

US007261857B2

(12) United States Patent Suslick et al.

(54) COLORIMETRIC ARTIFICIAL NOSE HAVING AN ARRAY OF DYES AND

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METHOD FOR ARTIFICIAL OLFACTION

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(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35

U.S.C. 154(b) by 534 days.

This patent is subject to a terminal dis-

claimer.

(21) Appl. No.: 10/279,788

(22) Filed: Oct. 24, 2002

(65) **Prior Publication Data**

US 2003/0143112 A1 Jul. 31, 2003

Related U.S. Application Data

- (63) Continuation-in-part of application No. 09/705,329, filed on Nov. 3, 2000, now Pat. No. 6,495,102, which is a continuation-in-part of application No. 09/532, 125, filed on Mar. 21, 2000, now Pat. No. 6,368,558.
- (51) **Int. Cl. G01N 21/00** (2006.01)

See application file for complete search history.

(10) Patent No.: US 7,261,857 B2

(45) **Date of Patent:** *Aug. 28, 2007

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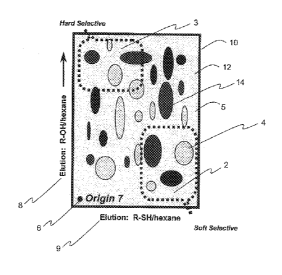
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(57) ABSTRACT

The present invention involves an artificial nose having an array comprising at least a first dye and a second dye in combination and having a distinct spectral response to an analyte. In one embodiment, the first and second dyes are from the group comprising chemoresponsive dyes, and the second dye is distinct from the first dye. In one embodiment, the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phthalocyanine, and salen, or their metal complexes. In another embodiment, the second dye is selected from the group of dyes consisting of acid-base indicator dyes and solvatochromic dyes. The present invention is particularly useful in detecting metal ligating vapors. Further, the array of the present invention can be connected to a visual display device.

22 Claims, 21 Drawing Sheets (13 of 21 Drawing Sheet(s) Filed in Color)



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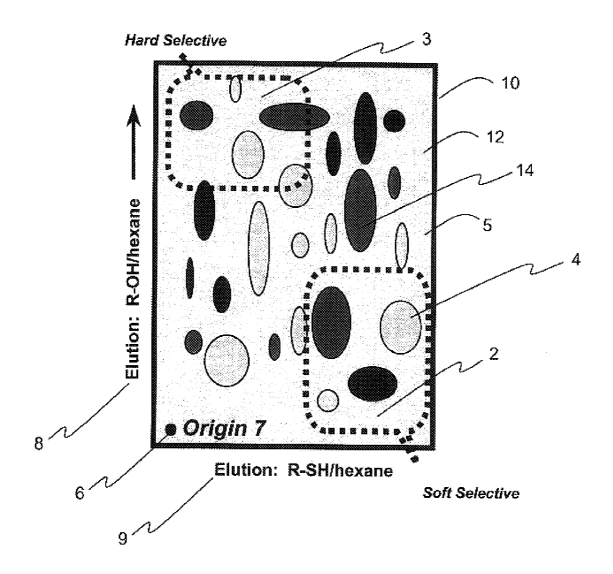
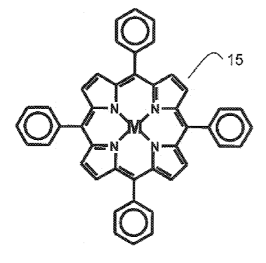


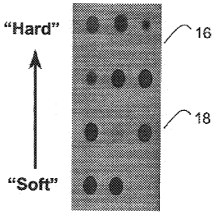
FIG. 1



Aug. 28, 2007

M(TPP)

<u>Metal</u>	Z/r Ratio (A-1)
Sn ⁴⁺	5.80
Co ₃₊	5.50
Cr³+	4.88
Min³+	4.65
Fe³÷	4.65
Co ²⁺	3.08
Cu²+	2.74
Ru²+	2.71
.Zn ²⁺	2.70
Ag ²⁺	2.13



Sn4+ Co3+ Cr3+ Mn³⁺ Fe³⁺ Co²⁺ Cu2+ Ru2+ Zn2+ Ag²⁺ 2H⁺ (FB)

FIG. 2A

FIG. 2B

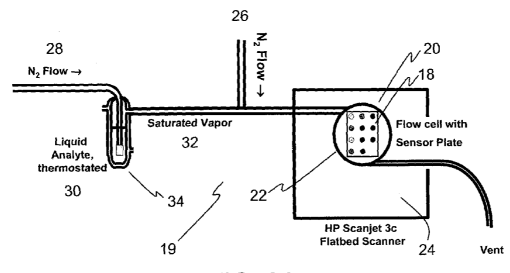


FIG. 3A

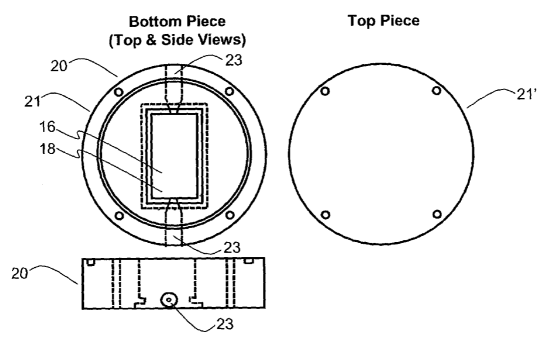


FIG. 3B

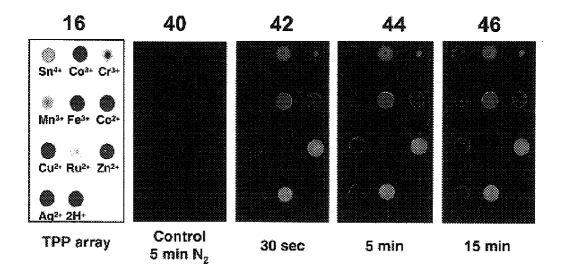


FIG. 4

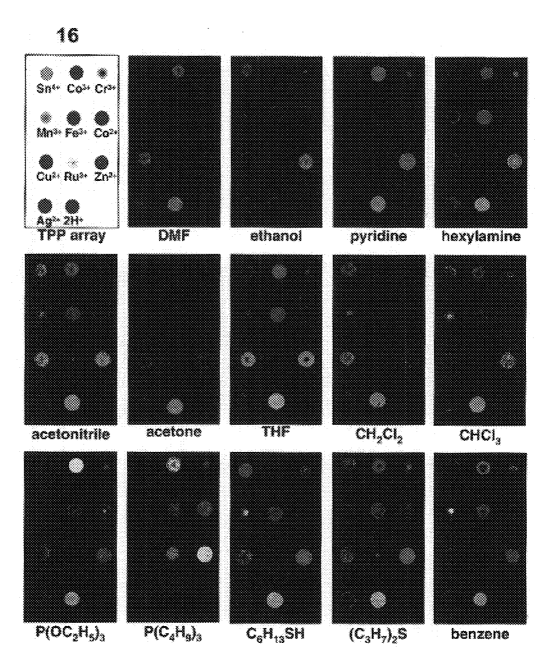


FIG. 5

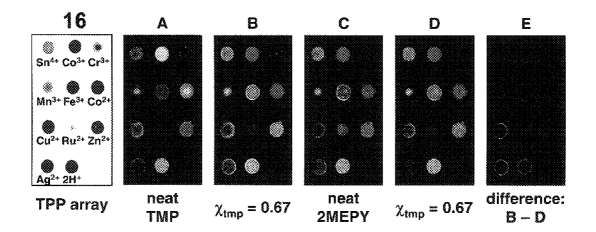


FIG. 6

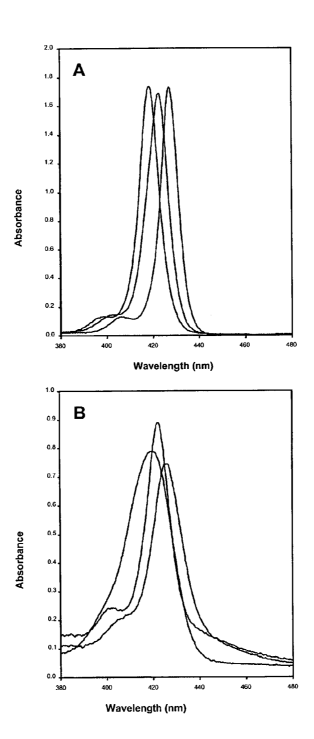


FIG. 7

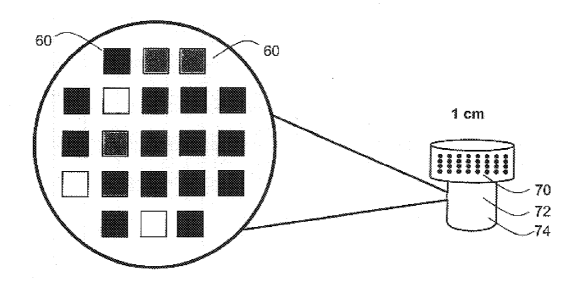


FIG. 8

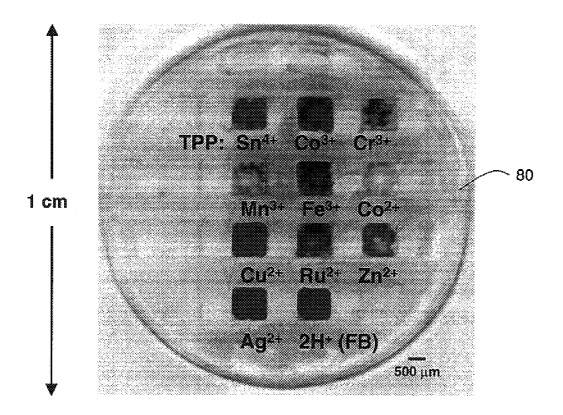


FIG. 9

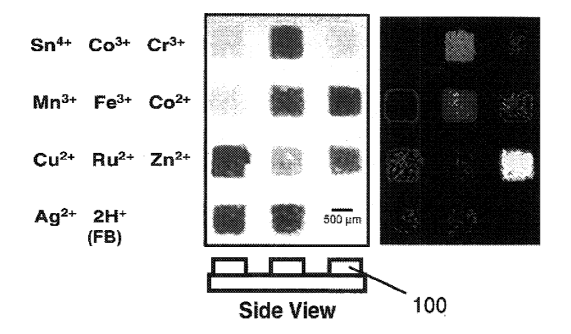


FIG. 10

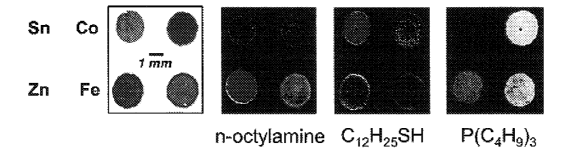


FIG. 11

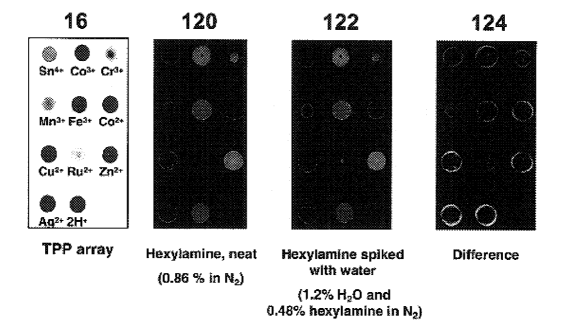
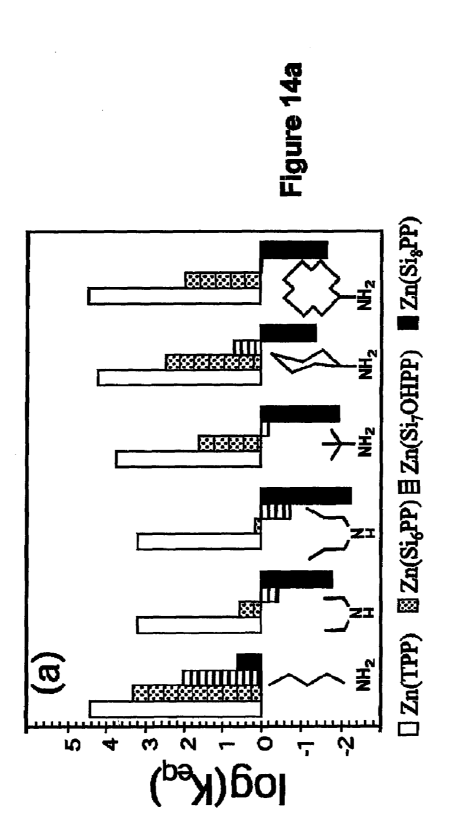
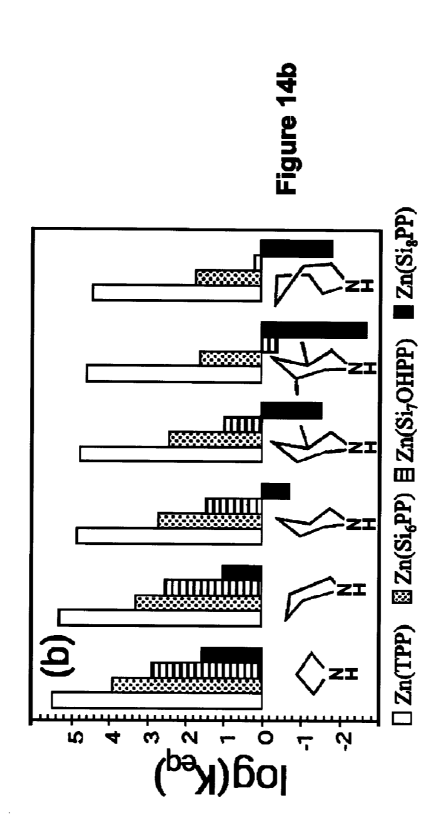
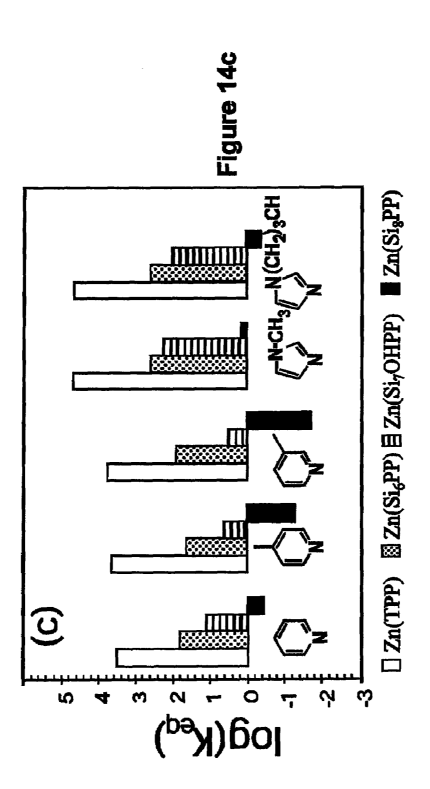


FIG. 12

Figure 13







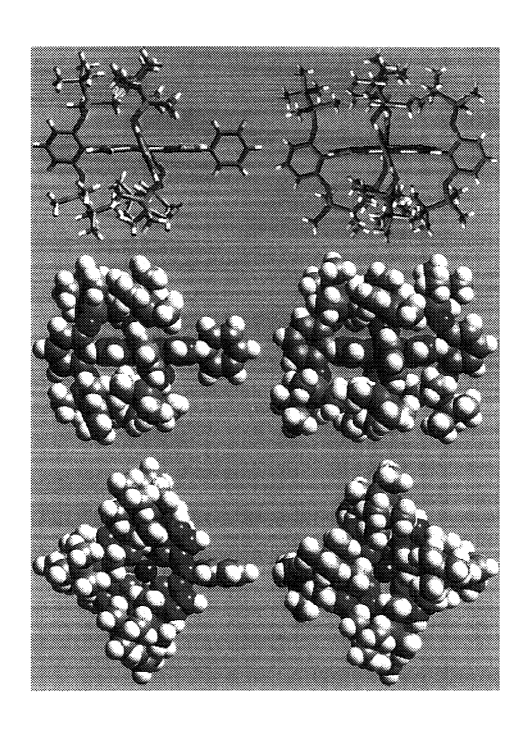


FIG. 15

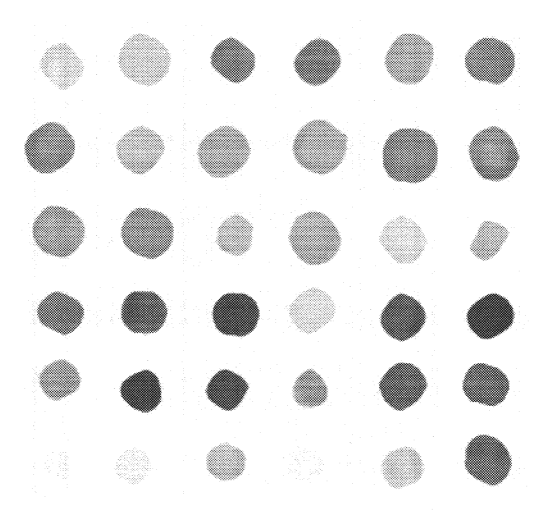


FIG. 16

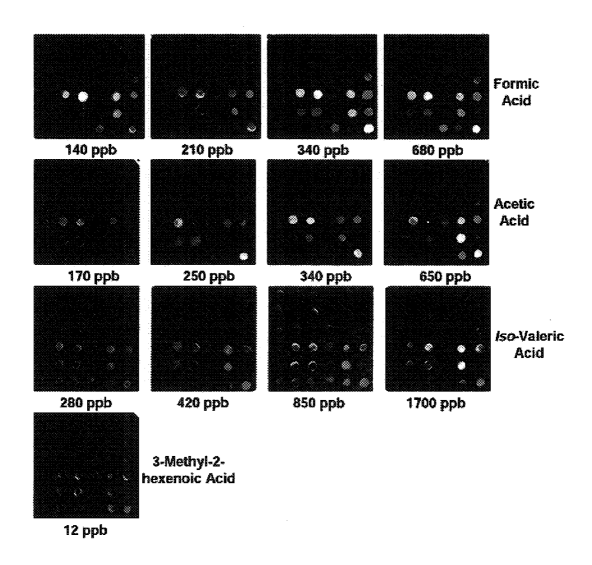


FIG. 17

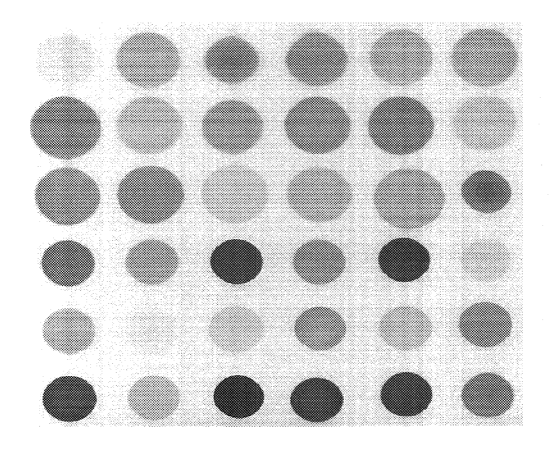
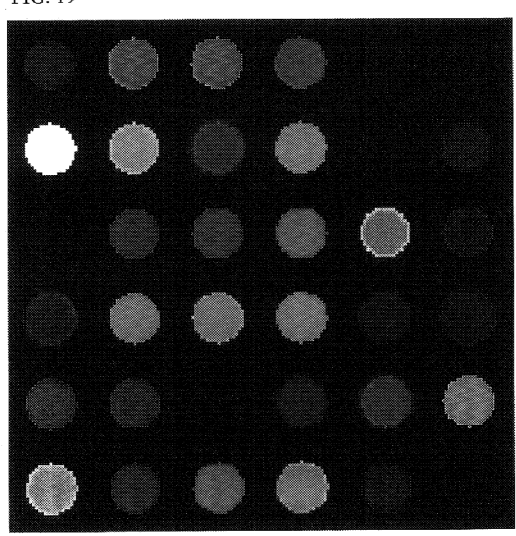


