(54) Title: MATRIX-FREE DESORPTION IONIZATION MASS SPECTROMETRY USING TAILORED MORPHOLOGY LAYER DEVICES

(57) Abstract: There is disclosed an apparatus for providing an ionized analyte for mass analysis by photon desorption comprising at least one layer (11) for contacting an analyte, and a substrate (10) on which said layer (11) is deposited. Upon irradiation of said apparatus, said analyte desorbs and ionizes for analysis by mass spectrometry. The layer of layers of said apparatus comprise a continuous film a discontinuous film or any combination thereof.
MATRIX-FREE DESORPTION IONIZATION MASS SPECTROMETRY
USING TAILORED MORPHOLOGY LAYER DEVICES

BACKGROUND OF THE INVENTION

Field of the Invention

The present invention relates to tailored-morphology material systems and their use in molecular mass analysis by electromagnetic energy desorption-ionization mass spectrometry. Areas of interest for this technology include, but are not limited to, chemical research and manufacturing, pharmaceutical research and manufacturing, bio-medical research and screening, head-space and environmental monitoring, and other applications involving molecular analysis.

Description of the Prior Art

Light desorption-ionization mass spectrometry is a very common and powerful technique for mass analysis of molecules. It is a technique which can be broadened to include the whole spectrum of electromagnetic energy for the desorption-ionization step. However, with recent demands in throughput and small molecule screening, the most popular and widely used laser-based technique, known as MALDI (matrix-assisted laser desorption-ionization), has limitations. MALDI was developed in the mid-eighties and is still being refined today for the analysis of a wide range of compounds with emphasis on proteins, peptides and other molecules in the range of 500 – 200,000 amu. In MALDI, the analyte (the molecules or compounds to be analyzed) is mixed in with an organic UV absorbent "matrix". This matrix provides a "soft" method of desorbing large
molecules by allowing excess energy in the analyte to be transferred to the matrix molecules during the desorption process. The matrix also provides an environment suitable for the protonation of the analyte molecules, giving them a single, positively charged state. However, for small molecules (approximately 500 amu and below, such as drug molecules), the matrix molecules themselves provide background in the signal and complicate spectrum analysis. Furthermore, with modern demands in automation, throughput and reproducibility, the addition of the matrix to the analyte and its preparation become issues particularly in the case of throughput. These limitations were recognized during the onset of MALDI, leading to the study of non-matrix methods.

The first studies in matrix-less light desorption from a surface used metals and glasses as a media to immobilize the analyte molecules. These materials had non-textured morphologies, i.e., essentially they were non-porous and had a flat (continuous) surface. In a study using this approach, two incident light beams were used, one to desorb and one to ionize the molecules. Zhan, Q. et al., Amer. Soc. Mass Spectrom. 8, 525-531 (1997). This approach is termed two-photon ionization. Other similar methods used ion beams and thermal sources for these tasks. Problems with all these matrix-less light desorption techniques reported in the literature include a high degree of molecular fragmentation and a very limited mass range. These studies, and recent comments, maintain that smooth (non-porous) surfaces do not work effectively for matrix-less laser mass desorption. (See for example, Wei, J., et al., Nature. 399, 243-246 (1999). A recent report supports the understanding that smooth surfaces do not function effectively for matrix-less laser mass desorption. Kruse, R., et al., Anal. Chem. 73, 3639-3645, 2001.

It has been shown that matrix-less laser mass desorption could be effective if done on a textured surface created with the use of
electrochemically etched porous silicon. Wei, J., et al., Nature. 399, 243-246 (1999). With this material as a substrate for laser desorption ionization, significant improvement in non-matrix techniques for molecular analysis has been reported. Also, it was reported that electrochemically etched porous silicon provided mass detection in the range of 0 – 8000 amu with little fragmentation and little low mass noise. However, other results using this material raised concerns about the low mass collection of hydrocarbons and other contaminants leading to “dirty” low mass signals. Shen, Z., et al., Anal. Chem. 73, 612-619 (2001). The use of HOME-HF electrochemically etched Si, GaAs and GaN, which requires metallic patterning and a wet etching step leading to a porous microstructure, has also been reported for matrix-less laser mass desorption. Kruse, R., et al., Anal. Chem. 73, 3639-3645, 2001.

Further limitations of the electrochemically etched materials are their limited useful lifetimes for mass desorption-ionization applications (<3 weeks) which occur for these materials because of etchants trapped in the material during its manufacturing process. The processing of these etching approaches involves the galvanic etching of a crystal conductive substrate in a hydrofluoric acid based solution. Although the fundamental theory of the mechanism of desorption-ionization of molecules using these techniques is currently under investigation, research groups using these materials reported the importance of the porous structure to the success of mass desorption-ionization and reported that solid, smooth (i.e., non-textured) silicon and silicon dioxide coated silicon did not generate ion signals; i.e., were not useful for light desorbed mass spectroscopy.

In other work, liquid matrix materials combined with UV light adsorbing particles have been used in recent laser desorption/ionization experiments as an alternative to traditional MALDI matrix materials. Dale et al, Anal Chem, 68, 3321-3329 (1996) used a glycerol/graphite slurry to desorb
detect proteins and peptides. This methodology proved less efficient in ionization than traditional MALDI and provided a very noisy spectrum from the glycerol contamination.

The use of a new material, deposited column/void network silicon, for laser desorption-ionization has eliminated several disadvantages associated with electrochemically etched material approaches. Cuiffi, J., et al., *Anal. Chem.* 73, 1293-1295 (2001). This reported technique of using deposited column/void network materials for mass desorption-ionization produced similar mass ranges and sensitivity to electrochemically etched material, but the film itself did not degrade over time. Furthermore the manufacturability of a deposited film system offers several advantages in cost, production throughput, contamination control, uniformity, and signal reproducibility. This deposited material also offers the further unique feature of having the capability to be deposited on a number of inexpensive substrates, including bio-degradable materials, plastics, and glass. On the other hand, electrochemically etched material always must be on a conducting substrate. In addition, Cuiffi et al. reported, for the first time, that solid (continuous) films of crystalline silicon and thermal silicon dioxide coated crystal silicon did give effective mass desorption-ionization spectroscopy signals. Cuiffi, J., et al., *Anal. Chem.* 73, 1293-1295 (2001).

