture. SARs of 10 – 50, 20 – 40, 30 – 40, 10 – 15, and 25 – 35 for example, can be selectively achieved based on the composition of the starting synthesis mixture and/or adjusting other process variables. The SAR of zeolites may be determined by conventional analysis. This ratio is meant to represent, as closely as possible, the ratio in the rigid atomic framework of the zeolite crystal and to exclude silicon or aluminium in the binder or, in cationic or other form, within the channels. It will be appreciated that it may be extremely difficult to directly measure the SAR of zeolite after it has been combined with a binder material. Accordingly, the SAR has been expressed hereinabove in
term of the SAR of the parent zeolite, i.e., the zeolite used to prepare the catalyst, as measured prior to the combination of this zeolite with the other catalyst components.

The foregoing one-pot synthesis procedure can result in zeolite crystals of uniform size and shape with relatively low amounts of agglomeration. In addition, the synthesis procedure can result in zeolite crystals having a mean crystalline size of about 0.1 to about 10 μm, for example about 0.5 to about 5 μm, about 0.1 to about 1 μm, about 1 to about 5 μm, about 3 to about 7 μm, and the like. In certain embodiments, large crystals are milled using a jet mill or other particle-on-particle milling technique to an average size of about 1.0 to about 1.5 micron to facilitate washcoating a slurry containing the catalyst to a substrate, such as a flow-through monolith.

Crystal size is the length of one edge of a face of the crystal. Direct measurement of the crystal size can be performed using microscopy methods, such as SEM and TEM. Other techniques for determining mean particle size, such as laser diffraction and scattering can also be used. In addition to the mean crystal size, catalyst compositions preferably have a majority of the crystal sizes are greater than about 0.1 μm, preferably between about 0.5 and about 5 μm, such as about 0.5 to about 5 μm, about 0.7 to about 5 μm, about 1 to about 5 μm, about 1.5 to about 5.0 μm, about 1.5 to about 4.0 μm, about 2 to about 5 μm, or about 1 μm to about 10 μm.

Catalysts of the present invention are particularly applicable for heterogeneous catalytic reaction systems (i.e., solid catalyst in contact with a gas reactant). To improve contact surface area, mechanical stability, and/or fluid flow characteristics, the catalysts can be disposed on and/or within a substrate, preferably a porous substrate. In certain embodiments, a washcoat containing the catalyst is applied to an inert substrate, such as corrugated metal plate or a honeycomb cordierite brick. Alternatively, the catalyst is kneaded along with other components such as fillers, binders, and reinforcing agents, into an extrudable paste which is then extruded through a die to form a honeycomb brick. Accordingly, in certain embodiments provided is a catalyst article comprising a JMZ-2 catalyst described herein coated on and/or incorporated into a substrate.

Certain aspects of the invention provide a catalytic washcoat. The washcoat comprising the JMZ-2 catalyst described herein is preferably a solution, suspension, or slurry. Suitable coatings include surface coatings, coatings that penetrate a portion of the substrate, coatings that permeate the substrate, or some combination thereof.
A washcoat can also include non-catalytic components, such as fillers, binders, stabilizers, rheology modifiers, and other additives, including one or more of alumina, silica, non-zeolite silica alumina, titania, zirconia, ceria. In certain embodiments, the catalyst composition may comprise pore-forming agents such as graphite, cellulose, starch, polyacrylate, and polyethylene, and the like. These additional components do not necessarily catalyse the desired reaction, but instead improve the catalytic material's effectiveness, for example, by increasing its operating temperature range, increasing contact surface area of the catalyst, increasing adherence of the catalyst to a substrate, etc. In preferred embodiments, the washcoat loading is >0.3 g/in$^3$ (>4.9 g/l), such as >1.2 g/in$^3$ (>19.7 g/l), >1.5 g/in$^3$ (>24.6 g/l), >1.7 g/in$^3$ (>27.9 g/l) or >2.00 g/in$^3$ (>32.8 g/l), and preferably < 3.5 g/in$^3$ (<57.4 g/l), such as < 2.5 g/in$^3$ (41.0 g/l). In certain embodiments, the washcoat is applied to a substrate in a loading of about 0.8 to 1.0 g/in$^3$ (about 13.1 to 16.4 g/l), 1.0 to 1.5 g/in$^3$ (16.4 to 24.6 g/l), or 1.5 to 2.5 g/in$^3$ (24.6 to 41.0 g/l).