FIG. 18

FIG. 19



COLORIMETRIC ARTIFICIAL NOSE HAVING AN ARRAY OF DYES AND METHOD FOR ARTIFICIAL OLFACTION

CONTINUING APPLICATION DATA

This application is a Continuation-in-Part of U.S. application Ser. No. 09/705,329, filed on Nov. 3, 2000, now U.S. Pat. No. 6,495,102, which is a Continuation-in-Part of U.S. application Ser. No. 09/532,125, filed on Mar. 21, 2000, now 10 U.S. Pat. No. 6,368,558.

This invention was made with Government support under Contract Nos. HL25934 awarded by the National Institutes of Health & Contract No. DAAG55-97-1-2211 awarded by the Department of the Army. The Government has certain 15 rights in the invention.

FIELD OF THE INVENTION

The present invention relates to methods and apparatus 20 for artificial olfaction, e.g., artificial noses, for the detection of odorants by a visual display.

BACKGROUND OF THE INVENTION

There is a great need for olfactory or vapor-selective detectors (i.e., "artificial noses") in a wide variety of applications. For example, there is a need for artificial noses that can detect low levels of odorants and/or where odorants may be harmful to humans, animals or plants. Artificial noses that can detect many different chemicals are desirable for personal dosimeters in order to detect the type and amount of odorants exposed to a human, the presence of chemical poisons or toxins, the spoilage in foods, the presence of flavorings, or the presence of vapor emitting items, such as plant materials, fruits and vegetables, e.g., at customs portals.

Conventional artificial noses have severe limitations and disadvantages and are not considered generally useful for such purposes. Limitations and disadvantages of conventional artificial noses include their need for extensive signal transduction hardware, and their inability to selectively target metal-coordinating vapors and toxins. In addition, artificial noses which incorporate mass sensitive signal transduction or polar polymers as sensor elements are susceptible to interference by water vapor. This limitation is significant in that it can cause variable response of the detector with changes ambient humidity. See F. L. Dickert, O. Hayden, Zenkel, M. E. *Anal. Chem.* 71, 1338 (1999).

Initial work in the field of artificial noses was conducted 50 by Wilkens and Hatman in 1964, though the bulk of research done in this area has been carried out since the early 1980's. See, e.g., W. F. Wilkens, A. D. Hatman. *Ann. NY Acad. Sci.*, 116, 608 (1964); K. Pursaud, G. H. Dodd. *Nature*, 299, 352-355 (1982); and J. W. Gardner, P. N. Bartlett. *Sensors* 55 and Actuators B, 18-19, 211-220 (1994).

Vapor-selective detectors or "artificial noses" are typically based upon the production of an interpretable signal or display upon exposure to a vapor emitting substance or odorant (hereinafter sometimes referred to as an "analyte"). 60 More specifically, typical artificial noses are based upon selective chemical binding or an interface between a detecting compound of the artificial nose and an analyte or odorant, and then transforming that chemical binding into a signal or display, i.e., signal transduction.

Polymer arrays having a single dye have been used for artificial noses. That is, a series of chemically-diverse poly2

mers or polymer blends are chosen so that their composite response distinguishes a given odorant or analyte from others. Examples of polymer array vapor detectors, including conductive polymer and conductive polymer/carbon black composites, are discussed in: M. S. Freund, N. S. Lewis, *Proc. Natl. Acad. Sci. USA* 92, 2652-2656 (1995); B. J. Doleman, R. D. Sanner, E. J. Severin, R. H. Grubbs, N. S. Lewis, *Anal. Chem.* 70, 2560-2564 (1998); T. A. Dickinson, J. White, J. S. Kauer, D. R. Walt, *Nature* 382, 697-700 (1996)(polymer array with optical detection); A. E. Hoyt, A. J. Ricco, H. C. Yang, R. M. Crooks, *J. Am. Chem. Soc.* 117, 8672 (1995); and J. W. Grate, M. H. Abraham, *Sensors and Actuators B* 3, 85-111 (1991).

Other interface materials include functionalized self-assembled monolayers (SAM), metal oxides, and dendrimers. Signal transduction is commonly achieved with mass sensitive piezoelectric substrates, surface acoustic wave (SAW) transducers, or conductive materials. Optical transducers (based on absorbance or luminescence) have also been examined. Examples of metal oxide, SAM, and dendrimerbased detectors are discussed in J. W. Gardner, H. V. Shurmer, P. Corcoran, *Sensors and Actuators B* 4, 117-121 (1991); J. W. Gardner, H. V. Shurmer, T. T. Tan, *Sensors and Actuators B* 6, 71-75 (1992); and R. M. Crooks, A. J. Ricco, *Acc. Chem. Res.* 31, 219-227 (1998). These devices also use a single dye.

Techniques have also been developed using a metalloporphyrin for optical detection of a specific, single gas such as oxygen or ammonia, and for vapor detection by chemically interactive layers on quartz crystal microbalances. See A. E. Baron, J. D. S. Danielson, M. Gouterman, J. R. Wan, J. B. Callis, Rev. Sci. Instrum. 64, 3394-3402 (1993); J. Kavandi, et al., Rev. Sci. Instrum. 61, 3340-3347 (1990); W. Lee, et al., J. Mater. Chem. 3, 1031-1035 (1993); A. A. Vaughan, M. G. Baron, R. Narayanaswamy, Anal Comm. 33, 393-396 (1996); J. A. J. Brunink, et al., Anal. Chim. Acta 325, 53-64 (1996); C. Di Natale, et al., Sensors and Actuators B 44, 521-526 (1997); and C. DiNatale, et al., Mat. Sci. Eng. C 5, 209-215 (1998). However, these techniques either require extensive signal transduction hardware, or, as noted above, are limited to the detection of a specific, single gas. They are also subject to water vapor interference problems, as discussed previously.

While typical systems to date have demonstrated some success in chemical vapor detection and differentiation, these systems have focused on the detection of non-metal binding or non-metal ligating solvent vapors, such as arenes, halocarbons and ketones. Detection of metal-ligating vapors (such as amines, thiols, and phosphines) has been much less explored. Further, while some single porphyrin based sensors have been used for detection of a single strong acid, there is a need for sensor devices that will detect a wide variety of vapors.

To summarize, there are a number of limitations and drawbacks to typical artificial noses and single porphyrin based sensors. As noted above typical artificial noses are not designed for metal binding and metal ligating vapors, such as amines, thiols, and phosphines. Further, typical artificial noses require extensive signal transduction hardware, and are subject to interference from water vapor. As noted above, single porphyrin based sensors have been used for detection of a single strong acid, but cannot detect a wide variety of vapors. Thus, there is a need for new artificial noses and methods that overcome these and other limitations of prior artificial noses and single porphyrin based sensors and methods.

SUMMARY OF THE INVENTION

The present invention comprises an array of dyes including at least a first dye and a second dye which in combination provide a spectral response distinct to an analyte or odorant.

The dyes of the present invention produce a response in the spectrum range of about 200 nanometers to 2,000 nanometers, which includes the visible spectrum of light. It has now been discovered that an array of two or more dyes responds to a given ligating species with a unique color pattern spectrally and in a time dependent manner. Thus, dyes in the array of the present invention are capable of changing color in a distinct manner when exposed to any one analyte or odorant. The pattern of colors manifested by the multiple dyes is indicative of a specific or given analyte. In other words, the pattern of dye colors observed is indicative of a particular vapor or liquid species.

In a preferred embodiment, the dyes of the array are porphyrins. In another preferred embodiment, the porphyrin dyes are metalloporphyrins. In a further preferred embodiment, the array will comprise ten to fifteen distinct metalloporphyrins in combination. Metalloporphyrins are preferable dyes in the present invention because they can coordinate metal-ligating vapors through open axial coordination sites, and they produce large spectral shifts upon binding of or interaction with metal-ligating vapors. In addition, porphyrins, metalloporphyrins, and many dyes show significant color changes upon changes in the polarity of their environment; this so-called solvatochromic effect will give net color changes even in the absence of direct bonding between the vapor molecules and the metal ions. Thus, metalloporphyrins produce intense and distinctive changes in coloration upon ligand binding with metal ligating vapors.

The present invention provides a means for the detection or differentiation and quantitative measurement of a wide range of ligand vapors, such as amines, alcohols, and thiols. Further, the color data obtained using the arrays of the present innovation may be used to give a qualitative fingerprint of an analyte, or may be quantitatively analyzed to allow for automated pattern recognition and/or determination of analyte concentration. Because porphyrins also exhibit wavelength and intensity changes in their absorption bands with varying solvent polarity, weakly ligating vapors (e.g., arenes, halocarbons, or ketones) are also differentiable.

Diversity within the metalloporphyrin array may be obtained by variation of the parent porphyrin, the porphyrin metal center, or the peripheral porphyrin substituents. The parent porphyrin is also referred to as a free base ("FB") 50 porphyrin, which has two central nitrogen atoms protonated (i.e., hydrogen cations bonded to two of the central pyrrole nitrogen atoms). A preferred parent porphyrin is depicted in FIG. 2A, with the substitution of a two hydrogen ion for the metal ion (depicted as "M") in the center of the porphyrin. 55 In FIG. 2A, TTP stands for 5,10,15,20-tetraphenylporphyrinate(-2).

In accordance with the present invention, calorimetric difference maps can be generated by subtracting unexposed and exposed metalloporphyrin array images (obtained, for 60 example, with a common flatbed scanner or inexpensive video or charge coupled device ("CCD") detector) with image analysis software. This eliminates the need for extensive and expensive signal transduction hardware associated with previous techniques (e.g., piezoelectric or semiconductor sensors). By simply differencing images of the array before and after exposure to analytes, the present invention

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provides unique color change signatures for the analytes, for both qualitative recognition and quantitative analysis.

Sensor plates which incorporate vapor sensitive combinations of dyes comprise an embodiment of the present invention which is economical, disposable, and can be utilized to provide qualitative and/or quantitative identification of an analyte. In accordance with the present invention, a catalog of arrays and the resultant visual pattern for each analyte can be coded and placed in a look-up table or book for future reference. Thus, the present invention includes a method of detecting an analyte comprising the steps of forming an array of at least a first dye and a second dye, subjecting the array to an analyte, inspecting the first and second dyes for a spectral response, and comparing the spectral response with a catalog of analyte spectral responses to identify the analyte.

Because sensing is based upon either covalent interaction (i.e., ligation) or non-covalent solvation interactions between the analyte and the porphyrin array, a broad spectrum of chemical species is differentiable. While long response times (e.g., about 45 minutes) are observed at low analyte concentrations of about 1 ppm with reverse phase silica gel plates, use of impermeable solid supports (such as polymer- or glass-based micro-array plates) substantially increases the low-level response to less than 5 minutes.

Thus, it is an object of the present invention to provide methods and devices for artificial olfaction, vapor-selective detectors or artificial noses for a wide variety of applications. It is another object of the present invention to provide methods of detection and artificial noses that can detect low levels of odorants and/or where odorants may be harmful to living human, animal or plant cells. It is also an object of the present invention to provide methods of olfactory detection and artificial noses that can detect and quantify many different chemicals for dosimeters that can detect chemical poisons or toxins, that can detect spoilage in foods, that can detect flavorings and additives, and that can detect plant materials, e.g., fruits and vegetables.

Another object of the present invention is to provide for the detection of analytes using data analysis/pattern recognition techniques, including automated techniques.

Another object of the invention is to provide an artificial nose comprising an array, the array comprising at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflectance response to an analyte wherein the first dye and the second are selected from the group of dyes consisting of chemoresponsive dyes, and the second dye is distinct from the first dye. In one embodiment, the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phtahlocyanine, and salen and their metal complexes. In another embodiment, the second dye is selected from the group consisting of acid-base indicator dyes and solvatochromic dyes.

Another object of the invention is to provide a method of detecting an analyte comprising the steps of: (a) forming an array of at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflectance response to an analyte wherein the first dye and the second dye are selected from the group consisting of chemoresponsive dyes, and the second dye is distinct from the first dye, (b) subjecting the array to an analyte, (c) inspecting the array for a distinct and direct spectral absorbance or reflectance response, and (d) correlating the distinct and direct spectral

response to the presence of the analyte. In one embodiment, the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phtahlocyanine, and salen and their metal complexes. In another embodiment, the second dye is selected from the group consisting of acid-base 5 indicator dyes and solvatochromic dyes.

Another object of the invention is to provide an artificial tongue comprising an array, the array comprising at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflectance response to an analyte wherein the first dye and the second are selected from the group of dyes consisting of chemoresponsive dyes, and the second dye is distinct from the first dye. In one embodiment, the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phtahlocyanine, and salen and their metal complexes. In another embodiment, the second dye is selected from the group consisting of acid-base indicator dyes and solvatochromic dyes.

BRIEF DESCRIPTION OF THE DRAWINGS

The file of this patent contains at least one drawing executed in color. Copies of this patent with color 25 drawing(s) will be provided by the Patent and Trademark Office upon request and payment of the necessary fee.

- FIG. 1 illustrates an embodiment of the optical sensing plate of the present invention using a first elution in the y axis and a second elution in the x axis of the plate. In this 30 embodiment the first elution R—OH/hexane and the second elution is R—SH/hexane.
- FIG. 2A illustrates an embodiment of the invention using metalloporphyrins as the sensing dyes.
- FIG. 2B illustrates an embodiment of the invention using 35 metalloporphyrins as the sensing dyes.
- FIG. 3A illustrates a vapor exposure apparatus for demonstration of the present invention.
- FIG. 3B illustrates a vapor exposure apparatus for demonstration of the present invention.
- FIG. 4 illustrates the color change profile in a metalloporphyrin array of FIG. 2 when used in the vapor exposure apparatus of FIG. 3A to detect n-butylamine. Metalloporphyrins were immobilized on reverse phase silica gel plates.
- FIG. 5 illustrates a comparison of color changes at saturation for a wide range of analytes. Each analyte was delivered to the array as a nitrogen stream saturated with the analyte vapor at 20° C. DMF stands for dimethylformamide; THF stands for tetrahydrofuran.
- FIG. 6 illustrates two component saturation responses of $_{50}$ mixtures of 2-methylpyridine and trimethylphosphite. Vapor mixtures were obtained by mixing two analyte-saturated $\rm N_2$ streams at variable flow ratios.
- FIG. 7 illustrates a comparison of Zn(TPP) spectral shifts upon exposure to ethanol and pyridine (py) in methylene 55 chloride solution (A) and on the reverse phase support (B).
- FIG. **8** illustrates another embodiment of the present invention, and more particularly, an small array comprising microwells built into a wearable detector which also contains a portable light source and a light detector, such as a 60 charge-coupled device (CCD) or photodiode array.
- FIG. 9 illustrates another embodiment of the present invention, and more particularly, a microwell porphyrin array wellplate constructed from polydimethylsiloxane (PDMS).
- FIG. 10 illustrates another embodiment of the present invention, and more particularly, a microplate containing

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machined teflon posts, upon which the porphyrin array is immobilized in a polymer matrix (polystyrene/dibutylphthalate).