The material systems of the present invention consist of one or more deposited film layers and a substrate on which they are deposited. The material system could also be grown (e.g., Si, SiGe alloy, Ge wafer materials) or casted (Si, SiGe, Ge sheet materials) and also function as the substrate. Unlike the previously reported techniques, our deposited material systems offer the flexibility of a number of deposition methods and encompass a broad range of material and morphological choices. These material systems can be uniquely tailored for mass spectrometry
applications through choice of the substrate, deposition techniques and materials, deposition parameters and pre- or post deposition physical and chemical modification, which are unavailable in the techniques of electrochemically etched porous silicon whether used with one or two-photon ionization. Specifically, the substrate materials available with our technique are chosen from a group consisting of polymers, plastics, biodegradable materials, semiconductors, metals, ceramics, insulators, glasses or combinations thereof. Electrochemically or HOME-HF etched porous materials require a conducting semiconductor substrate, and are fundamentally based on a subtractive electric current-driven etching process.

The materials of the present invention can be deposited. This can be done by one or a combination of the additive process comprising physical vapor deposition, chemical vapor deposition, molecular beam epitaxy, plasma assisted physical vapor deposition, plasma enhanced chemical vapor deposition, sol-gel, molecular self-assembly, electroplating, tape casting, spin casting, casting, liquid deposition, or assembly from liquid chemical precursors. The morphology of these materials, which is determined by the production technique and parameters, are application-specific and can range from a continuous (void free) solid with no surface texturing to high surface area to volume ratio (i.e., the deposited column-void nanotextured silicon film), or any intermediate morphologies.

The material systems of the present invention can also be altered by pre or post deposition physical or chemical modifications, which affect the morphology, surface chemistries and bulk material chemistries of the films.

Given these advantages, our material systems are easily integrated, when compared to other matrix-less light desorption/ ionization
techniques, with microelectronics, micro-fluidics and other micro and nanofabricated sensing devices.

Matrix free desorption/ionization mass spectrometry available using the tailorable morphology of our materials, has a variety of applications. The flexible nature of the substrate material composition, film composition, or both, permitted in our approach allows this technology to be used in atmospheric and reduced pressure desorption and ionization systems as a disposable consumable or reusable target. The composition and methods of production utilized in this technique allow for easy integration with microfabrication processes and microelectronic devices, such as microfluidics, microarrays, CMOS technology and thin film transistors. The matrix less desorption and ionization makes automated, high throughput sample analysis an attractive use of this technique.

The present invention presents a variety of structures that further expand the possibilities of molecular detection using light desorption-ionization, by providing low-cost, easily manufactured, tailor able material systems and techniques.

**SUMMARY OF THE INVENTION**

The present invention is directed to a class of layered structures comprising one or more layers, with tailored application-specific morphology, for use in light desorption-ionization mass spectrometry. This class of structures holds analytes and allows them to be desorbed and ionized in the presence of a light source for subsequent mass analysis. Analytes are preferably in an amount less than one millimole. Analyte is selected from the group consisting of organic chemical compositions, inorganic chemical compositions, biochemical compositions, cells, microorganisms, peptides, polypeptides, proteins, lipids, carbohydrates, drug
candidate molecules, drug molecules, drug metabolites, combinatorial chemistry products, nucleic acids, and any combinations thereof.

In order to perform [these two] the functions of desorption and ionization, the film structure and substrate of this invention must (1) effectively couple and absorb the incident electromagnetic energy (e.g., light), (2) transfer the energy from the incident energy into the analyte for desorption/ionization, and (3) provide the necessary surface and surroundings for the analytes to be desorbed and ionized. The structures of this invention may also be patterned or textured for tasks including increasing the surface area, enhancing ionization, enhancing optical absorption, and localizing the analyte. Also, chemical additives to the deposited films or analyte solution may also be used to enhance ionization and analyte detection. Finally, these material systems can be easily integrated with macro-scale and micro-scale devices. These aspects are explained and further detailed below. The class of structures of the present invention encompasses devices with one or more deposited films and a substrate to which they are adhered. The class of structures of this invention also encompasses structures with one or more layers from grown or caste materials.

Each layer in the device, including the substrate may perform one or more tasks. Two necessary tasks are the absorption of the light (done by the “absorption layer”) and holding of the analyte (done by the “immobilization layer”). Other tasks may include enhancing optical coupling of the light into the absorber via increasing optical path length and/or optical impedance matching, enhancing thermal energy transfer into the analyte via high thermal conductivity, controlling drop drying and crystallization and providing a source of ionizing or ionizing enhancing reagents. A layer may also be present to apply a bias to the analyte-bearing layer during the light impingement step.
The present invention discloses an apparatus for providing an ionized analyte for mass analysis by light desorption mass spectrometry comprising at least one layer for contacting an analyte, and a substrate on which said layer is deposited, wherein said analyte upon irradiation of said apparatus with a light source desorbs and ionizes for analysis by mass spectrometry. The substrate is selected from the group consisting of semiconductors, glasses, plastics, polymers, biodegradable or biocompatible materials, metals, ceramics, insulators, organic materials, and any combinations thereof. At least one layer is selected from the group consisting of metals, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, and any combinations thereof. At least one layer may be a continuous (non-textured) film, a textured (columnar or columnar-void) film, or any combinations thereof, and is deposited by physical vapor deposition, chemical vapor deposition, liquid deposition, molecular beam epitaxy, plasma assisted chemical vapor deposition, sol-gels, nebulization, electroplating, tape casting, spin coating, self-assembly, assembly from liquid chemical precursors, printing, and any combinations thereof.

The present invention also discloses a method for providing an ionized analyte for analysis of mass comprising providing an apparatus comprising at least one layer for contacting an analyte wherein said layer is deposited on a substrate, contacting an amount of an analyte containing entities such as molecules whose mass or masses are to be determined with said deposited layer, and irradiating said apparatus to desorb and ionize said analyte. Also, the present invention discloses a method for determining a physical property of an analyte component comprising providing an apparatus comprising at least one layer for contacting an analyte and a substrate on which said layer is deposited; positioning an amount of an analyte on the layer used for contacting an analyte of said
apparatus; irradiating said apparatus having said contacted analyte; desorbing and ionizing at least one component of said analyte; and analyzing said ionized analyte component for a physical property, preferably mass to charge ratio of the ionized analyte.

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**BRIEF DESCRIPTION OF THE DRAWINGS**

Fig. 1 is a schematic representation of the difference between MALDI (top) and the method of the present invention (bottom).

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Fig. 2 is a mass spectrum obtained using a silicon dioxide layer on top of silicon.

Fig. 3 is a mass spectrum obtained using a deposited germanium thin film.

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Fig. 4 is a mass spectrum obtained using a high surface to volume silicon material.

Fig. 5 a-f show various material system embodiments of the present invention.