Two of the most common substrate designs are plate and honeycomb. Preferred substrates, particularly for mobile applications, include flow-through monoliths having a so-called honeycomb geometry that comprise multiple adjacent, parallel channels that are open on both ends and generally extend from the inlet face to the outlet face of the substrate and result in a high-surface area-to-volume ratio. For certain applications, the honeycomb flow-through monolith preferably has a high cell density, for example about 600 to 800 cells per square inch (93.5 to 124 cell cm$^{-3}$), and/or an average internal wall thickness of about 0.18 – 0.35 mm, preferably about 0.20 – 0.25 mm. For certain other applications, the honeycomb flow-through monolith preferably has a low cell density of about 150 – 600 cells per square inch(23.3 to 93.5 cells cm$^{-3}$), more preferably about 200 – 400 cells per square inch (31 to 62 cells cm$^{-3}$). Preferably, the honeycomb monoliths are porous. In addition to cordierite, silicon carbide, silicon nitride, ceramic, and metal, other materials that can be used for the substrate include aluminium nitride, silicon nitride, aluminium titanate, α-alumina, mullite, e.g., acicular mullite, pollucite, a thermet such as Al$_2$OsZFe, Al$_2$O$_3$/Ni or B$_4$CZFe, or composites comprising segments of any two or more thereof. Preferred materials include cordierite, silicon carbide, and alumina titanate.

Plate-type catalysts have lower pressure drops and are less susceptible to plugging and fouling than the honeycomb types, which is advantageous in high efficiency stationary applications, but plate configurations can be much larger and more expensive. A Honeycomb
configuration is typically smaller than a plate type, which is an advantage in mobile applications, but has higher pressure drops and plug more easily. In certain embodiments the plate substrate is constructed of metal, preferably corrugated metal.

In certain embodiments, the invention is a catalyst article made by a process described herein. In a particular embodiment, the catalyst article is produced by a process that includes the steps of applying a JMZ-2 catalyst composition, preferably as a washcoat, to a substrate as a layer either before or after at least one additional layer of another composition for treating exhaust gas has been applied to the substrate. The one or more catalyst layers on the substrate, including the JMZ-2 catalyst layer, are arranged in consecutive layers. As used herein, the term “consecutive” with respect to catalyst layers on a substrate means that each layer is contact with its adjacent layer(s) and that the catalyst layers as a whole are arranged one on top of another on the substrate.

In certain embodiments, the JMZ-2 catalyst is disposed on the substrate as a first layer and another composition, such as an oxidation catalyst, reduction catalyst, scavenging component, or NOx storage component, is disposed on the substrate as a second layer. In other embodiments, the JMZ-2 catalyst is disposed on the substrate as a second layer and another composition, such as such as an oxidation catalyst, reduction catalyst, scavenging component, or NOx storage component, is disposed on the substrate as a first layer. As used herein the terms “first layer” and “second layer” are used to describe the relative positions of catalyst layers in the catalyst article with respect to the normal direction of exhaust gas flow-through, past, and/or over the catalyst article. Under normal exhaust gas flow conditions, exhaust gas contacts the first layer prior to contacting the second layer. In certain embodiments, the second layer is applied to an inert substrate as a bottom layer and the first layer is top layer that is applied over the second layer as a consecutive series of sub-layers. In such embodiments, the exhaust gas penetrates (and hence contacts) the first layer, before contacting the second layer, and subsequently returns through the first layer to exit the catalyst component. In other embodiments, the first layer is a first zone disposed on an upstream portion of the substrate and the second layer is disposed on the substrate as a second zone, wherein the second zone is downstream of the first.