- FIG. 11 illustrates another embodiment of the present invention, showing a microplate of the type shown in FIG. 10, consisting of a minimized array of four metalloporphyrins, showing the color profile changes for n-octylamine, dodecanethiol, and tri-n-butylphosphine, each at 1.8 ppm.
- FIG. 12 illustrates the immunity of the present invention to interference from water vapor.
- FIG. 13 illustrates the synthesis of siloxyl-substituted bis-pocket porphyrins in accordance with the present invention.
- FIGS. **14**a, **14**b, and **14**c illustrate differences in K_{eq} for various porphyrins.
- FIG. 15 illustrates molecular models of $Zn(Si_6PP)$ (left column) and $Zn(Si_8PP)$ (right column).
- FIG. **16** illustrates an array containing illustrative examples of porphyrin, metalloporphyrin, acid-base indicator, and solvatochromatic dyes.
 - FIG. 17 illustrates the response of the array described in FIG. 16 to acid vapors, specifically formic acid, acetic acid, iso-valeric acid, and 3-methyl-2-hexenoic acid.
 - FIG. **18** illustrates a preferred array containing illustrative examples of porphyrin, metalloporphyrin, acid-base indicator, and solvatochromatic dyes.
 - FIG. 19 illustrates the response of the array described in FIG. 18 to acetone.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Production of the Sensor Plate of the Present Invention

A sensor plate 10 fabricated in accordance with the present invention is shown in FIG. 1. Sensor plate 10 comprises a two-dimensionally spatially resolved array 12 of various sensing elements or dyes 14 capable of changing color upon interaction (e.g., binding, pi-pi complexation, or polarity induced shifts in color). As shown in FIG. 1, a library of such dyes 14 can be given spatial resolution by two-dimensional chromatography or by direct deposition, including, but not limited to, ink-jet printing, micropipette spotting, screen printing, or stamping. In FIG. 1, metalloporphyrin mixture 6 is placed at origin 7. Next, the metalloporphyrin mixture 6 is eluted through a silica gel or reversedphase silica gel 5 in sensor plate 10, and the metalloporphyrins are spatially resolved from each other and immobilized in silica gel 5 as depicted by the oval and circular shapes 4 as shown in FIG. 1. Sensor plate 10 can be made from any suitable material or materials, including but not limited to, chromatography plates, paper, filter papers, porous membranes, or properly machined polymers, glasses, or metals.

FIG. 1 also illustrates an embodiment of the optical sensing plate of the present invention using a first elution 8 in the y axis and a second elution 9 in the x axis of sensor plate 10. In this embodiment, the first elution 8 is R—OH/hexane and the second elution 9 is R—SH/hexane. The order of the first and second elutions can be reversed. The first and second elutions are used to spatially resolve the metalloporphyrin mixture 6 in silica gel 5. As shown in FIG. 1, the upper left hand quadrant 3 is characterized by metalloporphyrins that are "hard" selective, i.e., having a metal center having a high chemical hardness, i.e., a high charge density. As shown in FIG. 1, the lower right hand quadrant 2 is characterized by metalloporphyrins that are "soft" selective,

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i.e., having a metal center having a low chemical hardness, i.e., a low charge density. In accordance with the present invention, the array can be a spatially resolved collection of dyes, and more particularly a spatially resolved combinatorial family of dyes.

In accordance with the present invention, a porphyrin—metalloporphyrin sensor plate was prepared and then used to detect various odorants. More specifically, solutions of various metalated tetraphenylporphyrins in either methylene chloride or chlorobenzene were spotted in 1 µL aliquots onto two carbon ("C2", i.e, ethyl-capped) reverse phase silica thin layer chromatography plates (Product No. 4809-800, by Whatman, Inc., Clifton, N.J.) to yield the sensor array 16 seen in FIG. 2B. As shown in FIG. 2B and summarized in Table 1 below, the dyes have the following colors (the exact colors depend, among other things, upon scanner settings).

TABLE 1

	(Summarizing Colors of Dyes in	FIG. 2B)
Sn ⁴⁺ - Green Mn ³⁺ - Green	Co ³⁺ - Red Fe ³⁺ - Dark Red	Cr ³⁺ - Deep Green Co ²⁺ - Red
Cu ²⁺ - Red Ag ²⁺ - Red	Ru ²⁺ - Light Yellow 2H ⁺ (Free Base "FB") - Red	Zn ²⁺ - Greenish Red

A metalloporphyrin 15, sometimes referred to as M(TPP), of the present invention is depicted in FIG. 2A. FIG. 2A also depicts various metals of the metalloporphyrins 15 of the present invention, and corresponding metal ion charge to 30 radius ratio (i.e., Z/r Ratio) in reciprocal angstroms. The Z/r Ratio should preferably span a wide range in order to target a wide range of metal ligating analytes. These metalloporphyrins have excellent chemical stability on the solid support and most have well-studied solution ligation chemistry. 35 Reverse phase silica was chosen as a non-interacting dispersion medium for the metalloporphyrin array 16 depicted in FIG. 2B, as well as a suitable surface for diffuse reflectance spectral measurements. More importantly, the reverse phase silica presents a hydrophobic interface, which virtually eliminates interference from ambient water vapor. After spotting, sensor plates 18 like the one depicted in FIG. 2B were dried under vacuum at 50° C. for 1 hour prior to use. Thus, immobilization of the metalloporphyrins on a reverse phase silica support is obtained. While ten (10) different 45 metalloporphyrins are shown in FIG. 2A, those of skill in the art will recognize that many other metalloporphyrins are useful in accordance with the present invention. Those of skill in the art will further recognize that in accordance with the broad teachings of the present invention, any dyes 50 capable of changing color upon interacting with an analyte, both containing and not containing metal ions, are useful in the array of the present invention.

Colorimetric Analysis Using the Sensor Plate

For the detection and analysis of odorants in accordance with the present invention, one needs to monitor the absorbance of the sensor plate at one or more wavelengths in a spatially resolved fashion. This can be accomplished with an imaging spectrophotometer, a simple flatbed scanner (e.g. a Hewlett Packard Scanjet 3c), or an inexpensive video or CCD camera.

FIG. 3A illustrates a vapor exposure apparatus 19 of the present invention. FIG. 3B illustrates top and side views of bottom piece 21 and a top view of top piece 21' of a vapor 65 exposure flow cell 20 of the present invention. In an embodiment of the present invention for purposes of demonstration,

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each sensor plate 18 was placed inside of a stainless steel flow cell 20 equipped with a quartz window 22 as shown in FIGS. 3A and 3B. Scanning of the sensor plate 18 was done on a commercially available flatbed scanner 24 (Hewlett Packard Scanjet 3c) at 200 dpi resolution, in full color mode. Following an initial scan, a control run with a first pure nitrogen flow stream 26 was performed. The array 16 of plate 18 was then exposed to a second nitrogen flow stream 28 saturated with a liquid analyte 30 of interest. As shown in FIG. 3A, the nitrogen flow stream 28 saturated with liquid analyte 30 results in a saturated vapor 32. Saturated vapor 32, containing the analyte 30 of interest were generated by flowing nitrogen flow stream 28 at 0.47 L/min. through the neat liquid analyte 30 in a water-jacketed, glass fritted bubbler 34. Vapor pressures were controlled by regulating the bubbler 34 temperature. As shown in FIG. 3B, vapor channels 23 permit vapor flow to sensor plate 18.

EXAMPLE 1

Scanning at different time intervals and subtracting the red, green and blue ("RGB") values of the new images from those of the original scan yields a color change profile. This is shown for n-butylamine in FIG. 4, in which color change profiles of the metalloporphyrin sensor array 16 as a function of exposure time to n-butylamine vapor. Subtraction of the initial scan from a scan after 5 min. of N_2 exposure was used as a control, giving a black response, as shown. 9.3% n-butylamine in N_2 was then passed over the array and scans made after exposure for 30 s, 5 min., and 15 min. The red, green and blue ("RGB") mode images were subtracted (absolute value) to produce the color change profiles illustrated. Virtually all porphyrins are saturated after 30 seconds of exposure, yielding a color fingerprint unique for each class of analytes, which is illustrated in FIG. 4.

More specifically, subtraction of the initial scan 40 from a scan after 5 min. of N₂ exposure was used as a control, giving a black response, as shown in FIG. 4. A nitrogen flow stream containing 0.093% n-butylamine was then passed over the array 16 and scans 42, 44, and 46 were made after exposure for 30 seconds, 5 minutes, and 15 minutes, respectively. The RGB mode images were subtracted (absolute value) using Adobe PhotoshopTM (which comprises standard image analyzing software), with contrast enhancement by expanding the pixel range (a 32 value range was expanded to 256 each for the R, G, and B values). Subtraction of exposed and unexposed images gives color change patterns that vary in hue and intensity. Because differentiation is provided by an array of detectors, the system has parallels the mammalian olfactory system. As shown in FIG. 4 and summarized in Table 2 below, the dyes have the following colors in scans 42, 44, and 46.

TABLE 2

	(Summarizing Co	olors of Dyes in FIG. 4, Sc	ans 42, 44, and 46)
60	Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - No Change Ag ²⁺ - No Change	Co ³⁺ - Green Fe ³⁺ - Red Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") — Light Blue	Cr^{3+} - Green Co^{2+} - Faint Green Zn^{2+} - Light Green

As summarized in Table 3 below, for the TTP array 16 depicted on the left-hand side of FIG. 4, the dyes have the following colors.

TABLE 3

Sn ⁴⁺ - Greenish Yellow	Co ³⁺ - Red	Cr3+ - Yellow with Dark Red
		Center
Mn ³⁺ - Greenish Yellow	Fe ³⁺ - Dark Red	Co ²⁺ - Red
Cu ²⁺ - Red	Ru ²⁺ - Light Yellow	Zn ²⁺ - Red
Ag ²⁺ - Red	2H+ (Free Base "FB") - Red	

EXAMPLE 2

Visible spectral shifts and absorption intensity differences occur upon ligation of the metal center, leading to readily observable color changes. As is well known to those with skill in the art, the magnitude of spectral shift correlates with the polarizability of the ligand; hence, there exists an electronic basis for analyte distinction. Using metal centers that span a range of chemical hardness and ligand binding affinity, a wide range of volatile analytes (including soft ligands, such as thiols, and harder ligands, such as amines) are differentiable. Because porphyrins have been shown to exhibit wavelength and intensity changes in their absorption

bands with varying solvent polarity, it is contemplated that the methods and apparatus of the present invention can be used to calorimetrically distinguish among a series of weakly ligating solvent vapors (e.g., arenes, halocarbons, or ketones), as shown for example in FIG. 5.

A comparison of color changes at saturation for a wide range of analytes is shown in FIG. 5. Each analyte is identified under the colored array 16 that identifies each analyte.). DMF stands for the analyte dimethylformamide, and THF stands for the analyte tetrahydrofuran. As shown in FIG. 5 and summarized in Table 4 below, the colors of each dye in response to a particular analyte are as follows.

TABLE 4

	TADLE 4	
	Analyte: DMF	
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - Blue Ag ²⁺ - No Change	Co ³⁺ - Green Fe ³⁺ - No Change Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - Blue Analyte: Ethanol	Cr ³⁺ - No Change Co ²⁺ - No Change Zn ²⁺ - No Change
Sn ⁴⁺ - Dark Blue Mn ³⁺ - No Change Cu ²⁺ - No Change Ag ²⁺ - No Change	Co ³⁺ - No Change Fe ³⁺ - No Change Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - No Change Analyte: Pyridine	Cr ³⁺ - Red Co ²⁺ - No Change Zn ²⁺ - Blue
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - No Change Ag ²⁺ - No Change	Co ³⁺ - Green Fe ³⁺ - No Change Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - Blue Analyte: Hexylamine	Cr ³⁺ - Dark Green Co ²⁺ - No Change Zn ²⁺ - Green
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - Blue Ag ²⁺ - Dark Blue	Co ³⁺ - Dark Green Fe ³⁺ - Red Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - Blue Analyte: Acetonitrile	Cr ³⁺ - Green Co ²⁺ - No Change Zn ²⁺ - Green
$\mathrm{Sn^{4+}}$ - Blue $\mathrm{Mn^{3+}}$ - Yellow $\mathrm{Cu^{2+}}$ - Blue $\mathrm{Ag^{2+}}$ - No Change	Co ³⁺ - Dark Green Fe ³⁺ - Dark Green Ru ²⁺ - Blue (faint dot) 2H ⁺ (Free Base "FB") - Blue Analyte: Acetone	Cr ³⁺ - No Change Co ²⁺ - No Change Zn ²⁺ - Blue
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - Dark Blue Ag ²⁺ - No Change	Co ³⁺ - No Change Fe ³⁺ - No Change Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - Blue Analyte: THF	Cr ³⁺ - Red (small dot) Co ²⁺ - No Change Zn ²⁺ - Dark Blue
Sn^{4+} - Dark Blue Mn^{3+} - Blue (small dot) Cu^{2+} - Blue Ag^{2+} - No Change	Co^{3+} - Green Fe^{3+} - Dark Green Ru^{2+} - No Change $2H^+$ (Free Base "FB") - Blue Analyte: CH_2Cl_2	Cr^{3+} - Red Co^{2+} - No Change Zn^{2+} - Blue
Sn ⁴⁺ - Dark Blue Mn ³⁺ - Yellow and Red (small dot)	Co ³⁺ - No Change Fe ³⁺ - No Change	Cr ³⁺ - No Change Co ²⁺ - No Change
Cu ²⁺ - Dark Blue Ag ²⁺ - No Change	Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - Blue	Zn ²⁺ - No Change

TABLE 4-continued

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	Analyte: CHCl ₃	
Sn^{4+} - Dark Blue Mn^{3+} - Yellow Cu^{2+} - Dark Blue (very faint) Ag^{2+} - Blue (very faint)	Co^{3+} - Dark Green Fe^{3+} - Dark Green (very faint) Ru^{2+} - No Change $2H^+$ (Free Base "FB") - Blue Analyte: $P(OC_2H_5)_3$	Cr^{3+} - Yellow (circle) Co^{2+} - No Change Zn^{2+} - Blue
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - Dark Blue (faint) Ag ²⁺ - Blue (very faint)	Co^{3+} - Yellow Fe^{3+} - Dark Green (very faint) Ru^{2+} - No Change $2H^+$ (Free Base "FB") - Blue Analyte: $P(C_4H_9)_3$	Cr^{3+} - Dark Green Co^{2+} - Greenish Yellow Zn^{2+} - Greenish Blue
Sn^{4+} - No Change Mn^{3+} - No Change Cu^{2+} - No Change Ag^{2+} - No Change	Co ³⁺ - Yellow and Red Fe ³⁺ - Dark Green (faint) Ru ²⁺ - Dark Blue 2H ⁺ (Free Base "FB") - No Change Analyte: C ₆ H ₁₃ SH	Cr^{3+} - Deep Red Co^{2+} - Red (with some yellow) Zn^{2+} - Yellow
Sn ⁴⁺ - Green	Co ³⁺ - No Change	Cr ³⁺ - Yellow circle surrounded by greenish blue circle
$\mathrm{Mn^{3+}}$ - Yellow $\mathrm{Cu^{2+}}$ - Dark Blue (faint) $\mathrm{Ag^{2+}}$ - Blue (very faint)	Fe ³⁺ - Dark Green Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - Blue Analyte: (C ₃ H ₇) ₂ S	Co ²⁺ - No Change Zn ²⁺ - Green
Sn ⁴⁺ - Dark Blue (faint) Mn ³⁺ - No Change Cu ²⁺ - Dark Blue (faint) Ag ²⁺ - Blue (very faint)	Co ³⁺ - Deep Green Fe ³⁺ - Dark Green Ru ²⁺ - Green 2H ⁺ (Free Base "FB") - Blue Analyte: Benzene	Cr^{3+} - Green Co^{2+} - Dark Green (very faint) Zn^{2+} - Green
$\mathrm{Sn^{4+}}$ - No Change $\mathrm{Mn^{3+}}$ - Yellow (some green) $\mathrm{Cu^{2+}}$ - No Change $\mathrm{Ag^{2+}}$ - No Change	Co ³⁺ - Green Fe ³⁺ - Dark Green Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") - Blue	Cr ³⁺ - Yellow (very faint) Co ²⁺ - No Change Zn ²⁺ - Dark Green

The degree of ligand softness (roughly their polarizability) increases from left to right, top to bottom as shown in FIG. 1. Each analyte is easily distinguished from the others, and there are family resemblances among chemically similar species (e.g., pyridine and n-hexylamine). Analyte distinction originates both in the metal-specific ligation affinities and in their specific, unique color changes upon ligation. Each analyte was delivered to the array as a nitrogen stream saturated with the analyte vapor at 20° C. (to ensure complete saturation, 30 min. exposures to vapor were used. Although these fingerprints were obtained by exposure to saturated vapors (thousands of ppm), unique patterns can be identified at much lower concentrations.

The metalloporphyrin array 16 has been used to quantify single analytes and to identify vapor mixtures. Because the images' color channel data (i.e., RGB values) vary linearly with porphyrin concentration, we were able to quantify single porphyrin responses to different analytes. Color channel data were collected for individual spots and plotted, for example, as the quantity $(R_{plt}-R_{spt})/(R_{plt})$, where R_{plt} was the red channel value for the initial silica surface and R_{spt} the average value for the spot. For example, Fe(TFPP)(Cl) responded linearly to octylamine between 0 and 1.5 ppm. 65 Other porphyrins showed linear response ranges that varied with ligand affinity (i.e., equilibrium constant).

EXAMPLE 3

The array of the present invention has demonstrated interpretable and reversible responses even to analyte mixtures of strong ligands, such as pyridines and phosphites, as is shown in FIG. 6. Color change patterns for the mixtures are distinct from either of the neat vapors. Good reversibility was demonstrated for this analyte pair as the vapor mixtures were cycled between the neat analyte extremes, as shown in FIG. 6, which shows the two component saturation responses to mixtures of 2-methylpyridine ("2MEPY") and trimethylphosphite ("TMP"). Vapor mixtures were obtained by mixing the analyte-saturated N₂ streams at variable flow ratios. A single plate was first exposed to pure trimethylphosphite vapor in N₂ (Scan A), followed by increasing mole fractions of 2-methylpyridine up to pure 2-methylpyridine vapor (Scan C), followed by decreasing mole fractions of 2-methylpyridine back to pure trimethylphosphite vapor. In both directions, scans were taken at the same mole fraction trimethylphosphite and showed excellent reversibility; scans at mole fractions at 67% trimethylphosphite $(\chi_{tmp}=0.67, Scans B \text{ and D})$ and of their difference map are shown (Scan E). Response curves for the individual porphyrins allow for quantification of the mixture composition. The colors of each dye upon exposure to the analytes TMP and 2MEPY are shown in FIG. 6 and are summarized in Table 5 below.