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Fig. 6 Transmission and reflectance spectra of the Halogenated acidic polymer – carbon black composite film. Reflectance is with respect to barium sulfate.

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Fig. 7. Desorption ionization mass spectrum taken from the surface of the Carbon black – Halogenated acidic polymer film.
DETAILED DESCRIPTION OF THE INVENTION

Referring now to Fig. 1, there is shown a schematic representation of the MALDI and the method of the present invention. Ultra-violet light (337 nm) 1 impinges matrix 2 to provide sample and matrix ions and neutrals 3 to reach detector 4.

Referring now to Fig. 5 which shows devices of various materials, Fig. 5a shows a device having a high surface area to volume ratio film or layer of columnar silicon 11 on a plastic substrate 10. The functions of the columnar silicon layer 11 are absorption, optical coupling, and immobilization. Advantages of this device include, but are not limited to, one-step production, inexpensive substrate material, and high molecular immobilization.

Fig. 5b shows a device having a layer of silicon dioxide 15 on a layer of amorphous silicon 12, on a layer of metal 13, on a glass substrate 14. The function of the amorphous silicon layer 12 is light absorption. The metal 13 and silicon dioxide 15 provide optical coupling; also, the silicon dioxide layer 15 provides immobilization. Advantages of this device are that it is reusable and provides little low mass noise.

Fig. 5c shows a device having a layer of silicon dioxide 17 on a substrate of crystal silicon 16. The function of the crystal silicon substrate 16 is light absorption. The functions of the silicon dioxide layer 17 are optical coupling and immobilization.

Fig. 5d shows a device having a layer of amorphous silicon 19 on a textured plastic substrate 18. The functions of the amorphous silicon layer
19 are light absorption and analyte immobilization. The function of the textured plastic substrate 18 is optical coupling.

Fig. 5e shows a device having a layer of amorphous silicon 21 on a glass substrate 20. The functions of the amorphous silicon layer 21 are light absorption and analyte immobilization. Neither the glass substrate 20 nor the amorphous silicon layer 21 provides optical coupling. The advantages of this device include, but are not limited to, one-step production of manufacture, and low mass noise.

Fig. 5f shows a device having a high surface area to volume ratio silicon dioxide (porous SiO$_2$) layer 24 on a layer of amorphous silicon 23 on a glass substrate 22. The device is preferably illuminated from below, i.e., from the glass substrate layer 22. The function of the amorphous silicon layer 23 is light absorption of the back illumination. The function of the porous SiO$_2$ layer 24 is analyte immobilization. The function of the glass substrate layer 22 is optical coupling. Advantages of this device include, but are not limited to, elimination of direct light exposure of the analyte by providing illumination from below, and high molecular immobilization.

Fig. 6 is a graphic representation of transmission and reflectance spectra a halogenated acid polymer - carbon black composite film of the present invention. Reflectance is with respect to barium sulfate. Exceptionally low reflectance and almost no transmission indicated that the composite film absorbs most of the light impinging on it in the visible and ultraviolet range.

Fig. 7 is a graphic representation of a desorption ionization mass spectrum taken from the surface of a carbon black - halogenated acidic polymer film of the invention.
Absorption layer

5 A necessary role that one or more layers, which may include the substrate, must play is the absorption of the incident photons and the conversion of this light into species-desorbing energy. The optical properties required by this absorption layer are determined by the electromagnetic energy (e.g., light) source used. For example, silicon makes an excellent absorber for an ultra-violet light source such as common nitrogen laser (wavelength of 337nm), but may not work effectively with an infra-red source (depending on the wavelength) because of its optical bandgap structure. On the other hand, Ge will work for the shorter infra-red wavelengths. In general this absorber layer composition is selected to match the wavelength or wavelengths of the impinging electromagnetic radiation. Materials that can be used for this layer, with appropriate optical properties, include conductors, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, or composites thereof. Proper choice of this layer can allow the electromagnetic energy source to be a variety of possibilities including light emitting diodes or lasers.

A micro-composite or nano-composite polymer film can also be used for this layer, which may include the substrate, in matrix-less photon desorption mass analysis devices or apparatuses. For example, a polymer including but not limited to, an acidic halogenated polymer, or mixtures of halogenated polymers, may be added to an inorganic or organic material, or combinations thereof. A preferred composite embodiment comprises a fluorinated, acidic polymer and carbon black. Composite polymer films or layers are conveniently prepared by mixing the selected organic polymer and inorganic/organic material (i.e.,
fluorinated acidic polymer and carbon black) in a suitable solvent; and thereafter forming a composite film from the mixture. The composite film may be formed by methods known in the art such as molding, casting, spin casting, spraying, and any combinations thereof. An advantage of using composite polymer films is that desirable properties of materials essential to laser desorption ionization are consolidated. For example, when a fluorinated, acidic polymer/amorphous carbon composite comprising carbon black particles embedded or suspended in the polymer matrix is used, carbon black particles efficiently absorb the laser energy and convert it to heat while the acidic polymer provides a means of holding the carbon particles together and can function to mediate proton transfer as a donating medium for the analyte. Modifying the polymer content or polymer chemistry also offers a means to control the surface energy of immobilization layer and therefore modify the adsorption, drying and crystallization of the analyte.

Immobilization layer

The one or more necessary layers which are in contact with the sample atoms and/or molecules must hold the sample and allow it to effectively desorb and ionize, enabling it to interact with any ionizing/ionization enhancing agents if necessary. This layer may be composed of the same or similar material as the light coupling, thermal coupling, or absorption layer or may differ in both chemical composition and physical morphology. The morphology of this layer may range from solid, flat-surface (non-textured) material to high surface area to volume ratio very highly nano-textured material. The morphology of the film may be used to affect the mechanics and kinetics of analyte application, adsorption and concentration, and/or the adsorption and concentration of the ionizing and ionization enhancement agents. This affects signal properties, including but not limited to, sensitivity and resolution.
The chemical composition of this layer or layers can also affect signal response by modifying the interaction of the atomic and/or molecular species and other compounds with each other and the layer. Chemical composition of this layer may change species adsorption, desorption, ionization, and conductivity and molecular affinity of the layer. The chemical composition of this layer may also affect its ability to be cleansed of noise (non-analyte) molecules during analyte positioning. The bulk and surface chemistries may be specifically tailored, by controlling layer processing (e.g., casting, deposition) chemistry or by post layer processing modification for controlling the aforementioned properties. We have demonstrated that hydrophobic or hydrophilic and acidic or basic surfaces influence analyte desorption and ionization by modifying analyte, ionizing agent and surface interaction. The reduction of van der Walls and hydrogen bonding via surface chemistry also may enhance analyte desorption/ionization. The immobilization layer or layers that are in contact with the sample atoms and/or molecules may be comprised of conductors, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, or composites thereof.