In another embodiment, the catalyst article is produced by a process that includes the steps of applying a JMZ-2 catalyst composition, preferably as a washcoat, to a substrate as a first
zone, and subsequently applying at least one additional composition for treating an exhaust gas to the substrate as a second zone, wherein at least a portion of the first zone is downstream of the second zone. Alternatively, the JMZ-2 catalyst composition can be applied to the substrate in a second zone that is downstream of a first zone containing the additional composition. Examples of additional compositions include oxidation catalysts, reduction catalysts, scavenging components (e.g., for sulfur, water, etc.), or NOx storage components.

To reduce the amount of space required for an exhaust system, individual exhaust components in certain embodiments are designed to perform more than one function. For example, applying an SCR catalyst to a wall-flow filter substrate instead of a flow-through substrate serves to reduce the overall size of an exhaust treatment system by allowing one substrate to serve two functions, namely catalytically reducing NOx concentration in the exhaust gas and mechanically removing soot from the exhaust gas. Accordingly, in certain embodiments, the substrate is a honeycomb wall-flow filter or partial filter. Wall-flow filters are similar to flow-through honeycomb substrates in that they contain a plurality of adjacent, parallel channels.

However, the channels of flow-through honeycomb substrates are open at both ends, whereas the channels of wall-flow substrates have one end capped, wherein the capping occurs on opposite ends of adjacent channels in an alternating pattern. Capping alternating ends of channels prevents the gas entering the inlet face of the substrate from flowing straight through the channel and existing. Instead, the exhaust gas enters the front of the substrate and travels into about half of the channels where it is forced through the channel walls prior to entering the second half of the channels and exiting the back face of the substrate.

The substrate wall has a porosity and pore size that is gas permeable, but traps a major portion of the particulate matter, such as soot, from the gas as the gas passes through the wall. Preferred wall-flow substrates are high efficiency filters. Wall flow filters for use with the present invention preferably have an efficiency of least 70%, at least about 75%, at least about 80%, or at least about 90%. In certain embodiments, the efficiency will be from about 75 to about 99%, about 75 to about 90%, about 80 to about 90%, or about 85 to about 95%. Here, efficiency is relative to soot and other similarly sized particles and to particulate concentrations typically found in conventional diesel exhaust gas. For example, particulates in diesel exhaust can range in size
from 0.05 microns to 2.5 microns. Thus, the efficiency can be based on this range or a sub-range, such as 0.1 to 0.25 microns, 0.25 to 1.25 microns, or 1.25 to 2.5 microns.

Porosity is a measure of the percentage of void space in a porous substrate and is related to backpressure in an exhaust system: generally, the lower the porosity, the higher the backpressure. Preferably, the porous substrate has a porosity of about 30 to about 80%, for example about 40 to about 75%, about 40 to about 65%, or from about 50 to about 60%.

The pore interconnectivity, measured as a percentage of the substrate’s total void volume, is the degree to which pores, void, and/or channels, are joined to form continuous paths through a porous substrate, i.e., from the inlet face to the outlet face. In contrast to pore interconnectivity is the sum of closed pore volume and the volume of pores that have a conduit to only one of the surfaces of the substrate. Preferably, the porous substrate has a pore interconnectivity volume of at least about 30%, more preferably at least about 40%.

The mean pore size of the porous substrate is also important for filtration. Mean pore size can be determined by any acceptable means, including by mercury porosimetry. The mean pore size of the porous substrate should be of a high enough value to promote low backpressure, while providing an adequate efficiency by either the substrate per se, by promotion of a soot cake layer on the surface of the substrate, or combination of both. Preferred porous substrates have a mean pore size of about 10 to about 40 μm, for example about 20 to about 30 μm, about 10 to about 25 μm, about 10 to about 20 μm, about 20 to about 25 μm, about 10 to about 15 μm, and about 15 to about 20 μm.