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TABLE 5

	Scan A, Analyte: Neat TMP	
Sn ⁴⁺ - Dark Blue Mn ³⁺ - Yellow with red center Cu ²⁺ - Dark Blue Ag ²⁺ - Green (very faint)	Co ³⁺ - Yellow Fe ³⁺ - Dark Green Ru ²⁺ - No Change 2H ⁺ (Free Base "FB") — Reddish Blue Scan B, Analyte: TMP,x _{TMP} = 0.0	Cr^{3+} - No Change Co^{2+} - Greenish Yellow Zr^{2+} - Blue
Sn ⁴⁺ - Blue Mn ³⁺ - Yellow and Green Cu ²⁺ - Dark Blue Ag ²⁺ - Greenish Blue	Co ³⁺ - Green Fe ³⁺ - Green and Yellow Ru ²⁺ - Purple (very faint) 2H ⁺ (Free Base "FB") — Reddish Blue Scan C, Analyte: Neat 2MEPY	Cr ³⁺ - Green (small dot) Co ²⁺ - Green with red center Zn ²⁺ - Blue
Sn ⁴⁺ - Blue Mn ³⁺ - Yellow and Green with Red center Cu ²⁺ - Dark Blue Ag ²⁺ - Green with some Blue	Co ³⁺ - Green Fe ³⁺ - Red with some Yellow Ru ²⁺ - Deep Blue 2H ⁺ (Free Base "FB") — Reddish Blue Scan D, Analyte: TMP,x _{TMP} = 0.	Cr^{3+} - No Change Co^{2+} - Green Zn^{2+} - Green with some Blue
Sn ⁴⁺ - Blue Mn ³⁺ - Yellow and Green Cu ²⁺ - Dark Blue Ag ²⁺ - Greenish Blue (very faint)	Co ³⁺ - Green Fe ³⁺ - Green and Yellow Ru ²⁺ - Purple (very faint) 2H ⁺ (Free Base "FB") — Reddish Blue Scan E	Cr ³⁺ - No Change Co ²⁺ - Green Zn ²⁺ - Blue
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - Blue (very faint) Ag ²⁺ - Blue (very faint)	Co ³⁺ - No Change Fe ³⁺ - No Change Ru ²⁺ - Blue (small dot) 2H ⁺ (Free Base "FB") - Green	Cr^{3+} - No Change Co^{2+} - No Change Zn^{2+} - No Change

EXAMPLE 4

In an effort to understand the origin of the color changes upon vapor exposure, diffuse reflectance spectra were obtained for single porphyrin spots before and after exposure to analyte vapors. Porphyrin solutions were spotted in 40 50 μL aliquots onto a plate and allowed to dry under vacuum at 50° C. Diffuse reflectance spectra of the plate were then taken using a UV-visible spectrophotometer equipped with an integrating sphere. Unique spectral shifts were observed upon analyte exposure, which correlated well with those seen from solution ligation. For example, Zn(TPP) exposure to ethanol and pyridine gave unique shifts which were very similar to those resulting from ligand exposure in solution. FIG. 7 shows a comparison of Zn(TPP) spectral shifts upon exposure to ethanol and pyridine (py) in methylene chloride solution (A) and on the reverse phase support (B). In both A and B, the bands correspond, from left to right, to Zn(TPP), Zn(TPP)(C₂H₅OH), and Zn(TPP)(py), respectively. Solution spectra (A) were collected using a Hitachi U-3300 spectrophotometer; Zn(TPP), C₂H₅OH, and py concentrations were approximately 2 μM , 170 mM, and 200 μM , respectively. Diffuse reflectance spectra (B) were obtained with an integrating sphere attachment before exposure to analytes, after exposure to ethanol vapor in N₂, and after exposure to pyridine vapor in N₂ for 30 min. each using the flow cell.

Improvement to Low Concentration Response

Color changes at levels as low as 460 ppb have been observed for octylamine vapor, albeit with slow response times due to the high surface area of the silica on the plate 65 18. The surface area of C2 plates is $\approx 350 \text{ m}^2/\text{gram}$. Removal of excess silica gel surrounding the porphyrin spots from the

plate 18 led to substantial improvements in response time for suppose the strace levels of octylamine. Because the high surface area of the reverse phase silica surface is primarily responsible for the increased response time, other means of solid support or film formation can be used to improve low concentration response.

Further, the present invention contemplates miniaturization of the array using small wells **60** (<1 mm), for example in glass, quartz, or polymers, to hold metalloporphyrin or other dyes as thin films, which are deposited as a solution, by liquid droplet dispersion (e.g., airbrush or inkjet), or deposited as a solution of polymer with metalloporphyrin.

These embodiments are depicted in FIGS. **8**, **9**, and **10**. FIG. **8** illustrates the interfacing of a microplate **60** into an assembly consisting of a CCD **70**, a microplate **72** and a light source **74**. FIG. **9** illustrates another embodiment of the present invention, and more particularly, a microwell porphyrin array wellplate **80** constructed from polydimethylsiloxane (PDMS). The colors of the dyes shown in FIG. **9** are summarized below in Table 6.

TABLE 6

	Sn ⁴⁺ - Dark Red Mn ³⁺ - Green Cu ²⁺ - Deep Red	Co ³⁺ - Dark Red Fe ³⁺ - Dark Red Ru ²⁺ - Dark Red	Cr ³⁺ - Dark Green Co ²⁺ - Yellowish Green Zn ²⁺ - Red with some Yellow
)	Ag ²⁺ - Red	2H ⁺ (Free Base "FB") — Red	

FIG. 10 demonstrates deposition of metalloporphyrin/polymer (polystyrene/dibutylphthalate) solutions upon a plate, which includes a series of micro-machined Teflon® posts 100 having the same basic position relative to each

other as shown in FIG. 2A and FIG. 2B. The colors for the dyes in the middle of FIG. 10 are summarized in Table 7 below.

TABLE 7

Sn ⁴⁺ - Yellow	Co ³⁺ - Orange	Cr ³⁺ - Yellow
Mn ³⁺ - Yellow	Fe ³⁺ - Orange	Co ²⁺ - Orange
Cu ²⁺ - Orange	Ru ²⁺ - Dark Yellow	Zn ²⁺ - Orange
Ag ²⁺ - Orange	2H ⁺ (Free Base "FB") - Red	

The colors for the dyes on the right hand side of FIG. 10 are summarized in Table 8 below.

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The low ppm levels of octylamine, an analyte of interest, were generated from temperature-regulated octylamine/dodecane solutions with the assumption of solution ideality. The dodecane acts as a diluent to lower the level of octylamine vapor pressure for the purposes of this demonstration of the invention.

EXAMPLE 6

FIG. 12 illustrates the immunity of the present invention to interference from water vapor. The hydrophobicity of the

TABLE 8

Sn ⁴⁺ - No Change	Co ³⁺ - Green	Cr ³⁺ - Red
Mn ³⁺ - Blue	Fe ³⁺ - Red	Co2+ - Red, Green, Blue, and
		Yellow
Cu ²⁺ - Green with some Blue	Ru ²⁺ - Blue (very faint)	Zn ²⁺ - Yellow with some Red
Ag ²⁺ - Green with some Blue	2H+ (Free Base "FB") - Green	
	with some Blue	

EXAMPLE 5

FIG. 11 shows the color profile changes from a microplate of the type shown in FIG. 10. The microplate, consisting of a minimized array of four metalloporphyrins, i.e., Sn(TPP) (Cl₂), Co(TPP)(Cl), Zn(TPP), Fe(TFPP)(Cl), clockwise from the upper left (where TFPP stands for 5,10,15,20-tetrakis(pentafluorophenyl)porphyrinate). The color profile changes are shown in FIG. 11 after exposure to low levels of n-octylamine, dodecanethiol ($C_{12}H_{25}$ SH), and trinbutylphosphine ($P(C_4H_9)_3$), each at 1.8 ppm, which is summarized in Table 9 below.

reverse phase support greatly any possible effects from varying water vapor in the atmosphere to be tested. For instance, as shown in FIG. 12, a color fingerprint generated from exposure of the array to n-hexylamine (0.86% in N₂) was identical to that for n-hexylamine spiked heavily with water vapor (1.2% H₂O, 0.48% hexylamine in N₂). See scans 120, 122 and 124. The ability to easily detect species in the presence of a large water background represents a substantial advantage over mass-sensitive sensing techniques or methodologies that employ polar polymers as part of the sensor array. The color patterns shown in FIG. 12 are summarized in Table 10 below.

TABLE 9

	Dyes on Teflon ®
Sn - Dark Yellow	Co - Red
Zn - Red	Fe - Orange with Red outline
Dyes	exposed to n-octylamine
Sn - No Change	Co - Green (very faint)
Zn - Red	Fe - Green
Dye	es exposed to C ₁₂ H ₂₅ SH
Sn - Red	Co - Green with some red, yellow and blue
	(faint)
Zn - Red with some green and yellow	Fe - Blue (very faint)
Dye	es exposed to P(C ₄ H ₉) ₃
Sn - No Change	Co - Yellow with red center and some red periphery
Zn - Green	Fe - Yellow with some Green and Blue

TABLE 10

Scan 120							
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - No Change Ag ²⁺ - No Change	Co^{3+} - Green Fe^{3+} - Red Ru^{2+} - No Change $2H^+$ (Free Base "FB") - Dark Blue	Cr^{3+} - Green Co^{2+} - No Change Zn^{2+} - Green					
Sn ⁴⁺ - No Change Mn ³⁺ - No Change Cu ²⁺ - No Change Ag ²⁺ - No Change	Co^{3+} - Green Fe^{3+} - Red Ru^{2+} - Green (small dot) $2H^+$ (Free Base "FB") - Dark Blue	Cr^{3+} - Green Co^{2+} - No Change Zn^{2+} - Green					
Sn ⁴⁺ - Bluish Circle Mn ³⁺ - Bluish Circle Cu ²⁺ - Bluish Circle Ag ²⁺ - Bluish Circle	Co ³⁺ - Bluish Circle Fe ³⁺ - Bluish Circle Ru ²⁺ - Bluish Circle 2H ⁺ (Free Base "FB") - Bluish Circle	Cr ³⁺ - Bluish Circle Co ²⁺ - Bluish Circle Zn ²⁺ - Bluish Circle					

Additional Features of the Preferred Embodiments of the Invention

Having demonstrated electronic differentiation, an important further goal is the shape-selective distinction of analytes 25 (e.g., n-hexylamine vs. cyclohexylamine). Functionalized metalloporphyrins that limit steric access to the metal ion are candidates for such differentiation. For instance, we have been able to control ligation of various nitrogenous ligands to dendrimer-metalloporphyrins and induce selectivities over a range of more than 10⁴. As an initial attempt toward shape-selective detection, we employed the slightly-hindered tetrakis(2,4,6-trimethoxyphenyl)porphyrins (TTMPP) in our sensing array. With these porphyrins, fingerprints for t-butylamine and n-butylamine showed subtle distinctions, as did those for cyclohexylamine and n-hexylamine. Using more hindered metalloporphyrins, it is contemplated that the present invention can provide greater visual differentiation. Such porphyrins include those whose periphery is decorated with dendrimer, siloxyl, phenyl, t-butyl and other bulky substituents, providing sterically constrained pockets on at least one face (and preferably both) of the porphyrin.

In a similar fashion, it is contemplated that the sensor plates of the present invention can be used for the detection of analytes in liquids or solutions, or solids. A device that detects an analyte in a liquid or solution or solid can be referred to as an artificial tongue. Proper choice of the metal complexes and the solid support must preclude their dissolution into the solution to be analyzed. It is preferred that the surface support repel any carrier solvent to promote the detection of trace analytes in solution; for example, for analysis of aqueous solutions, reverse phase silica has advantages as a support since it will not be wetted directly by water.

Alternative sensors in accordance with the present invention may include any other dyes or metal complexes with intense absorbance in the ultraviolet, visible, or near infrared spectra that show a color change upon exposure to analytes. These alternative sensors include, but are not limited to, a variety of macrocycles and non-macrocycles such as chlorins and chlorophylls, phthalocyanines and metallophthalocyanines, salen-type compounds and their metal complexes, or other metal-containing dyes.

The present invention can be used to detect a wide variety 65 of analytes regardless of physical form of the analytes. That is, the present invention can be used to detect any vapor

emitting substance, including liquid, solid, or gaseous forms, and even when mixed with other vapor emitting substances, such solution mixtures of substances.

The present invention can be used in combinatorial libraries of metalloporphyrins for shape selective detection of substrates where the substituents on the periphery of the macrocycle or the metal bound by the porphyrin are created and then physically dispersed in two dimensions by (partial) chromatographic or electrophoretic separation.

The present invention can be used with chiral substituents on the periphery of the macrocycle for identification of chiral substrates, including but not limited to drugs, natural products, blood or bodily fluid components.

The present invention can be used for analysis of biological entities based on the surface proteins, oligosacharides, antigens, etc., that interact with the metalloporphyrin array sensors of the present invention. Further, the sensors of the present invention can be used for specific recognition of individual species of bacteria or viruses.

The present invention can be used for analysis of nucleic acid sequences based on sequence specific the surface interactions with the metalloporphyrin array sensors. The sensors of the present invention can be used for specific recognition of individual sequences of nucleic acids. Substituents on the porphyrins that would be particularly useful in this regard are known DNA intercalating molecules and nucleic acid oligomers.

The present invention can be used with ordinary flat bed scanners, as well as portable miniaturized detectors, such as CCD detectors with microarrays of dyes such as metalloporphyrins.

The present invention can be used for improved sensitivity, automation of pattern recognition of liquids and solutions, and analysis of biological and biochemical samples.

Superstructure Bonded to the Periphery of the Porphyrin

The present invention includes modified porphyrins that have a super structure bonded to the periphery of the porphyrin. A super structure bonded to the periphery of the porphyrin in accordance with the present invention includes any additional structural element or chemical structure built at the edge of the porphyrin and bonded thereto.

The super structures can include any structural element or chemical structure characterized in having a certain selectivity. Those of skill in the art will recognize that the super structures of the present invention include structures that are

shape selective, polarity selective, inantio selective, regio selective, hydrogen bonding selective, and acid-base selective. This structures can include siloxyl-substituted substituents, nonsiloxyl-substituted substituents and nonsiloxyl-substituted substituents, including but not limited to aryl 5 substituents, alkyl substituents, and organic, organometallic, and inorganic functional group substituents.

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Superstructure Bis-Pocket Porphyrins

A number of modified porphyrins have been synthesized to mimic various aspects of the enzymatic functions of heme proteins, especially oxygen binding (myoglobin and hemoglobin) and substrate oxidation (cytochrome P-450). See Suslick, K. S.; Reinert, T. J. J. Chem. Ed. 1985, 62, 974; Collman, J. P.; Zhang, X.; Lee, V. J.; Uffelman, E. S.; Brauman, J. I. Science 1993,261, 1404; Collman, J. P.; 15 Zhang, X. in Comprehensive Supramolecular Chemistry; Atwood, J. L.; Davies, J. E. D.; MacNicol, D. D.; Vogtel, F. Eds.; Pergamon: New York, 1996; vol. 5, pp. 1-32; Suslick, K. S.; van Deusen-Jeffries, S. in Comprehensive Supramolecular Chemistry; Atwood, J. L.: Davies, J. E. D.: MacNi- 20 col, D. D.; Vogtel, F. Eds.; Pergamon: New York, 1996; vol. 5, pp. 141-170; Suslick, K. S. in Activation and Functionalization of Alkanes; Hill, C. L., ed.; Wiley & Sons: New York, 1989; pp. 219-241. The notable property of many heme proteins is their remarkable substrate selectivity; the 25 development of highly regioselective synthetic catalysts, however, is still at an early stage. Discrimination of one site on a molecule from another and distinguishing among many similar molecules presents a difficult and important challenge to both industrial and biological chemistry. See Met- 30 alloporphyrins in Catalytic Oxidations; Sheldon, R. A. Ed. Marcel Dekker: New York, 1994). Although the axial ligation properties of simple synthetic metalloporphyrins are well documented in literature, see Bampos, N.; Marvaud, V.; Sanders, J. K. M. Chem. Eur. J. 1998, 4, 325; Stibrany, R. 35 T.; Vasudevan, J.; Knapp, S.; Potenza, J. A.; Emge, T.; Schugar, H. J. J. Am. Chem. Soc. 1996, 118, 3980, size and shape control of ligation to peripherally modified metalloporphyrins has been largely unexplored, with few notable exceptions, where only limited selectivities have been 40 observed. See Bhyrappa, P.; Vaijayanthimala, G.; Suslick, K. S. J. Am. Chem. Soc. 1999, 121, 262; Imai, H.; Nakagawa, S.; Kyuno, E. J. Am. Chem. Soc. 1992, 114, 6719

The present invention includes the synthesis, characterization and remarkable shape-selective ligation of silylethermetalloporphyrin scaffolds derived from the reaction of 5,10,15,20-tetrakis(2',6'-dihydroxyphenyl)porphyrinatozine (II) with t-butyldimethylsilyl chloride, whereby the two faces of the Zn(II) porphyrin were protected with six, seven, or eight siloxyl groups. This results in a set of three 50 porphyrins of nearly similar electronics but with different steric encumbrance around central metal atom present in the porphyrin. Ligation to Zn by classes of different sized ligands reveal shape selectivities as large as 10^7 .