This layer of the matrix-less devices of the present invention for photon desorption mass analysis may be a composite polymer containing film. The film may be either a micro-composite or a nano-composite film. Such composite polymer films include, but are not limited to, a suitable acidic halogenated polymer, or mixtures of acidic halogenated polymers, and a suitable organic or inorganic material, or combinations thereof. A preferred composite embodiment comprises acidic fluorinated polymer and the material carbon black.

Optical coupling layer
Another task that may be performed by one or more layers of the material system of this invention is coupling the incident light more effectively into the absorption layer. Several techniques can be used including optical impedance matching, anti-reflection coating, increasing the optical path length, and combinations thereof. Materials that can be used for this layer, with appropriate properties, include metals, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, or composites thereof.

Substrate

For the material system of this invention, the substrate may play one or more of the roles mentioned above or simply serve as a support medium. The only necessary qualification of the substrate is that it must be compatible with the processing used to create subsequent layers. Materials that can be used for this layer include metals, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, or composites thereof.

One or more of the different layers of the devices of the invention may be a conductor capable of being biased during the impingement of the desorbing light. Such layer biasing may be used to affect ionization of the analyte.

Processing methods

The layers of these morphology tailored structures can come from grown or caste materials. They can be deposited films produced by a variety of methods including but not limited to the following: PVD such as sputtering, evaporation, and PEPLVD, CVD PECVD, ECR-PECVD, MOCVD electroplating, so-gel, tape casting, spin coating, nebulization,
deposition, self-assembly, casting, liquid deposition, or assembly from liquid precursors, and any combinations thereof.

Composite polymer layers or films of the devices or apparatuses of the present invention may be prepared by methods known in the art. For example, organic polymers or polymers are mixed with organic or inorganic material or materials in a suitable solvent. A preferred organic polymer is a crosslinked halogenated acidic polymer. A preferred inorganic material is carbon black. The mixture is then formed into a film with removal of the solvent using methods known in the art such as molding, casting, spin casting, spray, and any combinations thereof.

Device structure and layer organization

The layer options and requirements detailed above enable the material structures of this invention to be uniquely tailored for optimal performance based on the analytes, their sample preparation, the type of electromagnetic energy source used for desorption-ionization, the mass analysis technique and integration techniques. However, several basic rules apply to the overall device structure. These rules apply whether the electromagnetic energy enters through the immobilization layer holding the analyte (the front) or from the back. First, the impinging photons must be able to enter the absorber layer or layers of the device. Second, the immobilization layer must enable the desorbed analytes to have access to the mass detector.

Device texturing and patterning

The texturing of one or more layers in the device of this invention can have several effects. Texturing of a reflective layer behind the absorber (on the opposite side of the device from the light source),
increases the optical path length and enhances optical absorption in thin absorber layers. Texturing of the immobilization layer can have various effects depending on the length scale considered. Micro- or nano-scale or both roughness allows more analyte and desorption-ionization enhancing agents to be present per incident impinging photon area. This can increase the sensitivity and longevity of analyte signal. On the nanoscale, the surface roughness will not only enhance signal sensitivity and longevity by increasing the absolute amount of analyte present, but may also act to enhance light coupling into the absorber by optical impedance matching. Texturing can be done by a variety of manufacturing techniques including pre-fabricated or molded substrates, physical roughening, laser ablation, lithographic processes, etching processes and textured film growth.

Patterning of one or more of the layers in the device can serve many purposes. The localization of analytes, which is important for automation and sample delivery purposes, can be done on a macroscopic or microscopic scale with wells either pre-formed on the substrate or produced during subsequent film growth and processing. Furthermore, patterning of the immobilization layer via differences in hydrophobicity, hydrophilicity, chemical affinity, charge and polarity can localize and preferentially bind desired analytes. Patterning of a metallic grid on the immobilization layer can remove charge buildup during the desorption/ionization process in machines requiring a grounded stage. Finally, with the integration of these devices into micro-fluidic systems, patterning can be used to define the device location. Patterning can also be done by a variety of manufacturing techniques including pre-fabricated or molded substrates, physical scribing, laser ablation and lithographic processes.

Photon Impingement Protocols
To extend the temporal duration or amount of analyte signal
generation, it may be necessary to have the impinging photons execute a
pattern in each given analyte-containing region. Such patterns would
allow more analyte to be desorbed and may involve multiple paths across
a given region. These protocols would be pre-programmed.

Chemical additives

Chemical additives, which are allowed to interact with the analyte
molecules during the desorption-ionization process, can act to enhance
analyte detection. In order to increase the proportion of charged analyte
species to neutrals, the surrounding environment can be made more
acidic or basic. To create a more "soft" ionization process known
molecules can be used with the analyte to act as a cooling media by which
excess thermal energy may be transferred from the analyte during
desorption. This greatly reduces the fragmentation of large molecules
during mass analysis. Other chemical additives can act to condition or
purify the surrounding media by chelating metal and salt ions known to
reduce sensitivity and cause adducts. Hydrated molecules such as
hygroscopic salts or other water containing molecules can provide a
source of ions prior to or during desorption. Finally, chemical additives
such as surfactants and detergents can change the way analyte molecules
and contaminates interact with each other and the immobilization layer
surface. This can be useful in cleaning the surface during sample
application, preventing agglomeration of analyte and preventing strong
adherence of the analyte to the immobilization layer.

These chemical additives can be introduced into the process in the
analyte preparation step, pre-coated on the immobilization layer,
chemically attached to the immobilization layer, or introduced during desorption-ionization via fluidic or gaseous transport.

Mixed phase films

Matrix free desorption ionization mass spectrometry can also be mediated by mixed phase or composite material surfaces. In particular, when the desorbing/ionizing layer material is available in less expensive powder form, a cost effective approach is to form layers of this material from powder such that the particles are fixed in/by a resin material. The second ("glue") material may serve more than just fixing the particles together, it may also function as the radiation absorber and/or ionization enhancer. It is possible that the particles and the "glue" material may have completely distinct roles essential to desorption/ionization. For instance, the particle material may be a strong absorber while a poor ionization enhancer. In contrast, the second material may be a poor absorber but be or contain effective ionization enhancer(s). On the other hand, the superior properties of the two materials (radiation absorption and ionization enhancement) can be brought together in a composite by mixing them. In this way a superior desorbing/ionizing layer can be obtained. The composite layer could be comprised of more than two materials (either in particulate or glue form) to better tailor its superior desorbing/ionizing properties. A low cost method of making a composite film is mixing its components in a liquid solvent, and then placing the liquid mixture onto a substrate (e.g., casting, spinning, spray, brush, dipping, printing etc. techniques).