In general, the production of an extruded solid body containing the JMZ-2 catalyst involves blending the JMZ-2 catalyst, a binder, an optional organic viscosity-enhancing compound into an homogeneous paste which is then added to a binder/matrix component or a precursor thereof and optionally one or more of stabilized ceria, and inorganic fibres. The blend is compacted in a mixing or kneading apparatus or an extruder. The mixtures have organic additives such as binders, pore formers, plasticizers, surfactants, lubricants, dispersants as processing aids to enhance wetting and therefore produce a uniform batch. The resulting plastic material is then moulded, in particular using an extrusion press or an extruder including an extrusion die, and the resulting mouldings are dried and calcined. The organic additives are "burnt out" during calcinations of the extruded solid body. A JMZ-2 catalyst may also be washcoated or otherwise applied to the extruded solid body.
as one or more sub-layers that reside on the surface or penetrate wholly or partly into the extruded solid body.

Extruded solid bodies containing JMZ-2 catalysts according to the present invention generally comprise a unitary structure in the form of a honeycomb having uniform-sized and parallel channels extending from a first end to a second end thereof. Channel walls defining the channels are porous. Typically, an external "skin" surrounds a plurality of the channels of the extruded solid body. The extruded solid body can be formed from any desired cross section, such as circular, square or oval. Individual channels in the plurality of channels can be square, triangular, hexagonal, circular etc. Channels at a first, upstream end can be blocked, e.g. with a suitable ceramic cement, and channels not blocked at the first, upstream end can also be blocked at a second, downstream end to form a wall-flow filter. Typically, the arrangement of the blocked channels at the first, upstream end resembles a checker-board with a similar arrangement of blocked and open downstream channel ends.

The binder/matrix component is preferably selected from the group consisting of cordierite, nitrides, carbides, borides, intermetallics, lithium aluminosilicate, a spinel, an optionally doped alumina, a silica source, titania, zirconia, titanio-titania-zirconia, zircon and mixtures of any two or more thereof. The paste can optionally contain reinforcing inorganic fibres selected from the group consisting of carbon fibres, glass fibres, metal fibres, boron fibres, alumina fibres, silica fibres, silica-alumina fibres, silicon carbide fibres, potassium titanate fibres, aluminium borate fibres and ceramic fibres.

The alumina binder/matrix component is preferably gamma alumina, but can be any other transition alumina, i.e., alpha alumina, beta alumina, chi alumina, eta alumina, rho alumina, kappa alumina, theta alumina, delta alumina, lanthanum beta alumina and mixtures of any two or more such transition aluminas. It is preferred that the alumina is doped with at least one non-aluminium element to increase the thermal stability of the alumina. Suitable alumina dopants include silicon, zirconium, barium, lanthanides and mixtures of any two or more thereof. Suitable lanthanide dopants include La, Ce, Nd, Pr, Gd and mixtures of any two or more thereof.

Sources of silica can include a silica sol, quartz, fused or amorphous silica, sodium silicate, an amorphous aluminosilicate, an alkoxy silane, a silicone resin binder such as methylphenyl silicone resin, a clay, talc or a mixture of any two or more thereof. Of this list, the
silica can be SiO₂ as such, feldspar, mullite, silica-alumina, silica-magnesia, silica-zirconia, silica-thoria, silica-berylia, silica-titania, ternary silica-alumina-zirconia, ternary silica-alumina-magnesia, ternary-silica-magnesia-zirconia, ternary silica-alumina-thoria and mixtures of any two or more thereof.

Preferably, the JMZ-2 catalyst is dispersed throughout, and preferably evenly throughout, the entire extruded catalyst body.