A family of siloxyl-substituted bis-pocket porphyrins 55 were prepared according to the scheme of FIG. 13. The abbreviations of the porhyrins that can be made in accordance with the scheme shown in FIG. 13 are as follows: Zn(TPP), 5,10,15,20-tetraphenylporphyrinatozinc(II);

Zn[(OH)₆PP], 5-phenyl-10,15,20-tris(2',6'-dihydroxyphe-60 nyl)porphyrinatozinc(II);

Zn[(OH)₈PP], 5,10,15,20-tetrakis(2',6'-dihydroxyphenyl) porphyrinatozinc(II);

Zn(Si₆PP), 5(phenyl)-10,15,20-trikis(2',6'-disilyloxyphenyl) porphyrinatozinc(II);

Zn(Si₇OHPP), 5,10,15-trikis(2',6'-disilyloxyphenyl)-20-(2'-hydroxy-6'-silyloxyphenyl)porphyrinatozinc(II);

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Zn(Si₈PP), 5,10,15,20-tetrakis(2',6'-disilyloxyphenyl)porphyrinatozinc(II). The synthesis of Zn[(OH)₆PP], Zn(Si₆PP), and Zn(Si₈PP) is detailed below. Zn[(OH)₆PP] and Zn[(OH)₈PP] were obtained (see Bhyrappa, P.; Vaijayanthimala, G.; Suslick, K. S. J. Am. Chem. Soc. 1999, 121, 262) from demethylation (see Momenteau, M.; Mispelter, J.; Loock, B.; Bisagni, E. J. Chem. Soc. Perkin Trans. 1, 1983, 189) of corresponding free base methoxy compounds followed by zinc(II) insertion. The methoxy porphyrins were synthesized by acid catalysed condensation of pyrrole with respective benzaldehydes following Lindsey procedures. See Lindsey, J. S.; Wagner, R. W. J. Org. Chem. 1989, 54, 828. Metalation was done in methanol with Zn(O₂CCH₃)₂. The t-butyldimethylsilyl groups were incorporated into the metalloporphyrin by stirring a DMF solution of hydroxyporphyrin complex with TBDMSiCl (i.e., t-butyldimethylsilyl chloride) in presence of imidazole. See Corey, E. J; Venkateswarlu, A. J. Am. Chem. Soc. 1972, 94, 6190. The octa (Zn(Si₈PP)), hepta (Zn(Si₇OHPP)), and hexa (Zn(Si₆PP)) silylether porphyrins were obtained from Zn[(OH)₈PP] and Zn[(OH)₆PP], respectively. The compounds were purified by silica gel column chromatography and fully characterized by UV-Visible, 1H-NMR, HPLC, and MALDI-TOF MS.

The size and shape selectivities of the binding sites of these bis-pocket Zn silylether porphyrins were probed using the axial ligation of various nitrogenous bases of different shapes and sizes in toluene at 25° C. Zn(II) porphyrins were chosen because, in solution, they generally bind only a single axial ligand. Successive addition of ligand to the porphyrin solutions caused a red-shift of the Soret band typical of coordination to zinc porphyrin complexes. There is no evidence from the electronic spectra of these porphyrins for significant distortions of the electronic structure of the porphyrin. The binding constants (K_{eq}) and binding composition (always 1:1) were evaluated using standard procedures. See Collman, J. P.; Brauman, J. I.; Doxsee, K. M.; Halbert, T. R.; Hayes, S. E.; Suslick, K. S. J. Am. Chem. Soc. 1978, 100, 2761; Suslick, K. S.; Fox, M. M.; Reinert, T. J. Am. Chem. Soc. 1984, 106,4522. The K_{eq} values of the silylether porphyrins with nitrogenous bases of different classes are compared with the sterically undemanding Zn(TPP) in FIGS. 14a, 14b, and 14c. It is worth noting the parallel between shape selectivity in these equilibrium measurements and prior kinetically-controlled epoxidation and hydroxylation. See Collman, J. P.; Zhang, X. in Comprehensive Supramolecular Chemistry; Atwood, J. L.; Davies, J. E. D.; MacNicol, D. D.; Vogtel, F. Eds.; Pergamon: New York, 1996; vol. 5, pp. 1-32; Suslick, K. S.; van Deusen-Jeffries, S. in Comprehensive Supramolecular Chemistry; Atwood, J. L.; Davies, J. E. D.; MacNicol, D. D.; Vogtel, F. Eds.; Pergamon: New York, 1996; vol. 5, pp. 141-170; Suslick, K. S. in Activation and Functionalization of Alkanes; Hill, C. L., ed.; Wiley & Sons: New York, 1989; pp. 219-241; Bhyrappa, P.; Young, J. K.; Moore, J. S.; Suslick, K. S. J. Am. Chem. Soc., 1996, 118, 5708-5711. Suslick, K. S.; Cook, B. R. J. Chem. Soc., Chem. Comm. 1987,200-202; Cook, B. R.; Reinert, T. J.; Suslick, K. S. J. Am. Chem. Soc. 1986,108,7281-7286; Suslick, K. S.; Cook, B. R.; Fox, M. M. J. Chem. Soc., Chem. Commun. 1985, 580-582. The selectivity for equilibrated ligation appears to be substantially larger than for irreversible oxidations of similarly shaped substrates.

The binding constants of silylether porphyrins are remarkably sensitive to the shape and size of the substrates relative to Zn(TPP). See FIGS. **14**a, **14**b, and **14**c. The binding

constants of different amines could be controlled over a range of 101 to 107 relative to Zn(TPP). It is believed that these selectivities originate from strong steric repulsions created by the methyl groups of the t-butyldimethylsiloxyl substituents. The steric congestion caused by these bulky silylether groups is pronounced even for linear amines and small cyclic amines (e.g., azetidine and pyrrolidine).

There are very large differences in K_{eq} for porphyrins having three versus four silvlether groups on each face (e.g., hexa- vs. octa-silylether porphyrins), as expected based on 10 obvious steric arguments (see FIGS. 14a, 14b, and 14c). Even between the hexa- over hepta-silylether porphyrins, however, there are still substantial differences in binding behavior. It is believed that this is probably due to doming of the macrocycle in the hexa- and hepta-silvlether porphy- 15 rins, which lessens the steric constraint relative to the octasilylether porphyrin. Such doming will be especially important in porphyrins whose two faces are not identical. The free hydroxy functionality of the hepta-silylether may play a role in binding of bi-functionalized ligands (e.g., free 20 amino acids); for the simple amines presented here, however, we have no evidence of any special effects.

These silylether porphyrins showed remarkable selectivities for normal, linear amines over their cyclic analogues. For a series of linear amines (n-propylamine through n-de- 25 cylamine), K_{eq} were very similar for each of the silylether porphyrins. In comparison, the relative K_{eq} for linear versus cyclic primary amines (FIG. **14***a*, n-butylamine vs. cyclohexylamine) were significantly different: $K_{eq}^{linear}/K_{eq}^{cyclic}$ ranges from 1 to 23 to 115 to >200 for Zn(TPP), $Zn(Si_6PP)$, 30 Zn(Si₇OHPP), and Zn(Si₈PP), respectively. The ability to discriminate between linear and cyclic compounds is thus established.

A series of cyclic 2° amines (FIG. 14b) demonstrate the remarkable size and shape selectivities of this family of 35 bis-pocket porphyrins. Whereas the binding constants to Zn(TPP) with those amines are virtually similar. In contrast, the K_{eq} values for silylether porphyrins strongly depend on the ring size and its peripheral substituents. The effect of these shape-selective binding sites is clear, even for compact 40 aromatic ligands with non-ortho methyl substituents (FIG.

The molecular structures of these silvlether porphyrins explains their ligation selectivity. The x-ray single crystal structure of Zn(Si₈PP) has been solved in the triclinic P1bar 45 space group. See Single crystal x-ray structure of Zn(Si8PP) shown in FIG. 15. As shown in FIG. 15, Zn(Si₆PP) (energy minimized molecular model) and Zn(Si₈PP) (single crystal x-ray structure) have dramatically different binding pockets. In the octasilylether porphyrin, the top access on both faces 50 of the porphyrin is very tightly controlled by the siloxyl pocket. In contrast, the metal center of the hexasilylether porphyrin is considerably more exposed for ligation.

FIG. 15 illustrates molecular models of Zn(Si₆PP) (left column) and $Zn(Si_8PP)$ (right column). The pairs of images 55 774 (m/z calcd. for $ZnC_{44}H_{28}O_6N_4=773$) in FAB-MS. from top to bottom are cylinder side-views, side-views, and top-views, respectively; space filling shown at 70% van der Waals radii; with the porphyrin carbon atoms shown in purple, oxygen atoms shown in red, silicon atoms in green, and Zn in dark red. The x-ray single crystal structure of 60 Zn(Si₈PP) is shown; for Zn(Si₆PP), an energy-minimized structure was obtained using Cerius 2 from MSI

In summary, a series of bis-pocket siloxyl metalloporphyrin complexes were prepared with sterically restrictive binding pockets on both faces of the macrocycle. Ligation to Zn 65 by various nitrogenous bases of different sizes and shapes were investigated. Shape selectivities as large as 10⁷ were

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found, compared to unhindered metalloporphyrins. Finetuning of ligation properties of these porphyrins was also possible using pockets of varying steric demands. The shape selectivities shown here rival or surpass those of any biological system.

EXAMPLES OF SYNTHESIS OF SUPER **STRUCRURES**

Synthesis of 5-phenyl-10,15,20-tris(2',6'-dihydroxyphenyl)-porphyrinatozinc(II), Zn[(OH)₆PP]:

The free base 5-phenyl-10,15,20-tris(2',6'-dimethoxyphenyl)-porphyrin was synthesized by Lewis acid catalyzed condensation of 2,6-dimethoxybezaldehyde and benzaldehyde with pyrrole (3:1:4 mole ratio) following the Lindsey procedure. See Lindsey, J. S.; Wagner, R. W. J. Org. Chem. 1989, 54, 828. The mixture of products thus formed was purified by silica gel column chromatography (if necessary, using CH₂Cl₂ as eluant). The isolated yield of the desired product was found to be 7%(wrt pyrrole used). The corresponding hydroxyporphyrins were obtained by demethylation with pyridine hydrochloride. See Momenteau, M.; Mispelter, J.; Loock, B.; Bisagni, E. J. Chem. Soc. Perkin Trans. 1, 1983, 189. After typical work-up known to those skilled in the art, the crude compound was purified by silica gel column chromatography using ethylacetate as eluant. The first fraction was Zn[(OH)₆PP], which was collected and the solvent was removed. The yield of the product was 90% (based on starting hydroxyporphryin). ¹H NMR of $H_2[(OH)_6PP]$ in acetone- d_6 (ppm): 8.96-8.79(m, 8H, b-pyrrole H), 8.24(m, 2H, o-H 5-Phenyl), 8.07 and 8.02(2s, 6H, —OH), 7.83(m, 3H, m,p-H 5-Phenyl), 7.50(t, 3H, p-H hydroxyphenyl), 6.90(d, 6H, m-H hydroxyphenyl), -2.69(s, 2H, imino-H). Elemental analysis, calcd. C₄₄H₃₀O₆N₄.H₂O: C=72.5, H=4.4 and N=7.7%. Found C=72.7, H=4.4 and N=7.4%. The compound showed molecular ion peak at 711 (m/z calcd. for $C_{44}H_{30}O_6N_4=710$) in FAB-MS.

The Zn derivative was obtain by stirring methanol solution of H₂[(OH)₆PP] with excess Zn(O₂CCH₃)₂2H₂O for 1 hour. Methanol was evaporated to dryness and the residue was dissolved in ethylacetate, washed with water, and the organic layer passed through anhyd. Na₂SO₄. The concentrated ethylacetate solution was passed through a silica gel column and the first band was collected as the desired product. The yield of the product was nearly quantitative. ¹H NMR of Zn(OH)₆PP in acetone-d₆ (ppm): 8.95-8.79(m, 8H, b-pyrrole H), 8.22(m, 2H, o-H 5-Phenyl), 7.79(m, 3H, m,p-H 5-Phenyl), 7.75 and 7.65(2s, 6H, —OH), 7.48(t, 3H,p-H hydroxyphenyl), 6.88(d, 6H, m-H hydroxyphenyl). Elemental analysis, calcd. for ZnC₄₄H₂₈O₆N₄.H₂O: C=66.7, H=3.8, N=7.1 and Zn=8.3%. Found C=66.4, H=3.8, N=6.7 and Zn=8.2%. The compound showed molecular ion peak at

Synthesis of 5-phenyl-10,15,20-tris(2',6'-disilyloxyphenyl)-porphyrinatozinc(II), Zn(Si₆PP):

The hexasilylether porphyrin was synthesized by stirring a DMF solution of 5-phenyl-10,15,20-tris(2',6'-dihydroxyphenyl)-porphyrinatozinc(II) (100 mg, 0.13 mmol) with t-butyldimethyl silylchloride (1.18 g, 7.8 mmol) in presence of imidazole (1.2 g, 17.9 mmol) at 60° C. for 24 h under nitrogen. After this period the reaction mixture was washed with water and extracted in CHCl3. The organic layer was dried over anhyd. Na₂SO₄. The crude reaction mixture was

loaded on a short silica gel column and eluted with mixture of CHCl₃/petether (1:1, v/v) to get rid of unreacted starting material and lower silylated products. The desired compound was further purified by running another silica gel column chromatography using mixture of CHCl₃/petether 5 (1:3, v/v) as eluant. The yield of the product was 60% based on starting hydroxyporphyrin.

 1 H NMR in chloroform-d (ppm): 8.94-8.82(m, 8H, b-pyrrole H), 8.20(m, 2H, o-H 5-Phenyl), 7.74(m, 3H, m,p-H 5-Phenyl), 7.49(t, 3H, p-H hydroxyphenyl), 6.91(t, 6H, 10 m —H hydroxyphenyl), -0.02 and -0.34(2s, 54H, t-butyl H), -0.43, -0.78 and -1.01(3s, 36H, methyl H). Elemental analysis, calcd. for $\rm ZnC_{80}H_{112}O_6N_4Si_6$: C=65.8, H=7.7, N=3.8, Si=11.5 and Zn=4.5%. Found C=65.5, H=7.7, N=3.8, Si=11.2 and Zn=4.4%. The low resolution MALDI-15 TOF mass spectrum showed molecular ion peak at 1457 (m/z calcd. for $\rm ZnC_{80}H_{112}O_6N_4Si_6$ =1458).

Synthesis of 5,10,15-tris(2',6'-disilyoxyphenyl)-20-(2'-hydr-oxy-6'-silyloxyphenyl)porphyrinatozinc(II), [Zn(Si₇OHPP)], and 5,10,15,20-tetrakis(2',6'-disily-loxyphenyl)porphy-rinato-zinc(II), [Zn(Si₈PP)]:

The synthesis of precursor porphyrin 5,10,15,20-tetrakis-(2',6'-dihydroxyphenyl)porphyrin and its Zn derivative was 25 accomplished as reported earlier. See Bhyrappa, P.; Vaijayanthimala, G.; Suslick, K. S. J. Am. Chem. Soc. 1999, 121, 262. The hepta-and octa-silylether porphyrins were synthesized by stirring DMF solution of 5,10,15,20-tetrakis(2',6'dihydroxyphenyl)porphyrinatozinc(II) (100 mg, 0.12 mmol) 30 with t-butyldimethyl silylchloride (1.45 g, 9.6 mmol) in presence of imidazole (1.50 g, 22.1 mmol) at 60° C. for 24 h under nitrogen. After usual work-up the mixture of crude products were loaded on a silica gel column and eluted with mixture of CHCl₃/pet. ether (1:1, v/v) to remove unreacted 35 starting material and lower silvlated products. The major product isolated from this column is a mixture of hepta- and octa-silylated porphyrins. The mixture thus obtained was further purified by another silica gel column chromatography using mixture of CHCl₃/pet. ether (1:3, v/v) as eluant. 40 The first two bands were isolated as octa- and heptasilylether porphyrin at 45% and 30% yield, respectively. Both the compounds were characterized by UV-Visible, ¹H NMR and MALDI-TOF spectroscopic techniques. The homogeneity of the sample was verified by HPLC.

For Zn(Si₇OHPP), 1 H NMR in chloroform-d (ppm): 8.91(m, 8H, b-pyrrole H), 7.50(m, 4H, p-H), 7.01-6.81(m, 8H, m-H), 0.11 to -0.03(12s, 105H, t-butyl) and methyl H). Elemental analysis, calcd. for ZnC₈₆H₁₂₆O₈N₄Si₇: C=64.3, H=7.8, N=3.5, Si=12.3 and Zn=4.1%. Found C=63.6, 50 H=8.1, N=3.5, Si=12.1 and Zn=3.9%. The low resolution MALDI-TOF mass spectrum showed molecular ion peak at 1604 (m/z calcd. for ZnC₈₆H₁₂₆O₈N₄Si₇=1604).

For Zn(Si₈PP), ¹H NMR in chloroform-d (ppm): 8.89(s, 8H, b-pyrrole H), 7.49(t, 4H, p-H), 6.92(d, 8H, m-H), 0.09(s, 55 72H, t-butyl H), -1.01(s, 48H, methyl H). Elemental analysis, calcd. for ZnC₉₂H₁₄₀O₈N₄Si₈: C=64.2, H=8.1, N=3.3, Si=13.1 and Zn=3.8%. Found C=63.5, H=8.4, N=3.3, Si=12.8 and Zn=4.0%. The low resolution MALDI-TOF mass spectrum showed molecular ion peak at 1719 (m/z 60 calcd. for ZnC₉₂H₁₄₀O₈N₄Si₈=1718).