As a specific example, amorphous carbon – halogenated acidic polymer composite films were prepared by spin casting. The carbon black produced from acetylene with an average particle size of 0.042 μm. The Halogenated acidic polymer solution with 5.0-5.4% polymer content by
weight was purchased from DuPont. Carbon black was mixed into the
Halogenated acidic polymer solution to an equal amount of polymer by
weight. The mixture was ultrasonicated for 6 h and stirred for 12 h before
spun on 1"×1" Corning 1737 glass substrates. A uniform film thickness of
1.8 μm was obtained at a spin rate of 2000 rpm in 40 s. The spin on was
followed by a 120 °C thermal anneal for 15 minutes. The resultant film
was found to be a very efficient light absorber in the visible and UV range
as seen from its exceptionally low reflectance and transmission
characteristics in Fig. 6. This is simply attributed to amorphous carbon’s
being a very efficient light absorber. On the other hand, the role of
Halogenated acidic polymer other than being a resin could be ionization
enhancement. This is because Halogenated acidic polymer, a
perfluorosulfonic acid/tetrafluoroethylene copolymer in the acid (H⁺) form,
is well known for its being an efficient proton storage, transport and
exchange properties. The conductivity of the film was measured to be
about 1.6 S/cm using glass substrates with coplanar metal contacts. Fig.
2 depicts a desorption ionization mass spectrum taken from the surface of
the Carbon black – Halogenated acidic polymer film after a 1.0 ng
reserpine was dried from a droplet on the surface. It is evident from Fig. 7
that a very clear analyte signal is obtainable.

Integration with preparation and application devices.

The deposited devices and mass analysis technique of this
invention have the unique ability to be integrated with a large number of
sample delivery and preparation techniques plus a large number of mass
analyzers. Preparation of the analyte molecules can be as simple as
placing a drop of the molecules on the immobilization layer surface and
allowing them to dry. It is also possible to allow the analyte molecules to
adsorb to the immobilization layer surface out of a gaseous or liquid
solution. This simple fluid handling can be performed by a number of
automated, high throughput handling systems. More complex schemes include the use of micro-fluidic, on chip, system that perform chromatography or purification. The deposited systems of this invention can be easily used in tandem with a chip-based system or integrated into the micro-fluidic device. The use of an integrated micro-fluidic system can also be used to deliver desorption-ionization enhancing agents to the analyte during mass analysis in order to prolong detection signal.

Many mass spectrometry techniques can be used to analyze the desorbed-ionized species. These may include but are not limited to: time of flight, quadrapole, ion trap, plasmon resonance or combinations thereof.

The present invention comprises a class of morphology-tailored structures (or material systems) for the mass analysis and a method of analysis of atoms, molecules and molecular compounds and complex structures such as adhered cells when coupled with light desorption-ionization mass spectrometry. These material systems act to hold analytes and allow them to desorb and ionize in the presence of light without the traditional organic or non-organic matrix. A schematic of the difference between traditional MALDI and the technique of this invention is given in Figure 1. The material systems of this invention are composed of one or more layers and a substrate to which they are adhered. The critical roles of adsorption, analyte immobilization, optical coupling, and substrate may be played by one or more material layers. one or more of these layers may be biased during photon exposure to influence desorption. The specifics of layer function and formation are detailed below.

Absorption layer. A necessary role that one or more layers, which may include the substrate, must play is the absorption of incident photons from the light source. The light source may range from IR to UV wavelengths and from coherent and in phase (laser) to non-coherent. The
choices of light source and absorber material are dependent on each other. The absorber must be able to absorb enough of the incident light to provide sufficient thermal energy to desorb the analytes from the immobilization layer, which may or may not be the absorber layer itself. The high light adsorption coefficients of semiconductor materials make the use low cost, light emitting diodes as a light source an attractive option, when compared to the traditional UV laser sources that are necessary for MALDI. A specific embodiment of this invention is the use of a 337nm UV light source and Si or Ge based absorber materials. These two materials, in amorphous through crystalline phases, absorb UV light very efficiently, and we have demonstrated this in Figures 2 and 3. This idea can easily be extrapolated to include all semiconductors in the binary, tertiary, mixed, and graded varieties. All other materials for use as an absorber, with appropriate optical properties are encompassed by the scope of this invention including: metals, semiconductors, insulators, ceramics, polymers, organic materials or composites thereof.

The only requirement of the position of the absorber layer in the material system of this invention is that the incident photons have access to this layer. A unique aspect of this invention is the ability to illuminate the device from any direction including through the substrate (rear of the device) and through the immobilization layer (front of the device). This is important if the analytes adsorb the light wavelengths used for desorption and ionization. For instance small molecules, peptides and proteins adsorb UV wavelengths efficiently, which can lead to thermal degradation and fragmentation of the analyte, reducing the sensitivity of detection. Unlike MALDI, in our system the analyte is not required to sit in the photon path, thus entirely avoiding any photon/analyte interactions.

**Immobilization layer.** One or more layers in the material system of this invention must come into contact with the sample atoms/and or
molecules. This layer must hold the sample and allow it to effectively
desorb and ionize in the presence of the energy generated by light
adsorption in absorber layer, which may also act the immobilization layer.
The material properties required of the immobilization layer range widely
and depend highly on its interaction with the sample species. Also, if the
immobilization layer is in the light path between the light source and the
absorber layer, it must have optical properties such that incident photons
are allowed to reach the absorber. Specific materials of this invention
used for the immobilization layer include but are not limited to silicon,
germanium, silicon dioxide, germanium oxide, and their alloyed forms. All
other materials for use as an immobilization layer, with appropriate
material properties are encompassed by the scope of this invention
including: metals, semiconductors, insulators, ceramics, polymers, organic
materials or composites thereof.

A. Morphology of the immobilization layer

The morphology and physical structure of this layer can be tailored
for specific applications. Three morphological structures of the
immobilization layer, specific to this invention, include nanometer range
texturing, micrometer range texturing, and a macro-scale flat surface. The
first two types of films we categorize as discontinuous films. The macro-
scale film is what we term a continuous film. The advantages and
disadvantages of these three film structures are given in Table 1. The
immobilization layer may be comprised of one or more of these
morphological features.