Where any of the above extruded solid bodies are made into a wall-flow filter, the porosity of the wall-flow filter can be from 30-80%, such as from 40-70%. Porosity and pore volume and pore radius can be measured e.g. using mercury intrusion porosimetry.

The JMZ-2 catalyst described herein can promote the reaction of a reductant, preferably ammonia, with nitrogen oxides to selectively form elemental nitrogen (N₂) and water (H₂O). Thus, in one embodiment, the catalyst can be formulated to favour the reduction of nitrogen oxides with a reductant (i.e., an SCR catalyst). Examples of such reductants include hydrocarbons (e.g., C3 – C6 hydrocarbons) and nitrogenous reductants such as ammonia and ammonia hydrazine or any suitable ammonia precursor, such as urea ((NH₂)₂CO), ammonium carbonate, ammonium carbamate, ammonium hydrogen carbonate or ammonium formate.

The JMZ-2 catalyst described herein can also promote the oxidation of ammonia. Thus, in another embodiment, the catalyst can be formulated to favour the oxidation of ammonia with oxygen, particularly a concentrations of ammonia typically encountered downstream of an SCR catalyst (e.g., ammonia oxidation (AMOX) catalyst, such as an ammonia slip catalyst (ASC)). In certain embodiments, the JMZ-2 catalyst is disposed as a top layer over an oxidative under-layer, wherein the under-layer comprises a platinum group metal (PGM) catalyst or a non-PGM catalyst. Preferably, the catalyst component in the underlayer is disposed on a high surface area support, including but not limited to alumina.

In yet another embodiment, an SCR and AMOX operations are performed in series, wherein both processes utilize a catalyst comprising the JMZ-2 catalyst described herein, and wherein the SCR process occurs upstream of the AMOX process. For example, an SCR formulation of the catalyst can be disposed on the inlet side of a filter and an AMOX formulation of the catalyst can be disposed on the outlet side of the filter.
Accordingly, provided is a method for the reduction of NO\(_x\) compounds or oxidation of NH\(_3\) in a gas, which comprises contacting the gas with a catalyst composition described herein for the catalytic reduction of NO\(_x\) compounds for a time sufficient to reduce the level of NO\(_x\) compounds and/or NH\(_3\) in the gas. In certain embodiments, provided is a catalyst article having an ammonia slip catalyst disposed downstream of a selective catalytic reduction (SCR) catalyst. In such embodiments, the ammonia slip catalyst oxidizes at least a portion of any nitrogenous reductant that is not consumed by the selective catalytic reduction process. For example, in certain embodiments, the ammonia slip catalyst is disposed on the outlet side of a wall flow filter and an SCR catalyst is disposed on the upstream side of a filter. In certain other embodiments, the ammonia slip catalyst is disposed on the downstream end of a flow-through substrate and an SCR catalyst is disposed on the upstream end of the flow-through substrate. In other embodiments, the ammonia slip catalyst and SCR catalyst are disposed on separate bricks within the exhaust system. These separate bricks can be adjacent to, and in contact with, each other or separated by a specific distance, provided that they are in fluid communication with each other and provided that the SCR catalyst brick is disposed upstream of the ammonia slip catalyst brick.

In certain embodiments, the SCR and/or AMOX process is performed at a temperature of at least 100 °C. In another embodiment, the process(es) occur at a temperature from about 150 °C to about 750 °C. In a particular embodiment, the temperature range is from about 175 to about 550 °C. In another embodiment, the temperature range is from 175 to 400 °C. In yet another embodiment, the temperature range is 450 to 900 °C, preferably 500 to 750 °C, 500 to 650 °C, 450 to 550 °C, or 650 to 850 °C. Embodiments utilizing temperatures greater than 450 °C are particularly useful for treating exhaust gases from a heavy and light duty diesel engine that is equipped with an exhaust system comprising (optionally catalysed) diesel particulate filters which are regenerated actively, e.g. by injecting hydrocarbon into the exhaust system upstream of the filter, wherein the zeolite catalyst for use in the present invention is located downstream of the filter.