ADDITIONAL FEATURES OF THE PREFERRED EMBEDIMENTS OF THE INVENTION

Having demonstrated electronic differentiation and shapeselective distinction of analytes that bind to metal ions in 24

metallodyes, an important further goal is the differentiation of analytes that do not bind or bind only weakly to metal ions. Such analytes include acidic compounds, such as carboxylic acids, and certain organic compounds lacking ligatable functionality, such as simple alkanes, arenes, some alkenes and alkynes (especially if sterically hindered), and molecules sterically hindered as to preclude effective ligation. One approach that has been developed to achieve this goal in accordance with the present invention is to include in the sensor array other chemoresponsive dyes, including pH sensitive dyes (i.e., pH indicator or acid-base indicator dyes that change color upon exposure to acids or bases), and/or solvatochromic dyes (i.e., dyes that change color depending upon the local polarity of their micro-environment).

It has been discovered that the addition of pH sensitive dyes and solvatochromic dyes to other arrays containing metalloporphyrins as described above expands the range of analytes to which the arrays are sensitive, improves sensitivities to some analytes, and increases the ability to discriminate between analytes.

The present invention includes an artificial nose comprising an array, the array comprising at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflectance response to an analyte wherein the first dye and the second dye are selected from the group consisting of chemoresponsive dyes, and the second dye is distinct from the first dye. In a preferred embodiment, the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phtahlocyanine, and salen and their metal complexes. In another preferred embodiment, the second dye is selected from the group of dyes consisting of acid-base indicator dyes and solvatochromic dyes.

The present invention includes a method of detecting an analyte comprising the steps of: (a) forming an array of at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflectance response to an analyte wherein the first dye and the second dye are selected from the group consisting of chemoresponsive dyes, and the second dye is distinct from the first dye; (b) subjecting the array to an analyte; (c) inspecting the array for a distinct and direct spectral absorbance or reflectance response; and (d) correlating the distinct and direct spectral response to the presence of the analyte. In a preferred method, the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phtahlocyanine, and salen and their metal complexes. In another preferred method, the second dye is selected from the group of acid-base indicator dyes and solvatochromic dyes.

The present invention includes an artificial tongue comprising an array, the array comprising at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflectance response to an analyte wherein the first dye and the second dye are selected from the group consisting of chemoresponsive dyes, and the second dye is distinct from the first dye. In a preferred embodiment, the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phtahlocyanine, and salen and their metal complexes. In another preferred embodiment, the second dye is selected from the group of dyes consisting of acid-base indicator dyes and solvatochromic dyes.

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Chemoresponsive dyes are those dyes that change color,

in either reflected or absorbed light, upon changes in their

chemical environment. Three general classes of chemore-

sponsive dyes are (1) Lewis acid/base dyes, (2) pH indicator

dyes, and (3) solvatochromic dyes.

Bromocresol Purple

Synonyms: 5',5" dibromo-m-cresolsulfonphthalein; Bromcresol Purple

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Molecular Formula: C₂₁H₁₆Br₂O₅S

Molecular Weight: 698.04

CAS: 115-40-2

pH = 5.2 yellow

=6.8 blue

Lewis acid/base dyes are those dyes that contain a Lewis acidic or basic center (where a Lewis acid is an electron pair acceptor and a Lewis base is an electron pair donor) and change color in response to changes in the Lewis acidity or basicity of their environment. A specific set of Lewis acid/base dyes includes dyes such as porphyrin, chlorin, chlorophyll, phtahlocyanine, and salen and their metal complexes.

pH indicator or acid-base indicator dyes are those that change color in response to changes in the proton acidity or basicity (also called Bronsted acidity or basicity) of their 15 environment. A specific set of pH indicator dyes include Chlorphenol Red, Bromocresol Green, Bromocresol Purple, Bromothymol Blue, Phenol Red, Thymol Blue, Cresol Red, Alizarin, Mordant Orange, Methyl Orange, Methyl Red, Congo Red, Victoria Blue B, Eosin Blue, Fat Brown B, 20 Benzopurpurin 4B, Phloxine B, Orange G, Metanil Yellow, Naphthol Green B, Methylene Blue, Safranine O, Methylene Violet 3RAX, Sudan Orange G, Morin Hydrate, Neutral Red, Disperse Orange 25, Rosolic Acid, Fat Brown RR, Cyanidin chloride, 3,6-Acridineamine, 6'-Butoxy-2,6-di- 25 amino-3,3'-azodipyridine, para-Rosaniline Base, Acridine Orange Base, Crystal Violet, and Malachite Green Carbinol Base.

Solvatochromic dyes are those that change color in response to changes in the general polarity of their environment, primarily through strong dipole-dipole interactions. To some extent, all dyes inherently are solvatochromic, although some are much more responsive than others. A specific set of highly responsive solvatochromic dyes include Reichardt's Dye and Nile Red.

It has been discovered that the following pH indicator (i.e., acid-base indicator) dyes and solvatochromic dyes are useful to expand the range of analytes to which the arrays containing metalloporphyrins are sensitive, improve sensitivities to some analytes, and increase the ability to discriminate between analytes. Those skilled in the art will recognize that other modifications and variations in the choice of such auxiliary dyes may be made in addition to those described and illustrated herein without departing from the spirit and scope of the present invention. Accordingly, the choice of dyes described and illustrated herein should be understood to be illustrative only and not limiting upon the scope of the present invention.

Chlorphenol Red

Molecular Formula: C₁₉H₁₂Cl₂O₅S

Molecular Weight: 423.28

CAS: 4430-20-0

Transition interval: pH 4.8 (yellow) to pH 6.7 (violet)

Bromocresol Green

 $Synonyms: \ \ 3',3'',5',5'' Tetrabromo-m-cresol sulfon phthale in;$

Bromcresol Green

Molecular Formula: $C_{21}H_{14}Br_4O_5S$

Molecular Weight: 698.04

CAS: 76-60-8

pH = 3.8 yellow

= 5.4 blue

Bromothymol Blue

5 Synonyms: 3',3"-Dibromothymolsulfonphthalein; Bromthy-

mol Blue

Molecular Formula: $C_{27}H_{28}Br_2O_5S$

Molecular Weight: 624.41

CAS: 76-59-5

pH = 6.0 yellow

= 7.6 blue

Phenol Red

Synonyms: Phenolsulfonphthalein Molecular Formula: C₁₉H₁₄O₅S

Molecular Weight: 354.38 CAS: 143-74-8

pH = 6.8 yellow

= 8.2 red

Thymol Blue

Synonyms: Thymolsulfonphthalein Molecular Formula: $C_{27}H_{30}O_5S$

Molecular Weight: 466.60

CAS: 76-61-9

pH = 1.2 red

= 2.8 yellow

= 8 yellow

=9.2 blue

Cresol Red

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55 Synonyms: Phenol, 4,4'-(1,1-dioxido-3H-2,1-benzoxathiol-

3-ylidene)bis[2-methyl-(9CI)] Molecular Formula: C₂₁H₁₈O₅S Molecular Weight: 382.43

CAS: 1733-12-6

⁶⁰ pH 1.8 (orange) to pH 2.0 (yellow); Transition interval

(alkaline): pH 7.0 (yellow) to pH 8.8 (violet)

Alizarin

Synonyms: 1,2-Dihydroxyanthraquinone, 9,10-Anthracene-

dione, 1,2-dihydroxy-(9CI) Molecular Formula: C₁₄H₈O₄ Molecular Weight: 240.22

pH = 5.5 yellow =6.8 red= 10.1 red= 12.1 violet

Mordant Orange 1

Synonyms: Alizarin Yellow R, C.I. 14030, 5-(4-nitropheny-

lazo)salicylic acid

Molecular Formula: C₁₃H₉N₃O₅ Molecular Weight: 287.23

CAS: 2243-76-7

Methyl Orange

Synonyms: 4-(p-[Dimethylamino]phenylazo)benzenesulfonic acid, sodium salt Acid Orange 52

Molecular Formula: C₁₄H₁₄N₃O₃SNa

Molecular Weight: 327.3 pH 3.0 (pink)—pH 4.4 (yellow)

Methyl Red

Synonyms: 4-Dimethylaminoazobenzene-2'carboxylicacid; 2-(4-Dimethylaminophenylazo)benzoic acid

Molecular Formula: $C_{15}H_{15}N_3O_2$ Molecular Weight: 269.31

CAS: 493-52-7

pH = 4.2 pink

= 6.2 yellow

Reichardt's Dye

Synonyms: [2,6-diphenyl-4-(2,4,6-triphenylpyridinio)phe-

nolate]

Molecular Formula: C41H29NO Molecular Weight: 551.69

CAS: 10081-39-7

Nile Red

9-(diethy-Synonyms: 5H-Benzo[a]phenoxazin-5-one, lamino)-(7CI, 8CI, 9CI), 9-(Diethylamino)-5H-benzo[a] phenoxazin-5-one; Nile Blue A oxazone

Molecular Formula: $C_{20}H_{18}N_2O_2$

Molecular Weight: 318.38

CAS: 7385-67-3

Congo Red

Molecular Formula: C₃₂H₂₄N₆O₆S₂.Na₂

Molecular Weight: 696.67

CAS: 573-58-0

pH range: blue 3.1-4.9 red

Victoria Blue B

Synonyms: Basic Blue 26, C.I. 44045 Molecular Formula: C₃₃H₃₂ClN₃ Molecular Weight: 506.10

CAS: 2580-56-5

Eosin Blue

Synonyms: (Acid Red 91, C.I. 45400, 4',5'-dibromo-2',7'dinitrofluorescein, disodium salt)

Molecular Formula: C₂₀H₈Br₂N₂O₉

Molecular Weight: 624.08

CAS: 548-24-3

Fat Brown B

Synonyms: Solvent red 3 Molecular Formula: C₁₈H₁₆N₂O₂

Molecular Weight: 292.3

CAS: 6535-42-8

Benzopurpurin 4B Synonyms: (C.I. 23500, Direct Red 2) Molecular Formula: C₃₄H₂₈N₆O₆S₂

Molecular Weight: 724.73

CAS: 992-59-6

pH range: violet 1.2-3.8 yellow

Phloxine B

Molecular Formula: $C_{20}H_4Br_4Cl_4O_5$

CAS: 18472-87-2

pH range: colorless 2.1-4.1 pink

Orange G

Synonyms: 1-Phenylazo-2-naphthol-6,8-disulfonic acid

disodium salt

Molecular Formula: $C_{16}H_{10}N_2Na_2O_7S_2$

Molecular Weight: 452.

pH range: yellow 11.5-14.0 pink

Metanil Yellow

Synonyms: (Acid Yellow 36, C.I. 13065)

Molecular Formula: C₁₈H₁₅N₃O₃S.Na

Molecular Weight: 375.38

CAS: 587-98-4

pH 1.5 (red) to pH 2.7 (yellow)

Naphthol Green B

Synonyms: (Acid Green 1, C.I. 10020)

Molecular Formula: C₁₀H₇NO₅S

Molecular Weight: 878.47

CAS: 19381-50-1

Methylene Blue

Synonyms: (Basic Blue 9, C.I. 52015)

40 Molecular Formula: $C_{16}H_{18}ClN_3S$

Molecular Weight: 373.90

CAS: 7220-79-3

Safranine O

Synonyms: (C.I. 50240, 3,7-diamino-2,8-dimethyl-5-phe-

nylphenazinium chloride) Molecular Formula: C₂₀H₁₉ClN₄ Molecular Weight: 350.85

CAS: 477-73-6

Methylene Violet 3RAX

[3-amino-7-(diethylamino)-5-phenylphenazinium chloride, C.I. 50206, N,N-diethylphenosafranine]

Molecular Formula: C22H23ClN4

Molecular Weight: 378.91

CAS: 4569-86-2

Sudan Orange G

Synonyms: [C.I. 11920, 4-(phenylazo)resorcinol, Solvent

Orange 1]

Molecular Formula: C₆H₅N=NC₆H₃-1,3-(OH)₂

Molecular Weight: 214.22

CAS: 2051-85-6

Morin Hydrate

65 Synonyms: (2',3,4',5,7-pentahydroxyflavone)

Molecular Formula: C₁₅H₁₀O₇

Molecular Weight: 302.24 Neutral Red

Molecular Formula: C₁₅H₁₆N₄.HCl Molecular Weight: 288.78

CAS: 553-24-2

pH = 6.8 red

= 8.0 yellow

Disperse Orange 25

Molecular Formula: $C_{17}H_{17}N_5$ 02

Molecular Weight: 323.36 CAS: 31482-56-1

D 1' 4 ' 1

Rosolic Acid

Molecular Formula: C₂₀H₁₆O₃ Molecular Weight: 290.32

CAS: 603-45-2

pH = 5.0 yellow

=6.8 pink

Fat Brown RR

Molecular Formula: C₁₆H₁₄N₄

Molecular Weight: 262.32

CAS: 6416-57-5

Cyanidin chloride

Molecular Formula: C15H11O6.Cl

Molecular Weight: 322.7

CAS: 528-58-5

3,6-Acridineamine

Molecular Formula: C₁₃H₁₁N₃

Molecular Weight: 209.25

CAS Number: 92-62-6

6'-Butoxy-2.6-diamino-3.3'-azodipyridine

Synonym: Azodipyridine

Molecular Formula: $C_{14}H_{18}N_6O$

Molecular Weight: 286.34

CAS: 617-19-6

para-Rosaniline Base

Synonym: Rosaniline

Molecular Formula: C₁₉H₁₉N₃O

Molecular Weight: 305.4

CAS: 25620-78-4

Acridine Orange Base

Molecular Formula: C₁₇H₁₉N₃

Molecular Weight: 265.36

CAS: 494-38-2

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Crystal Violet

Molecular Formula: $C_{25}H_{30}N_3.Cl$

Molecular Weight: 407.99

CAS: 548-62-9

pH = 0 yellow

= 1.8 blue

10

20

Malachite Green Carbinol Base

Molecular Formula: C₂₃H₂₆N₂O

Molecular Weight: 346.48

15 CAS: 510-13-4

pH = 0.2 yellow

= 1.8 blue-green

In a preferred embodiment, a low volatility liquid, e.g., a plasticizer, is used in an array of the present invention to keep the dyes in the array from crystallizing and to enhance then response of the array to an analyte. Examples of suitable low volatility liquids include, but are not limited to DOW CORNING 704 silicone diffusion pump fluid (Mo-

lecular Weight: 484.82, Density: 1.070, CAS Number: 3982-82-9), and diundecyl phthalate (Molecular Weight: 474.73, Density: 0.950, CAS Number: 3648-20-2, Formula: C₃₀H₅₀O₄, Boiling Point (° C.): 523 at 760 torr), dibutyl

 $C_{30}H_{50}O_4$, Boiling Point (° C.): 523 at 760 torr), dibutyl phthalate (Molecular Weight: 278.4, Density: 1.048, CAS Number: 84-74-2, Formula: $C_{16}H_{22}O_4$, Boiling Point (° C.): 340 at 760 torr), diisopropyl phthalate (Molecular Weight:

250.3, Density: 1.063, CAS Number: 605-45-8, Formula: C₁₄H₁₈O₄), squalane (Molecular Weight: 422.83, Density: 0.810, CAS Number: 111-01-3, Formula: C₃₀H₆₂, Boiling Point (° C.): 176 at 0.05 torr), triethylene glycol dimethyl ether (synonym: Trigluyme, Molecular Weight: 178.23,

Density: 0.986, CAS Number: 112-49-2, Formula: C₈H₁₈O₄, Boiling Point (° C.): 216 at 760 torr), and tetraethlyene glycol dimethyl ether (synonym: Tetraglyme, (Molecular Weight: 222.28, Density: 1.009, CAS Number: 143-24-8, Formula: C₁₀H₂₂O₅, Boiling Point (° C.): 275-276 at

⁴⁵ 760 torr).

FIG. 16 illustrates an array containing illustrative examples of porphyrin, metalloporphyrin, acid-base indicator, and solvatochromatic dyes. Typical sizes can range from 0.5 mm to 2 cm on a side. Linear, hexagonal, or rectangular arrays are also easily used. From left to right and top to bottom the identities and colors of the dyes used in the illustrative example of FIG. 16 are listed in Table 11 as follows (the exact colors depend, among other things, upon scanner settings).

TABLE 11

	(Summarizing the Dyes and Colors in FIG. 16, i.e., "Dye - Color")					
SnTPPCl ₂ - Light Green AgTPP -	CoTPP - Peach NiTPP -	CrTPPCl - Green InTPPCL -	MnTPPCl - Green IrTPPCl -	FeTPPCl - Light Brownish Green ZnTPP - Salmon	CuTPP - Salmon FeTFPPCl -	
Salmon	Pink	Tan	Pink		Olive	
ZnSi ₆ PP -	ZnSi ₇ OHPP -	ZnSi ₈ PP -	H ₂ TPP -	$\mathrm{H}_2\mathrm{FPP}$ - Light	Alizarin basic -	
Pink	Deep Pink	Pink	Carmel	Brown	Violet	
Me Red -	BCP - Dark	BCPbasic -	BTB - Dark	BTB basic - Blue	Ph Red basic -	
Orange	Green	Blue	Yellow		Lavender	

TABLE 11-continued

	(Summarizing	the Dyes and C	Colors in FIG.	16, i.e., "Dye - Col	or")
Nile Red - Violet	BCG - Blue	BCG basic - Blue	CresRed - Brownish Purple	CresRed basic - Purple	CP Red - Purple
R Dye - Light Blue	TB - Yellow	TB basic - Greenish Gray	MeOr - Yellow	MeOr basic - Orangish Brown	CP Red basic - Bluish Purple

where TPP=5,10,15,20-tetraphenylporphyrinate(-2);

Zn (Si₆PP)=5(phenyl)-10,15,29-trikis(2',6'-disilyloxyphenyl)porphyrinatozinc(II);

Zn(Si₇OHPP)=5,10,15trikis(2',6'-disilyloxyphenyl)-20-(2'-hydroxy-6'-silyloxyphenyl)porphyrinatozinc(II);

Zn(Si₈PP)=5,10,15,20-tetrakis(2',6'-disilyloxyphenyl)porphyrinatozinc(II);

Me Red=Methyl Red;

BCP=Bromocresol Purple;

BTB=Bromothymol Blue;

Ph Red=Phenol Red;

BCG=Bromocresol Green;

CresRed=Cresol Red;

CP Red=Chlorophenol Red;

R Dye=Reichardt's Dye;

TB=Thymol Blue;

MeOr=Methyl Orange; and

basic indicates the addition of KOH until the color of the basic form of the indicator dye was observed.