Table 1. Morphological structures of the immobilization layer of this
invention and their advantages and disadvantages
<table>
<thead>
<tr>
<th>Morphology</th>
<th>Properties</th>
<th>Advantages</th>
<th>Possible Disadvantages</th>
<th>Examples</th>
</tr>
</thead>
</table>
| Nanometer scale texture | - Ultra high surface area  
- High steric interaction with molecular species  
- Strong capillary forces | - Very high loading capacity of analyte  
- High adsorption of analyte species from wet or dry ambient  
- Excellent uniformity of analyte coverage | - High adsorption of ambient noise | - Nanoscale deposited column/void network material |
| Micrometer scale texture | - High surface area | - High loading capacity of analyte  
- Low adsorption of ambient noise | - Glancing angle deposited films | -                                        |
| Flat surface | - Low surface area | - Very low adsorption of ambient noise | - Poor uniformity of analyte coverage  
- Low sample loading density | - Evaporated, spun-on, or sputtered materials |
B. Chemical modifications and additives to the immobilization layer

The surface and bulk chemistry of this layer can also be tailored during layer processing or post layer processing for specific interactions with the analyte molecules, desorption/ionization enhancing species, and the immobilization layer surface. For example, for deposited layers the chemistry of the film can be modified by plasma, thermal or wet chemistries such as, but not limited to; RIE, CVD, PECVD, DVD, MOCVD, PVD and wet chemical modification. A specific embodiment of this invention is to use surface chemical modifications either during or after film deposition to control hydrophobicity and hydrophylicity of the film, such as the incorporation of carbon and fluorine while depositing a film or the growth of a thermal oxide. The chemistry of the immobilization layer can be tailored for certain molecules to improve their desorption and ionization efficiency. Other functional groups can also be used to tailor the interaction of the surface with the analyte molecules by altering hydrogen bonding, surface charge, van Der Walls interaction, polar and non-polar interactions, steric interaction, antigen/antibody reactions etc. The surface chemistry and energy can play a critical role in the manner an analyte interacts with the surface during adsorption. The manner in which a analyte crystallizes can play a large role in the efficiency with which it desorbs and ionizes. As an example, composite halogenated, acidic polymers provide an excellent surface for analyte crystallization, while the acidic groups provide a source of ions for the ionization process. Crosslinked polymers are a more thermally stable surface that provides spectrums with very little noise from polymeric breakdown. Carbon black/halogenated polymer composites possess an extremely hydrophobic surface composition, when compared to the hydrophobicity of the polymer surface alone. The water contact angle of these materials are in excess of 100 degrees.
Other chemical additives specifically enhance the ionization of the analyte molecules. The additives can be chemically bonded to the surface prior to analyte application, applied to the surface in solid liquid or gas phase, or applied into the analyte solution prior or during mass analysis. In order to improve ionization efficiency, a number of materials, including but not limited to additives that are salts, hydrated molecules, surfactants, detergents, chelators, acids and bases, may be added on the surface of the apparatus and dried prior to the addition of the analyte, or added to the analyte prior to or during contacting the analyte to the apparatus. A specific chemical modification for improving ionization efficiency is the control of surface pH to enhance either negative or positive ion spectra. For example this can be accomplished simply by allowing HCl or Trifluoroacetic acid (TFA) to dry on the immobilization layer prior to applying the analyte or attaching and acid or basic group to the immobilization layer surface using a silanization reaction. Hydrated salts such as MgCl₂ can provide a significant source of protons in a crystallized analyte for ionization. Chelating agents such as ammonium citrate remove salt ions which form adducts during mass analysis and also disperse analytes for more uniform spatial distribution. Other small molecules added to the analyte, such as amino acids interact with the analyte in the desorption plume and adsorb energy from the analyte reducing fragmentation during mass analysis. HCl and TFA may also be added to the analyte.

Another useful chemical modification can act to self-clean the device during sample application. By using a layer composition or thin surface coating that is soluble in the sample solvent, the coating will be dissolved, "cleaning" the surface of adhered contaminants such as hydrocarbons. A specific embodiment of this invention is the use of a water-soluble germanium oxide for its self-cleaning properties. Such an
oxide will be inherently present as soon as Ge is exposed to atmosphere. This nascent oxide may be augmented by oxide formed in situ by wet chemistry, thermal or plasma oxidation or deposited as a thin film by the deposition methods previously mentioned.

Optical coupling layer. Another task that may be performed by one or more layers of the material system of this invention is coupling the incident electromagnetic radiation more effectively into the absorption layer. Several techniques can be used, such as those used modern solar cell devices, including but not limited to, optical impedance matching, anti-reflection coating, increasing the optical path length, and combinations thereof. Specific techniques demonstrated in Figs. 2 and 4 include using silicon dioxide for an anti reflection coating and to serve as an immobilization layer, and using nano-structured silicon to act as an optical impedance matching medium as well as an absorber. Materials that can be used for this layer, with appropriate properties, include metals, semiconductors, insulators, ceramics, polymers, organic materials, or composites thereof.

Substrate. For the material system of this invention, the substrate to which the layers are adhered onto may play one or more of the roles mentioned above or simply serve as a support media. The only necessary qualification of the substrate is that it must be compatible with subsequent processing. The substrate may also be pre-patterned for sample preparation and localization. Specific embodiments demonstrated in this invention use inexpensive acrylic, and polyimide plastics, glasses and metal foils as substrates. In general, materials that can be used for this layer include metals, semiconductors, insulators, ceramics, polymers, organic materials, or composites thereof.

Deposition methods.
Deposited films were preferably used to demonstrate this invention. Such deposited films can be deposited by a variety of methods including but not limited to the following: PVD such as sputtering and evaporation, CVD, PECVD, ECR-PECVD, PEPVD, electroplating, sol-gel, tape casting, self-assembly, liquid deposition, nebulization deposition, and spin coating. Specific techniques of the present invention include evaporation, sputtering, PECVD, and combinations thereof. Films or layers of the present invention are not limited to deposited films.

It is understood that certain films, i.e., polymer composite films, that are suitable for matrix-free photon desorption mass analysis devices of the present invention are not required to be deposited. For example, microcomposite or nano-composite polymers films presented herein may be used. They can be prepared by processing methods known in the art such as mixing organic polymer(s) and inorganic materials(s) in a solvent, and thereafter forming a composite film from the mixture. Such composite films may be prepared, for example, by molding, casting, spin casting, spraying, and any combinations thereof, or other procedures known to produce such polymer composites.