According to another aspect of the invention, provided is a method for the reduction of NO\(_x\) compounds and/or oxidation of NH\(_3\) in a gas, which comprises contacting the gas with a catalyst described herein for a time sufficient to reduce the level of NO\(_x\) compounds in the gas.

Methods of the present invention may comprise one or more of the following steps: (a)
accumulating and/or combusting soot that is in contact with the inlet of a catalytic filter; (b) introducing a nitrogenous reducing agent into the exhaust gas stream prior to contacting the catalytic filter, preferably with no intervening catalytic steps involving the treatment of NO\textsubscript{x} and the reductant; (c) generating NH\textsubscript{3} over a NO\textsubscript{x} adsorber catalyst or lean NO\textsubscript{x} trap, and preferably using such NH\textsubscript{3} as a reductant in a downstream SCR reaction; (d) contacting the exhaust gas stream with a DOC to oxidize hydrocarbon based soluble organic fraction (SOF) and/or carbon monoxide into CO\textsubscript{2}, and/or oxidize NO into NO\textsubscript{2}, which in turn, may be used to oxidize particulate matter in particulate filter; and/or reduce the particulate matter (PM) in the exhaust gas; (e) contacting the exhaust gas with one or more flow-through SCR catalyst device(s) in the presence of a reducing agent to reduce the NOx concentration in the exhaust gas; and (f) contacting the exhaust gas with an ammonia slip catalyst, preferably downstream of the SCR catalyst to oxidize most, if not all, of the ammonia prior to emitting the exhaust gas into the atmosphere or passing the exhaust gas through a recirculation loop prior to exhaust gas entering/re-entering the engine.

In another embodiment, all or at least a portion of the nitrogen-based reductant, particularly NH\textsubscript{3}, for consumption in the SCR process can be supplied by a NO\textsubscript{x} adsorber catalyst (NAC), a lean NO\textsubscript{x} trap (LNT), or a NO\textsubscript{x} storage/reduction catalyst (NSRC), disposed upstream of the SCR catalyst, e.g., a SCR catalyst of the present invention disposed on a wall-flow filter. NAC components useful in the present invention include a catalyst combination of a basic material (such as alkali metal, alkaline earth metal or a rare earth metal, including oxides of alkali metals, oxides of alkaline earth metals, and combinations thereof), and a precious metal (such as platinum), and optionally a reduction catalyst component, such as rhodium. Specific types of basic material useful in the NAC include caesium oxide, potassium oxide, magnesium oxide, sodium oxide, calcium oxide, strontium oxide, barium oxide, and combinations thereof. The precious metal is preferably present at about 10 to about 200 g/ft\textsuperscript{3} (about 0.353 to about 7.063 g/l), such as 20 to 60 g/ft\textsuperscript{3} (0.706 to 2.119 g/l). Alternatively, the precious metal of the catalyst is characterized by the average concentration which may be from about 40 to about 100 grams/ft\textsuperscript{3} (about 1.413 to about 3.531 g/l).

Under certain conditions, during the periodically rich regeneration events, NH\textsubscript{3} may be generated over a NO\textsubscript{x} adsorber catalyst. The SCR catalyst downstream of the NO\textsubscript{x} adsorber catalyst may improve the overall system NO\textsubscript{x} reduction efficiency. In the combined system, the
SCR catalyst is capable of storing the released NH₃ from the NAC catalyst during rich regeneration events and utilizes the stored NH₃ to selectively reduce some or all of the NOₓ that slips through the NAC catalyst during the normal lean operation conditions.