Note: DOW CORNING 704 silicone diffusion pump fluid (Molecular Weight: 484.82, Density: 1.070, CAS Number: 3982-82-9) was added to all porphyrin solutions: 40 µl/ml.

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FIG. 17 illustrates the response of the array described in FIG. 16 to acid vapors, specifically formic acid, acetic acid, iso-valeric acid, and 3-methyl-2-hexenoic acid. As shown in FIG. 17 and summarized in Table 12 below, the color changes of each dye in response to a particular analyte are shown as color difference maps, as follows (the exact colors depend, among others things, upon scanner settings). The color changes are derived simply by comparing the before exposure and after exposure colors and subtracting the two images (i.e., the absolute value of the difference of the red values becomes the new red value in the color difference map; etc. for green values and blue values). If there is no change in the red, green, and blue color values of a dye in the after-exposure image, then the color difference map will show black (i.e., red value=green value=blue value=0).

TABLE 12

	(Summarizing th		or Changes in Fl Map Color")	IG. 17, i.e. "Dye -	Difference
		(Analyte:	Formic Acid 14	O ppb)	
${\rm SnTPPCl}_2 -$	CoTPP -	CrTPPCl	MnTPPCl	FeTPPCl -	CuTPP -
Black	Black (no	Black	Black	Faint Blue	Black (no
(no	change)	(no	(no	Periphery	change)
change)		change)	change)		
AgTPP -	NiTPP -	InTPPCL -	IrTPPCl -	ZnTPP -	FeTFPPCl -
Black (no	Black (no	Black	Black (no	Black (no	Black (no
change)	change)	(no	change)	change)	change)
7 a: pp	Z a: outp	change)	II EDD	II EDD	
ZnSi ₆ PP -	ZnSi ₇ OHPP -	ZnSi ₈ PP -	H ₂ TPP -	H ₂ FPP -	Alizarin basic -
Black (no	Black (no	Black (no	Black (no	Black (no	Dark Blue
change)	change)	change)	change)	change)	mi m i i i
Me Red -	BCP -	BCP basic -	BTB -	BTB basic -	Ph Red basic -
Black (no	Yellow	White	Black (no	Red	Green
change)			change)	Periphery	
				w/Yellow Center	
Nile Red -	BCG -	BCG	CresRed -	CresRed	CP Red -
Black (no	Black (no	basic -	Black (no	basic - Light	Black (no
change)	change)	Dark	change)	Green	change)
change)	change)	Purple	change)	Green	change)
R Dye -	TB - Black	TB basic -	MeOr -	MeOr basic -	CP Red basic -
Black (no	(no change)	Black (no	Green and	Dark	Yellow
change)	(no change)	change)	Purple	Purple	Periphery and
change)		change)	Turpic	Tuple	Purple center
		(Analyte:	Formic Acid 21	(dag (ruipie center
				· PP·	
SnTPPCl ₂ -	CoTPP -	CrTPPC1 -	MnTPPCl	FeTPPC1 -	CuTPP - Black
Black	Black (no	Black (no	Black (no	Black (no	(no change)
(no	change)	change)	change)	change)	
change)					
AgTPP -	NiTPP -	InTPPCL -	IrTPPC1 -	ZnTPP - Black	FeTFPPC1 -
Black (no	Black (no	Black (no	Black (no	(no change)	Black (no
change)	change)	change)	change)		change)

TABLE 12-continued

		11.122	o iz commu		
	(Summarizing th		r Changes in FIG Map Color")	ł. 17, i.e. "Dye - 1	Difference
ZnSi ₆ PP - Black (no change)	ZnSi ₇ OH PP - Black (no	ZnSi ₈ PP - Black (no change)	H ₂ TPP - Black (no change)	H ₂ FPP - Black (no change)	Alizarin basic - Black (no change)
Me Red - Black (no change)	change) BCP - Red	BCP basic - Yellow Periphery and Red Center	BTB - Black (no change)	BTB basic - Red	Ph Red basic - Green
Nile Red - Black (no change)	BCG - Black (no change)	BCG basic - Red periphery	CresRed - Black (no change)	CresRed basic - Green	CP Red - Black (no change)
R Dye - Black (no change)	TB - Black (no change)	TB basic - Black (no change)	MeOr - Black (no change)	MeOr basic - Black (no change)	CP Red basic - Yellow Periphery and
0 /	,		Formic Acid 340	0 /	Purple Center
a mppot	c mpp	c mpp.ci	M. Empor		c mpp pt t
SnTPPCl ₂ - Black (no change)	CoTPP - Black (no change)	CrTPPCl - Black (no change)	MnTPPCl - Black (no change)	FeTPPCl - Black (no change)	CuTPP - Black (no change)
AgTPP - Black (no	NiTPP - Black (no	InTPPCL - Black (no	IrTPPCl - Black (no	ZnTPP - Black (no	FeTFPPCl - Black (no change)
change) ZnSi ₆ PP - Black (no change)	change) ZnSi ₇ OHPP Black (no change)	change) ZnSi ₈ PP - Black (no change)	change) H ₂ TPP - Black (no change)	change) H ₂ FPP - Black (no change)	Alizarin basic - Green and Purple
Me Red - Black (no change)	BCP - Yellow	BCP basic - White	BTB - Black (no change)	BTB basic - Yellow	Ph Red basic - Green
Nile Red - Black (no change)	BCG - Red	BCG basic - Red and Purple	CresRed - Black (no change)	CresRed basic - Light Green	CP Red - Green
R Dye - Black (no change)	TB - Black (no change)	TB basic - Black (no change)	MeOr - Blue	MeOr basic - Purple	CP Red basic - White
0 /		0 /	Formic Acid 680	ppb)	
SnTPPCl ₂ - Black (no	CoTPP - Black (no change)	CrTPPCl - Black (no change)	MnTPPCl Black (no change)	FeTPPCl - Black (no change)	CuTPP - Black (no change)
change) AgTPP - Black (no	NiTPP - Black (no	InTPPCL - Black (no	IrTPPCl - Black (no	ZnTPP - Black (no	FeTFPPCl - Black (no change)
change) ZnSi ₆ PP -	change) ZnSi ₇ OHPP	change) ZnSi ₈ PP -	change) H ₂ TPP -	change) H ₂ FPP -	Alizarin basic -
Black (no change) Me Red -	Black (no change) BCP -	Black (no change) BCP basic -	Black (no change) BTB -	Black (no change) BTB basic -	Green and Purple Ph Red basic -
Black (no change)	Yellow	White	Black (no change)	Red Periphery and Yellow Center	Green
Nile Red - Black (no	BCG Red and Purple	BCG basic - Red and Purple	CresRed - Black (no change)	CresRed basic - Green	CP Red - Black (no change)
change) R Dye - Black (no change)	TB - Black (no change)	TB basic - Black (no change)	MeOr - Light blue	MeOr basic - Purple	CP Red basic - White
		-	Acetic Acid 170 p	opb)	
SnTPPCl ₂ - Black (no	CoTPP - Black (no change)	CrTPPCl - Black (no change)	MnTPPCl Black (no change)	FeTPPCl - Black (no change)	CuTPP - Black (no change)
change) AgTPP - Black (no	NiTPP - Black (no change)	InTPPCL - Black (no	IrTPPCl - Black (no	ZnTPP - Black (no change)	FeTFPPCl - Black (no
change) ZnSi ₆ PP - Black (no change) Me Red -	ZnSi ₇ OHPP - Black (no change) BCP - Red	change) ZnSi ₈ PP - Black (no change) BCP basic -	change) H ₂ TPP - Black (no change) BTB -	H ₂ FPP - Black (no change) BTB basic -	change) Alizarin basic - Black (no change) Ph Red basic -
Black (no change)	DCI - KCU	Orange	Black (no change)	Red	Black (no change)

TABLE 12-continued

	(Summarizing the	e Dves and Colo	r Changes in FIG	. 17, i.e. "Dye - 1	Difference
	(Sammarzing U	-	Map Color")	. 17, i.e. Dyc - 1	5.110101100
Nile Red -	BCG - Purple	BCG basic -	CresRed -	CresRed basic -	
Black (no	and Orange	Purple	Black (no	Black (no	Black (no
change)	mp pl l /	Orange	change)	change)	change)
R Dye - Black (no	TB - Black (no	TB basic - Black (no	MeOr -	MeOr basic -	CP Red
change)	change)	change)	Black (no change)	Black (no change)	basic - Black (no change)
change)			Acetic Acid 250 p		(no change)
SnTPPCl ₂ -	CoTPP -	CrTPPCl -	MnTPPCl -	FeTPPC1 -	CuTPP - Black (no
Black	Black (no	Black (no	Black (no	Black (no	change)
(no change)	change)	change)	change)	change)	
AgTPP -	NiTPP -	InTPPCL -	IrTPPCl -	ZnTPP -	FeTFPPCl - Black
Black (no	Black (no	Black (no	Black (no	Black (no	(no change)
change) ZnSi ₆ PP -	change) ZnSi ₇ OHPP	change) ZnSi ₈ PP -	change) H ₂ TPP -	change) H ₂ FPP -	Alizarin basic -
Black (no	Black (no	Black (no	Black (no	Black (no	Black (no change)
change)	change)	change)	change)	change)	Diama (in timingt)
Me Red -	BCP -	BCP basic -	BTB -	BTB basic -	Ph Red basic -
Black (no	Yellow with	Red	Black (no	Red	Green
change)	Red Center		change)		
Nile Red -	BCG -	BCG basic -	CresRed -	CresRed	CP Red - Black
Black (no	Orange	Red and	Black (no	basic -	(no change)
change)	TED DI I	Purple	change)	Black (no change)	CD D III '
R Dye - Black (no	TB - Black	TB basic - Black (no	MeOr - Black (no	MeOr basic - Black (no	CP Red basic - White
change)	(no change)	change)	change)	change)	winte
change)			Acetic Acid 340 p		
SnTPPCl ₂ -	CoTPP -	CrTPPCl -	MnTPPCl -	FeTPPC1 -	CuTPP - Black (no
Black	Black (no	Black (no	Black (no	Black (no	change)
(no change)	change)	change)	change)	change)	
AgTPP -	NiTPP -	InTPPCL -	IrTPPCl -	ZnTPP -	FeTFPPCl - Black
Black (no	Black (no	Black (no	Black (no	Black (no	(no change)
change) ZnSi ₆ PP -	change) ZnSi ₇ OHPP	change) ZnSi ₈ PP -	change) H ₂ TPP -	change) H ₂ FPP -	Alizarin basic -
Black (no	Black (no	Black (no	Black (no	Black (no	Black (no change)
change)	change)	change)	change)	change)	5.00 (av 1.00 gr)
Me Red -	BCP -	BCP basic -	BTB -	BTB basic -	Ph Red basic -
Black (no change)	Yellow	Yellow	Black (no change)	Ornage	Green
Nile Red -	BCG -	BCG basic -	CresRed -	CresRed	CP Red - Black
Black (no	Faint	Purple	Black (no	basic -	(no change)
change)	Orange and Purple	•	change)	Green	, 0,
R Dye -	TB - Black	TB basic -	MeOr -	MeOr basic -	CP Red basic -
Black (no	(no change)	Black (no	Black (no	Black (no	White
change)		change) (Analyte: A	change) Acetic Acid 650 p	change)	
SnTPPCl 2 -	CoTPP -	CrTPPCl -	MnTPPCl - n	FeTPPC1 -	CuTPP - Black (no
Black	Black (no	Black (no	Black (no	Black (no	change)
(no change)	change)	change)	change)	change)	change)
AgTPP -	NiTPP -	InTPPCL -	IrTPPCl -	ZnTPP -	FeTFPPCl - Black
Black (no	Black (no	Black (no	Black (no	Black (no	(no change)
change)	change)	change)	change)	change)	
ZnSi ₆ PP -	ZnSi ₇ OHPP	ZnSi ₈ PP -	H_2TPP -	H ₂ FPP -	Alizarin basic -
Black (no	Black (no	Black (no	Black (no	Black (no	Faint Green
change)	change)	change)	change)	change)	Dh Dad basis
Me Red - Black (no	BCP - Yellow and	BCP basic - Faint	BTB - Orange	BTB basic - Yellow	Ph Red basic - Green
change)	Orange)	Yellow	Orange	20110 **	CICCH
Nile Red -	BCG -	BCG basic -	CresRed -	CresRed	CP Red - Faint
Black (no	Black (no	Purple	Black (no	basic -	Green
change)	change)		change)	White	
R Dye -	TB - Black	TB basic -	MeOr -	MeOr basic -	CP Red basic -
Black (no	(no change)	Black (no	Black (no	Green	White
change)		change)	change)		

TABLE 12-continued

(Summarizing the Dyes and Color Changes in FIG. 17, i.e. "Dye - Difference Map Color")

	`	1	Map Color")	,	
		(Analyte: Is	o-Valeric Acid 28	30 ppb)	
SnTPPCl ₂ -	CoTPP -	CrTPPCl -	MnTPPCl -	FeTPPCl -	CuTPP - Black (no
Black	Black (no	Black (no	Black (no	Black (no	change)
(no	change)	change)	change)	change)	
change)	NETDD	In TRRCI	I-TDDCI	ZwTDD	FeTFPPC - Black
AgTPP - Black (no	NiTPP - Black (no	InTPPCL - Black (no	IrTPPCl - Black (no	ZnTPP - Black (no	(no change)
change)	change)	change)	change)	change)	(no entange)
ZnSi ₆ PP -	ZnSi ₇ OHPP	ZnSi ₈ PP -	H_2TPP -	H ₂ FPP -	Alizarin basic -
Black (no	Black (no	Black (no change)	Black (no	Black (no	Black (no change)
change) Me Red -	change) BCP - Red	BCP basic -	change) BTB -	change) BTB basic -	Ph Red basic -
Black (no	Black (no	Faint Red	Black (no	Orange	Orange
change)	change)		change)		
Nile Red - Black (no	BCG - Faint Purple	BCG basic - Red	CresRed - Black (no	CresRed basic - Dark	CP Red - Black (no change)
change)	Periphery	Periphery	change)	Green	(no entange)
R Dye -	TB - Red	TB basic -	MeOr -	MeOr basic -	CP Red basic -
Black (no	and Purple	Red	Green	Green	Green Periphery
change)	Periphery	Periphery (Analyte:	Center Iso-Valeric 420	Periphery pph)	
		(1 11111) (2.	Do valene 120	PPC)	
SnTPPCl ₂ -	CoTPP -	CrTPPCl -	MnTPPCl -	FeTPPC1 -	CuTPP - Black (no
Black (no	Black (no change)	Black (no change)	Black (no change)	Black (no change)	change)
change)	change)	change)	change)	change)	
AgTPP -	NiTPP -	InTPPCL -	IrTPPC1 -	ZnTPP -	FeTFPPCl - Black
Black (no	Black (no	Black (no	Black (no	Black (no	(no change)
change) ZnSi ₆ PP -	change) ZnSi ₇ OHPP	change) ZnSi ₈ PP -	change) H ₂ TPP -	change) H ₂ FPP -	Alizarin basic -
Black (no	Black (no	Black (no	Black (no	Black (no	Black (no change)
change)	change)	change)	change)	change)	
Me Red -	BCP - Red	BCP basic - Faint	BTB -	BTB basic -	Ph Red basic -
Black (no change)		Green and	Black (no change)	Orange and Yellow	Faint Orange and Green
8-7		orange			
Nile Red -	BCG -	BCG basic -	CresRed -	CresRed	CP Red - Black
Black (no change)	Orange	Orange Periphery	Black (no change	basic - Green	(no change)
R Dye -	TB - Black	TB basic -	MeOr -	MeOr basic -	CP Red basic -
Black (no	(no change)	Black (no	Green	Green	Green
change)		change)	771 ' 1 '100	-0 1)	
		(Analyte: 1s	o-Valeric Acid 85	50 ppb)	
$SnTPPCl_2$ -	CoTPP -	CrTPPCl -	MnTPPCl -	FeTPPC1 -	CuTPP - Black
Faint	Faint Purple	Faint	Faint Purple	Faint Purple	(no change)
blue	a rimpo	Purple		g	n mennou nu u
AgTPP - Faint Blue	NiTPP - Black (no	InTPPCL - Faint Pink	IrTPPC1 - Black (no	ZnTPP - Black (no	FeTFPPCl - Black (no change)
raint blue	change)	ramit rink	change)	change)	(no change)
ZnSi ₆ PP -	ZnSi ₇ OHPP	ZnSi ₈ PP -	H ₂ TPP -	H ₂ FPP -	Alizarin basic -
Faint Blue	Faint Blue	Black (no	Faint Blue	Black (no	Black (no change)
		change)		change)	
Me Red - Black (no	BCP - White and Red	BCP basic - Yellow	BTB - Blue and Red	BTB basic - Red and	Ph Red basic - Yellow and Red
change)	and Red	and Red	and Red	Yellow	renow and Red
Nile Red -	BCG - White,	BCG basic -	CresRed -	CresRed	CP Red - Faint
Black (no	Red and Blue	White	Purple	basic -	Orange
change)		and Red	Periphery	Light Green	
R Dye - Faint Red	TB - Light Blue	TB basic -	MeOr - Green and	MeOr basic - Light	CP Red basic - Light Green
raint Red	Periphery	Purple Periphery	Blue	Green	Light Green
	1 empirery	and Red	21110	314411	
		Center			
		(Analyte: Isc	-Valeric Acid 17	00 ppb)	
SnTPPCl ₂ -	CoTPP -	CrTPPCl -	MnTPPCl -	FeTPPC1 -	CuTPP - Black (no
Black	Black (no	Black (no	Black (no	Black (no	change)
(no	change)	change)	change)	change)	- /
change)	a rimpo		T 5770 - 1		n anna ci
AgTPP -	NiTPP -	InTPPCL -	IrTPPCl -	ZnTPP -	FeTFPPCl - Black
Black (no change)	Black (no change)	Black (no change)	Black (no change)	Black (no change)	(no change)
Jimige)	J. J	-1101150)	Junge)	ommige)	