Device texturing and patterning. The texturing of one or more layers in the device of this invention can have several effects. Texturing of a reflective layer behind the absorber (on the opposite side of the device from the light source), increases the optical path length and enhances optical absorption in thin absorber layers. Texturing of the immobilization layer can have various effects depending on the length scale considered. As shown in Table 1 above, micro-scale roughness allows more analyte and desorption-ionization enhancing agents to be present per incident laser area. This can increase the sensitivity and longevity of analyte signal. On the nano-scale, the surface roughness will not only enhance signal sensitivity and longevity, but may also act to enhance light coupling
into the absorber by impedance matching. The nanoscale texturing also allows effective adsorption of analytes from the gas or liquid phase and provides better uniformity of the analyte distribution for more reproducible signal than achieved using other morphologies. Texturing can be done by a variety of manufacturing techniques including pre-fabricated or molded substrates, physical roughening, laser ablation, lithographic processes, and textured film growth. Methods of textured film growth of this invention include nano-structured PE-CVD growth conditions, zone growth model surface texturing, and glancing angle deposition.

Patterning of one or more of the layers in the device can serve many purposes. The localization of analytes, which is important for automation and sample delivery purposes, can be done on a macroscopic or microscopic scale with wells either pre-formed on the substrate or produced during subsequent film growth and processing. For instance wells could be hot embossed into a plastic substrate. Localization could also be attained by the plasma deposition of polymers or by the selective removal of an oxide layer. It could be attained by using "soft" lithographic patterning, such as PDMS stamping of molecules. Furthermore, patterning of the immobilization layer to causes differences in hydrophobicity, chemical affinity, acidity, charge and polarity can localize and preferentially bind desired analytes. Patterning of a metallic grid on the immobilization layer can remove charge buildup during the desorption/ionization process in machines requiring a grounded stage.

Finally, with the integration of these devices into micro-fluidic systems, patterning can be used to place the devices where needed. Patterning can also be done by a variety of manufacturing techniques including pre-fabricated or molded substrates, physical scribing, stamping, embossing, laser ablation and lithographic processes.
Integration with preparation, application, and analysis devices.
The deposited devices and mass analysis technique of this invention have
the unique ability to be integrated with a large number of sample delivery
and preparation techniques plus a large number of mass analyzers.
Preparation of the analyte can be as simple as placing a drop of the
molecules on the immobilization layer surface and allowing it to dry. It is
also possible to allow the analyte to adsorb to the immobilization layer
surface out of a gaseous or liquid solution. This simple gas or fluid
handling can be performed by a number of automated, high throughput
sampling systems. More complex schemes include the use of computer
integrated micro-fluidic, on chip, systems that perform chromatography or
purification. The deposited systems of this invention can be easily used in
tandem with a chip-based system or integrated into the micro-fluidic
device. The use of an integrated micro-fluidic system can also be used to
deliver desorption-ionization enhancing agents, such as water, to the
analyte during mass analysis in order to prolong detection signal.

Many mass spectroscopic methods can be used to analyze the
desorbed-ionized species. These may include but are not limited to: time
of flight, quadrupole, ion trap, plasmon resonance or combinations thereof.

Device structure and layer organization. The layer options and
requirements detailed above enable the morphology-tailored material
structures of this invention to be uniquely designed for optimal
performance based on the analytes, their sample preparation, the type of
electromagnetic source used for desorption-ionization, the mass analysis
technique used and integration techniques employed. However, there are
basic rules that apply to the overall device structure. First, the impinging
photons must be able to enter the absorber layer or layers of the device.
Second, the immobilization layer must enable the desorbed analytes to
enter the necessary mass detection area. Fig. 5 provides a variety of specific device structures unique to this invention.

Although the present invention describes in detail certain embodiments, it is understood that variations and modifications exist known to those skilled in the art that are within the invention. Accordingly, the present invention is intended to encompass all such alternatives, modifications and variations that are within the scope of the invention as set forth in the following claims.
WHAT IS CLAIMED IS:

1. An apparatus for providing an ionized analyte for mass analysis by photon desorption comprising:

   at least one layer for contacting an analyte; and

2. The apparatus of claim 1, further comprising one or more layers deposited on said substrate that act to absorb and convert photons to energy sufficient to desorb and ionize said analyte.

3. The apparatus of claim 1, wherein said substrate upon irradiation absorbs and converts photon energy to energy sufficient to desorb and ionize said analyte.

4. The apparatus of claim 1, wherein said substrate is selected from the group consisting of semiconductors, glasses, plastics, polymers, metals, ceramics, insulators, organic materials, inorganic materials, or any combinations thereof.

5. The apparatus of claim 2, wherein said one or more layers is selected from the group consisting of metals, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, and any combinations thereof.
6. The apparatus of claim 4, wherein said deposited layer enhances the absorption of photons by optical impedance matching, by acting as an anti-reflective coating, by increasing the photon path length, or any combinations thereof.

7. The apparatus of claim 1, wherein said deposited layer contacting said analyte is selected from the group consisting of silicon, silicon dioxide, germanium, germanium oxide, indium, gallium, cadmium, selenium, tellurium, and alloys and compounds thereof, carbon, hydrogen, semiconductors, insulators, metals, ceramics, polymers, other inorganic material, organic material, or any combinations thereof.

8. The apparatus of claim 1, wherein said layer is deposited by physical vapor deposition, chemical vapor deposition, liquid deposition, molecular beam epitaxy, plasma assisted chemical vapor deposition, sol-gels, nebulization, spraying, electroplating, tape casting, spin coating, assembly from liquid chemical precursors, printing, self-assembly and any combinations thereof.

9. The apparatus of claim 2, wherein said layer is deposited by physical vapor deposition, chemical vapor deposition, liquid deposition, molecular beam epitaxy, plasma assisted chemical vapor deposition, sol-gels, nebulization, spraying, electroplating, tape casting, spin coating, assembly from liquid chemical precursors, printing, self-assembly and any combinations thereof.

10. The apparatus of claim 1, wherein said deposited layer is a continuous film, a discontinuous film or any combinations thereof.
11. The apparatus of claim 1, wherein said layer contacting said analyte is physically or chemically modified, surface functionalized, or patterned.

12. The apparatus of claim 11, wherein the surface of said layer is chemically modified to control acid behavior, basic behavior, hydrophobicity, hydrophlicity, and any combinations thereof.

13. The apparatus of claim 1, wherein the thickness of said layer is essentially uniform from 5 nm to 10 microns.

14. The apparatus of claim 1, wherein said layer contacting an analyte is non-textured, micro-scale textured, nano-scale textured, or any combinations thereof.

15. The apparatus of claim 1, wherein the analyte is in an amount greater than 1 attomole.

16. The apparatus of claim 1, further comprising a micro-fluidic apparatus, a nano-fluidic apparatus, or combination thereof.