The method for treating exhaust gas as described herein can be performed on an exhaust gas derived from a combustion process, such as from an internal combustion engine (whether mobile or stationary), a gas turbine and coal or oil fired power plants. The method may also be used to treat gas from industrial processes such as refining, from refinery heaters and boilers, furnaces, the chemical processing industry, coke ovens, municipal waste plants and incinerators, etc. In a particular embodiment, the method is used for treating exhaust gas from a vehicular lean burn internal combustion engine, such as a diesel engine, a lean-burn gasoline engine or an engine powered by liquid petroleum gas or natural gas.

In certain aspects, the invention is a system for treating exhaust gas generated by combustion process, such as from an internal combustion engine (whether mobile or stationary), a gas turbine, coal or oil fired power plants, and the like. Such systems include a catalytic article comprising the JMZ-2 catalyst described herein and at least one additional component for treating the exhaust gas, wherein the catalytic article and at least one additional component are designed to function as a coherent unit.

In certain embodiments, the system comprises a catalytic article comprising a JMZ-2 catalyst described herein, a conduit for directing a flowing exhaust gas, a source of nitrogenous reductant disposed upstream of the catalytic article. The system can include a controller for the metering the nitrogenous reductant into the flowing exhaust gas only when it is determined that the zeolite catalyst is capable of catalysing NOₓ reduction at or above a desired efficiency, such as at above 100 °C, above 150 °C or above 175 °C. The metering of the nitrogenous reductant can be arranged such that 60% to 200% of theoretical ammonia is present in exhaust gas entering the SCR catalyst calculated at 1:1 NH₃/NO and 4:3 NH₃/NO₂.

In another embodiment, the system comprises an oxidation catalyst (e.g., a diesel oxidation catalyst (DOC)) for oxidizing nitrogen monoxide in the exhaust gas to nitrogen dioxide can be located upstream of a point of metering the nitrogenous reductant into the exhaust gas. In one embodiment, the oxidation catalyst is adapted to yield a gas stream entering the SCR zeolite catalyst having a ratio of NO to NO₂ of from about 4:1 to about 1:3 by volume, e.g. at an exhaust
gas temperature at oxidation catalyst inlet of 250 °C to 450 °C. The oxidation catalyst can include at least one platinum group metal (or some combination of these), such as platinum, palladium, or rhodium, coated on a flow-through monolith substrate. In one embodiment, the at least one platinum group metal is platinum, palladium or a combination of both platinum and palladium. The platinum group metal can be supported on a high surface area washcoat component such as alumina, a zeolite such as an aluminosilicate zeolite, silica, non-zeolite silica alumina, ceria, zirconia, titania or a mixed or composite oxide containing both ceria and zirconia.

In a further embodiment, a suitable filter substrate is located between the oxidation catalyst and the SCR catalyst. Filter substrates can be selected from any of those mentioned above, e.g. wall flow filters. Where the filter is catalysed, e.g. with an oxidation catalyst of the kind discussed above, preferably the point of metering nitrogenous reductant is located between the filter and the zeolite catalyst. Alternatively, if the filter is un-catalysed, the means for metering nitrogenous reductant can be located between the oxidation catalyst and the filter.
CLAIMS:

1. A composition comprising a synthetic zeolite having an AEI framework structure and an in-situ transition metal dispersed within the cavities and channels of the zeolite.

2. The composition of claim 1, wherein the zeolite further comprises an ITE framework, provided that the AEI and ITE frameworks together are at least 70 weight percent of the zeolite based on the total weight of the zeolite, and wherein the AEI framework and ITE framework are present in a relative weight ratio of 1:99 to 99:1.

3. The composition of any of the preceding claims, wherein the AEI framework and ITE framework are present in a relative weight ratio of 4:1 to 20:1.

4. The composition of any of the preceding claims, wherein the transition metal is copper or iron.

5. The composition of any of the preceding claims, wherein the transition metal is present in an amount of 0.1 to 5 weight percent.

6. The composition of any of the preceding claims, wherein the aluminosilicate has a silica-to-alumina ratio of 10 to 50.