TABLE 12-continued

	(Summarizing t		or Changes in FI Map Color")	G. 17, i.e. "Dye -	Difference
ZnSi ₆ PP -	ZnSi ₇ OHPP	ZnSi ₈ PP -	H ₂ TPP -	H ₂ FPP -	Alizarin basic -
Black (no	Black (no	Black (no	Black (no	Black (no	Faint Purple
change)	change)	change)	change)	change)	
Me Red -	BCP - Red	BCP basic -	BTB -	BTB basic -	Ph Red basic -
Black (no change)		White	Black (no change)	White	White and Purple
Nile Red -	BCG - Red	BCG basic -	CresRed -	CresRed	CP Red - Black
Black	and Purple	White,	Black (no	basic -	(no change)
(no change)		Red, and Purple	change)	White	
R Dye -	TB - Black	TB basic -	MeOr -	MeOr basic -	CP Red basic -
Black (no	(no change)	Faint Red	Black (no	Faint	Green
change)	(0 /		change)	Green	
		(Analyte: 3-Met	hyl-2-hexenoic A	cid 12 ppb)	
SnTPPCl ₂ -	CoTPP -	CrTPPCl -	MnTPPCl -	FeTPPCl -	CuTPP - Black (no
Black	Black (no	Black (no	Black (no	Black (no	change)
(no	change)	change)	change)	change)	
change)					
AgTPP -	NiTPP -	InTPPCL -	IrTPPCl -	ZnTPP -	FeTFPPCl - Black
Black (no	Black (no	Black (no	Black (no	Black (no	(no change)
change)	change)	change)	change)	change)	
ZnSi ₆ PP -	ZnSi ₇ OHPP	ZnSi ₈ PP -	H ₂ TPP -	H ₂ FPP -	Alizarin basic -
Black (no change)	Black (no change)	Black (no change)	Black (no change)	Black (no change)	Black (no change)
Me Red -	BCP - Faint	BCP basic -	BTB -	BTB basic -	Ph Red basic -
Black (no	Purple	White	Black (no	Red	Purple and Green
change)	ruspic	and Purple	change)	Red	raipie and Green
Nile Red -	BCG -	BCG basic -	CresRed -	CresRed	CP Red - Black
Black (no	Faint Red	Faint	Black (no	basic - Light	(no change)
change)	and Purple	White and	change)	Blue and	
0 /		Purple	<i>U</i> /	Green	
R Dye -	TB - Black	TB basic -	MeOr -	MeOr basic -	CP Red basic -
Black (no	(no change)	Black (no	Black (no	Blue and	Green
change)		change)	change)	Green	

FIG. 18 illustrates a preferred array containing illustrative examples of porphyrin, metalloporphyrin, acid-base indicator, and solvatochromatic dyes. Typical sizes of the array can range from 0.5 mm to 2 cm on a side. Linear, hexagonal or rectangular arrays are also easily used. From left to right and

top to bottom the identities and colors of the dyes used in the illustrative example of FIG. 18 are listed in Table 13 as follows (the exact colors depend, among other things, upon scanner setting).

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TABLE 13

	(Summarizing the Dyes and Colors in FIG. 18, i.e., "Dye - Color")						
SnTPPCl ₂ - Light Green	CoTPP - Tan	CrTPPCl - Green with Dark Green Center	MnTPPCl - Green	FeTPPCl - Light Green	CuTPP - Light Pink		
$Zn(C_3F_7)_4P$ - Gray $ZnSi_6PP$ - Pink	ZnF ₂ PP - Light Pink ZnSi ₇ OHPP - Pink	InTPPCI - Reddish Beige ZnSi ₈ PP - Light Pink	ZnTMP - Pink H ₂ TPP - Light Reddish Beige	ZnTPP - Salmon H ₂ FPP - Greenish Yellow	FeTFPPCl - Beige Neutral Red Pink with Brown Center		
Methyl Red - Orange	Disperse Orange 25 - Pinkish Orange	Rosolic Acid - Red	Fat Brown RR - Dark	Cyanidin Chloride - Reddish Brown	Metanil Yellow - Light Yellow		
Nile Red -	Mordant	3,6-Acridineamine	Bromocresol	Azodipyridine - Yellow	Rosaniline -		
Light Purple	Orange 1 - Light Yellow	Yellow	Green - Dark Yellow		Pink		
Reichardt's Dye - Teal	Acridine Orange Base - Yellow	Crystal Violet - Dark Blue	Thymol Blue -Purple	Congo Red - Dark Red	Malachite Green Carbinol base - Light Blue		

Note:

DOW CORNING 704 silicone diffusion pump fluid (Molecular Weight: 484.82, Density: 1.070, CAS Number: 3982-82-9) was added to all porphyrin solutions: 40 µl/ml.

where

SnTPPCl₂ is 5,10,15,20-Tetraphenyl-21H,23H-porphine Tin

(W) Dichloride

Molecular Formula: C44H28SnCl2N4

Molecular Weight: 802 CAS: 26334-85-0;

CoTPP 5,10,15,20-Tetraphenyl-21H,23H-porphine is

Cobalt(II)

Molecular Formula: C44H28CoN4

Molecular Weight: 671 CAS: 14172-90-8:

CrTPPCl is 5,10,15,20-Tetraphenyl-21H,23H-porphine

Chromium(III) Chloride

Molecular Formula: C44H28CrClN4

Molecular Weight: 700 CAS: 28110-70-5;

MnTPPCl is 5,10,15,20-Tetraphenyl-21H,23H-porphine 20

Manganese(III) Chloride

Molecular Formula: C44H28ClMnN4

Molecular Weight: 703 CAS: 32195-55-4;

FeTPPC1 is 5,10,15,20-Tetraphenyl-21H,23H-porphine Iron ²⁵

(III) Chloride

Molecular Formula: C44H28ClFeN4

Molecular Weight: 704 CAS: 16456-81-8;

CuTPP is 5,10,15,20-Tetraphenyl-21H,23H-porphine Cop-

Molecular Formula: C44H28CuN4

Molecular Weight: 676 CAS: 14172-91-9;

Zn(C₃F₇)₄P is Meso tetra(heptafluoropropyl)porphine Zinc

Molecular Formula: C₃₂H₈ZnF₂₈N₄

Molecular Weight: 1044;

ZnF₂PP is 5,10,15,20-Tetrakis(2,6-difluorophenyl)-21H,

23H-porphine Zinc(II)

Molecular Formula: C₄₄H₂₀F₈N₄Zn

Molecular Weight: 820;

InTPPC1 is 5,10,15,20-Tetraphenyl-21H,23H-porphine

Indium(III) Chloride

Molecular Formula: C₄₄H₂₈ClInN₄

Molecular Weight: 763;

ZnTMP is 5,10,15,20-Tetrakis(2,4,6-trimethylphenyl)-21H,

23H-porphine Zinc(II)

Molecular Formula: C₅₆H₅₂N₄Zn

Molecular Weight: 846 CAS: 104025-54-9;

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ZnTPP is 5,10,15,20-Tetraphenyl-21H,23H-porphine Zinc

Molecular Formula: C₄₄H₂₈N₄Zn

Molecular Weight: 678

5 CAS: 14074-80-7;

FeTFPPC1 is 5,10,15,20-Tetrakis(pentafluorophenyl)-21H,

23H-porphine Iron(III) Chloride Molecular Formula: C₄₄H₈ClF₂₀FeN₄

Molecular Weight: 1063.85

CAS: 36965-71-6;

ZnSi₆PP is 5(phenyl)-10,15,20-trikis(2',6'-disilyloxyphenyl)

porphyrinatozinc(II)

Molecular Formula: ZnC₈₀H₁₁₂O₆N₄Si₆

15 Molecular Weight: 1458;

ZnSi₇OHPP is 5,10,15-trikis(2',6'-disilyloxyphenyl)-20-(2'-

hydroxy-6'-silyloxyphenyl)porphyrinatozinc(II)

Molecular Formula: ZnC₈₆H₁₂₆O₈N₄Si₇

Molecular Weight: 1604;

ZnSi₈PP is 5,10,15,20-tetrakis(2',6'-disilyloxyphenyl)por-

phyrinatozinc(II)

Molecular Formula: ZnC₉₂H₁₄₀O₈N₄Si₈

Molecular Weight: 1718;

H₂TPP is 5,10,15,20-Tetraphenyl-21H,23H-porphine

Molecular Formula: C₄₄H₃₀N₄ Molecular Weight: 614.75

CAS: 917-23-7;

30 H₂FPP is 5,10,15,20-Tetrakis(pentafluorophenyl)-21H,23H-

Molecular Formula: C₄₄H₁₀F₂₀N₄

Molecular Weight: 974.57

CAS: 25440-14-6:

³⁵ Azodipyridine is 6'-Butoxy-2,6-diamino-3,3'-azodipyridine

Molecular Formula: C₁₄H₁₈N₆O Molecular Weight: 286.34

CAS: 617-19-6;

Rosaniline is Para-Rosaniline Base

Molecular Formula: C₁₉H₁₉N₃O

Molecular Weight: 305.4

CAS: 25620-78-4

FIG. 19 illustrates the response of the array described in FIG. 18 to acetone. As shown in FIG. 18 and summarized in Table 14 below, the color changes of each dye in response to aceteone are as follows (the exact colors depend, among other things, upon scanner settings). The color changes are derived simply by comparing the before exposure and after exposure colors and subtracting the two images (i.e., the absolute value of the difference of the red values becomes the new red value in the color difference map; etc. for green values and blue values). If there is no change in the red, green, and blue color values of a dye in the after-exposure image, then the color difference map will show black (i.e., red value=green value=blue value=0).

TABLE 14

(Summarizing the Dyes and Colors in FIG. 19, i.e., "Dye - Color")						
SnTPPCl ₂	CoTPP -	CrTPPCl -	MnTPPCl	FeTPPCl -	CuTPP - Black	
Reddish	Lavender	Gray	Pink	Black (no	(no change)	
Brown				change)		
AgTPP -	NiTPP -	InTPPCL -	IrTPPCl -	ZnTPP -	FeTFPPCl - Dark	
White	Light Teal	Blue	Light	Black (no	Dark Cobalt	
			Green	change)		

TABLE 14-continued

(5	(Summarizing the Dyes and Colors in FIG. 19, i.e., "Dye - Color")						
ZnSi ₆ PP -	ZnSi ₇ OHPP	ZnSi ₈ PP -	H ₂ TPP -	H ₂ FPP -	Alizarin basic -		
Black (no change)	Aqua	Dark Teal	Green	White Periphery and Blue Center	Dark Purple		
Me Red -	BCP -	BCP basic -	BTB -	BTB basic -	Ph Red basic -		
Dark Blue	Green	Light Green	Light Green	Dark Blue	Royal Blue		
Nile Red -	BCG - Tan	BCG basic -	CresRed -	CresRed	CP Red - Gold		
Olive		Black (no change)	Dark Pink	basic - Blue			
R Dye - Light Pink	TB - Brown	TB basic - Green	MeOr - Light Green	MeOr basic - Dark Blue	CP Red basic - Black (no change)		

Many modifications and variations may be made in the techniques and structures described and illustrated herein 20 plexes, and the second dye is distinct from the first dye and without departing from the spirit and scope of the present invention. Accordingly, the techniques and structures described and illustrated herein should be understood to be illustrative only and not limiting upon the scope of the present invention.

What is claimed is:

- 1. An artificial nose comprising an array, the array comprising at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a 30 distinct and direct spectral absorbance or reflectance response to distinct analytes, wherein the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phthalocyanine, and salen and their metal complexes, and the second dye is distinct from the first dye and selected 35 from the group of dyes consisting of acid-base indicator dyes and solvatochromic dyes.
- 2. The artificial nose of claim 1 wherein the first dye is a metalloporphyrin.
- 3. The artificial nose of claim 1 wherein the second dye is 40 an acid-base indicator dye.
- 4. The artificial nose of claim 1 wherein the second dye is a solvatochromic dye.
- 5. The artificial nose of claim 1 wherein the second dye is selected from the group consisting of Chlorphenol Red, 45 Bromocresol Green, Bromocresol Purple, Bromothymol Blue, Phenol Red, Thymol Blue, Cresol Red, Alizarin, Mordant Orange, Methyl Orange, Methyl Red, Reichardt's Dye, Nile Red, Congo Red, Victoria Blue B, Eosin Blue, Fat Brown B, Benzopurpurin 4B, Phloxine B, Orange G, Meta- 50 nil Yellow, Naphthol Green B, Methylene Blue, Safranine O, Methylene Violet 3RAX, Sudan Orange G, Morin Hydrate, Neutral Red, Disperse Orange 25, Rosolic Acid, Fat Brown RR, Cyanidin chloride, 3,6-Acridineamine, 6'Butoxy2,6diamino3,3'-azodipyridine, para-Rosaniline Base, Acridine 55 Orange Base, Crystal Violet, and Malachite Green Carbinol
- **6**. The artificial nose of claim **1** wherein the first dye is a porphyrin and has a periphery and a superstructure bonded to the periphery.
- 7. A method of detecting an analyte comprising the steps of: forming an array of at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflec- 65 tance response to distinct analytes wherein the first dye is selected from the group consisting of porphyrin, chlorin,

chlorophyll, phthalocyanine, and salen and their metal comselected from the group of acid-base indicator dyes and solvatochromic dyes, subjecting the array to an analyte, inspecting the array for a distinct and direct spectral absorbance or reflectance response, and correlating the distinct and direct spectral response to the presence of the analyte.

- 8. The method of claim 7 wherein the first dye is a metalloporphyrin.
- 9. The method of claim 7 wherein the second dye is an acid-base indicator dye.
- 10. The method of claim 7 wherein the second dye is a solvatochromic dye.
- 11. The method of claim 7 wherein the second dye is selected from the group consisting of Chlorphenol Red, Bromocresol Green, Bromocresol Purple, Bromothymol Blue, Phenol Red, Thymol Blue, Cresol Red, Alizarin, Mordant Orange, Methyl Orange, Methyl Red, Reichardt's Dye, Nile Red, Congo Red, Victoria Blue B, Eosin Blue, Fat Brown B, Benzopurpurin 4B, Phloxine B, Orange G, Metanil Yellow, Naphthol Green B, Methylene Blue, Safranine 0, Methylene Violet 3RAX, Sudan Orange G, Morin Hydrate, Neutral Red, Disperse Orange 25, Rosolic Acid, Fat Brown RR, Cyanidin chloride, 3,6-Acridineamine, 6'-Butoxy-2,6diamino-3,3'-azodipyridine, para-Rosaniline Base, Acridine Orange Base, Crystal Violet, and Malachite Green Carbinol
- 12. The method of claim 7 wherein the first dye is a porphyrin and has a periphery and a superstructure bonded to the periphery.
- 13. An artificial tongue comprising an array, the array comprising at least a first dye and a second dye deposited directly onto a single support in a predetermined pattern combination, the combination of the dyes in the array having a distinct and direct spectral absorbance or reflectance response to distinct analytes in solution or liquid analytes, or analytes in a solid or solid analytes, wherein the first dye is selected from the group consisting of porphyrin, chlorin, chlorophyll, phthalocyanine, and salen and their metal complexes, and the second dye is distinct from the first dye and selected from the group of dyes consisting of acid-base indicator dyes and solvatochromic dyes.
- 14. The artificial tongue of claim 13 wherein the first dye is a metalloporphyrin.
- 15. The artificial tongue of claim 13 wherein the second dye is an acid-base indicator dye.
- 16. The artificial tongue of claim 13 wherein the second dye is a solvatochromic dye.

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17. The artificial tongue of claim 13 wherein the second dye is selected from the group consisting of Chlorphenol Red, Bromocresol Green, Bromocresol Purple, Bromothymol Blue, Phenol Red, Thymol Blue, Cresol Red, Alizarin, Mordant Orange, Methyl Orange, Methyl Red, Reichardt's 5 Dye, Nile Red, Congo Red, Victoria Blue B, Eosin Blue, Fat Brown B, Benzopurpurin 4B, Phloxine B, Orange G, Metanil Yellow, Naphthol Green B, Methylene Blue, Safranine O, Methylene Violet 3RAX, Sudan Orange G, Morin Hydrate, Neutral Red, Disperse Orange 25, Rosolic Acid, Fat Brown 10 RR, Cyanidin chloride, 3,6-Acridineamine, 6'-Butoxy2,6diamino-3,3'-azodipyridine, para-Rosaniline Base, Acridine Orange Base, Crystal Violet, and Malachite Green Carbinol Base.

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- 18. The artificial tongue of claim 13 wherein the first dye is a porphyrin and has a periphery and a superstructure bonded to the periphery.
- 19. The artificial nose of claim 1 further comprising a low volatility liquid.
- 20. The method of claim 7 further comprising the step of adding a low volatility liquid to the array.
- 21. The artificial tongue of claim 13 further comprising a low volatility liquid.
- 22. The method of claim 7 further comprising the step of forming a table of responses of the array to a plurality of distinct analytes.

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