7. The composition of any of the preceding claims, wherein zeolite contains uniformly distributed ionic copper.

8. The composition of any of the preceding claims, wherein the zeolite contains less than 5 weight percent CuO based on the total weight of the zeolite.

9. A catalyst composition comprising a synthetic zeolite having an AEI framework structure, a SAR of 10 to 50, and containing 0.5 to 5 weight percent of non-framework copper based on the total weight of the zeolite.

10. The catalyst composition of claim 9, wherein the zeolite is free of post-synthesis exchanged copper.

11. A method for synthesizing a zeolite comprising:
preparing a reaction mixture comprising (a) at least one source of alumina, (b) at least one source of silica, (c) a transition-metal-amine organic templating agent, (d) an alkali metal hydroxide, and (e) distinct second organic templating agent, wherein each of the first and
second templating agents is suitable for forming an AEI framework structure and wherein the reaction mixture is essentially free of fluorine; heating the reaction mixture at crystallization conditions for a sufficient time to form zeolite crystals having an AEI framework and optionally an ITE framework and containing the transition metal.

12. The method of claim 11, wherein a majority of the transition metal is ionic metal dispersed in the crystallized zeolite.

13. The method of claim 11 or 12, wherein the transition-metal-amine organic templating agent is formed in-situ in the reaction mixture from individual metal and amine components.

14. The method of claim 11, 12 or 13, wherein the transition-metal-amine organic templating agent is Cu-TEPA.

15. The method of claim 14, wherein the second organic templating agent is N,N-Dimethyl-3,5-dimethylpiperidinium.

16. The method of claim 14, wherein the second organic templating agent is 1,1-Diethyl-2,6-dimethylpiperidinium.

17. The method of any of claims 11 to 16, wherein the zeolite has a SAR of 10 to 50 and contains 0.5 to 5 weight percent ionic copper.

18. The method of any of claims 11 to 17, wherein the zeolite is washed, dried, and calcined to form a Cu-AEI/ITE catalyst.

19. A zeolite composition prepared by preparing a reaction mixture comprising (a) at least one source of alumina, (b) at least one source of silica, (c) Cu-TEPA, (d) an alkali metal hydroxide, and (e) one of N,N-Dimethyl-3,5-dimethylpiperidinium or 1,1-Diethyl-2,6-dimethylpiperidinium; heating the reaction mixture at crystallization conditions for a sufficient time to form zeolite.

20. A catalyst article for treating exhaust gas comprising a catalyst composition according to any of claims 1 to 10 disposed on and/or within a honeycomb monolith substrate.

21. A method for treating an exhaust gas comprising contacting a combustion exhaust gas containing NOₓ and/or NH₃ with a catalyst article according to claim 20 to selectively
reduce at least a portion of the NO$_x$ into N$_2$ and H$_2$O and/or oxidize at least a portion of the NH$_3$. 
Application No: GB1421334.2
Claims searched: 1-21
Examiner: Mr Martin Price
Date of search: 1 June 2015

Patents Act 1977: Search Report under Section 17

Documents considered to be relevant:

<table>
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<th>Category</th>
<th>Relevant to claims</th>
<th>Identity of document and passage or figure of particular relevance</th>
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| X        | 1, 4-10, 19-21     | GB 2507902 A
Johnson Matthey - see e.g. claim 1 and Table 1               |
| X, E     | 1, 4-10, 19-21     | GB 2519648 A
Johnson Matthey - see e.g. claims 1, 2, 12                   |
| X        | 1, 4-10, 19-21     | WO 2014/141200 A1
Johnson Matthey - see e.g. claims 1-13 and the examples      |
| X        | 1, 4-10, 19-21     | WO 2013/159825 A1
Haldor Topsoe - see e.g. claim 1                              |

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Worldwide search of patent documents classified in the following areas of the IPC
B01D; B01J; C01B

The following online and other databases have been used in the preparation of this search report
EPODOC, WPI
## International Classification:

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