17. The apparatus of claim 1, further comprising a mass spectrometer for analysis of the mass of said analyte.

18. The apparatus of claim 17, wherein said mass analysis is by time of flight mass spectrometer, quadrapole mass spectrometer, ion trap device, or any combinations thereof.

19. The apparatus of claim 2, wherein one or more of said deposited layers is a continuous film, a discontinuous film, or any combinations thereof.
20. The apparatus of claim 1, wherein one or more of said contacting layers is physically or chemically modified, surface functionalized, or patterned.

21. The apparatus of claim 20, wherein the surface of said layer is chemically modified to control acid behavior, basic behavior, water content, hydrophobicity or hydrophylicity, and any combinations thereof.

22. The apparatus of claim 2, wherein the thickness of said layer is essentially uniform from 5 nm to 10 microns.

23. The apparatus of claim 1, wherein said layer contacting an analyte is non-textured, micro-scale textured, nano-scale textured, or any combinations thereof.

24. The apparatus of claim 1, wherein the analyte is in an amount less than 1 attomole.

25. The apparatus of claim 1, further comprising a micro-fluidic apparatus, a nano-fluidic apparatus, or combination thereof.

26. The apparatus of claim 1, further comprising a device for analysis of the mass of said analyte.

27. The apparatus of claim 26, wherein said device is a time of flight mass spectrometer, a quadrupole mass spectrometer, an ion trap device, or any combinations thereof.

28. A method for providing an ionized analyte for analysis of mass comprising:
providing an apparatus comprising at least one layer for contacting an analyte wherein said layer is deposited on a substrate;

contacting an amount of an analyte containing entities such as molecules whose mass or masses are to be determined with said deposited layer; and

irradiating said apparatus to desorb and ionize said analyte.

29. The method of claim 28, wherein said analyte is substantially free of a matrix.

30. The method of claim 28, wherein said analyte is selected from the group comprising organic chemical compositions, inorganic chemical compositions, biochemical compositions, cells, micro-organisms, peptides, polypeptides, proteins, lipids, carbohydrates, drug candidate molecules, drug molecules, drug metabolites, combinatorial chemistry products, nucleic acids, and any combinations thereof.

31. The method of claim 28, wherein said apparatus further comprises on or more layers deposited on said substrate that upon irradiating said apparatus absorb and convert photon energy sufficient to desorb and ionize said analyte.

32. The method of claim 28, wherein said substrate upon irradiation of said apparatus absorbs and converts photons to energy sufficient to desorb and ionize said analyte.

33. The method of claim 28, wherein said substrate is selected from the group consisting of semiconductors, glasses, plastics, polymers,
metals, ceramics, insulators, organic materials, inorganic materials, and any combinations thereof.

34. The method of claim 31, wherein said one or more deposited layers enhance the absorption of light by optical impedance matching, by acting as an anti-reflection coating, by increasing the photon path length, or by any combinations thereof.

35. The method of claim 31, wherein said one or more layers is selected from the group consisting of metals, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, and any combinations thereof.

36. The method of claim 28, wherein said deposited layer contacting said analyte is selected from the group consisting of silicon, silicon dioxide, germanium, germanium oxide, indium, gallium, cadmium, selenium, tellurium, and alloys and compounds thereof, carbon, hydrogen, semiconductors, insulators, metals, ceramics, polymers, other inorganic material, organic material, or any combinations thereof.

37. The method of claim 28, wherein said deposited layer of said apparatus is deposited by physical vapor deposition, chemical vapor deposition, liquid deposition, molecular beam epitaxy, plasma assisted chemical vapor deposition, sol-gels, nebulization, electroplating, tape casting, spin coating, assembly from liquid chemical precursors, printing, self-assembly, and any combinations thereof.

38. The method of claim 31, wherein said one or more layers is deposited by physical vapor deposition, chemical vapor deposition, liquid deposition, molecular beam epitaxy, plasma assisted chemical vapor deposition, sol-gels, nebulization, electroplating, tape casting, spin coating,
coating, assembly from liquid chemical precursors, printing, self-assembly, and any combinations thereof.

39. The method of claim 28, wherein said deposited layer of said apparatus contacting said analyte is a continuous film, a discontinuous film, or any combinations thereof.

40. The method of claim 28, wherein said deposited layer of said apparatus contacting said analyte is physically or chemically modified, surface functionalized, patterned, or any combinations thereof.

41. The method of claim 40, wherein said layer is chemically modified to control hydrophobicity or hydropilicity.

42. The method of claim 40, wherein said layer is chemically modified to control the surface pH of said layer.

43. The method of claim 28, wherein said layer contacting said analyte is non-textured, micro-scale textured, nano-scale textured, or any combinations thereof.

44. The method of claim 43, wherein said layer is textured by prefabricating textured substrates, physical roughening, laser ablation, lithographic processes, textured film growth, self-assembly deposition, or any combinations thereof.

45. The method of claim 28, wherein said analyte is in an amount less than 1 attomole.

46. The method of claim 28, wherein the thickness of said layer is essentially uniform from 5 nm to 10 microns.
47. The method of claim 28, further comprising adding an enhancing agent to said analyte prior to irradiating said apparatus.

48. The method of claim 47, wherein said enhancing agent is ammonium citrate, HCl, TFA, salts, hydrated molecules, surfactants, detergents, acids, bases, and any combinations thereof.

49. The method of claim 28, wherein said apparatus further comprises a micro-fluidic apparatus, a nano-fluidic apparatus, or combination thereof.

50. The method of claim 28, further comprising analyzing the mass of said ionized analyte by a device.

51. The method of claim 50, wherein said analyzing the mass of said ionized analyte is by time of flight mass spectroscopy, quadrapole mass spectroscopy, ion trap device, or any combinations thereof.

52. A method for determining a physical property of an analyte component comprising:

providing an apparatus comprising at least one layer for contacting an analyte and a substrate on which said layer is deposited;

positioning an amount of an analyte on the layer used for contacting an analyte of said apparatus;

irradiating said apparatus having said contacted analyte;

desorbing and ionizing at least one component of said analyte; and
analyzing said ionized at least one analyte component for a physical property.

53. The method of claim 52, wherein said physical property of said at least one analyte component is the mass to charge ratio (m/z) of the ionized analyte.

54. The method of claim 53, wherein said physical property is analyzed by mass spectrometry.

55. The method of claim 54, wherein said mass spectroscopy is time of flight, quadrupole, ion trap, or any combinations thereof.

56. The method of claim 52, wherein said analyte is substantially free of photon-absorbing matrix.

57. The method of claim 52, wherein said analyte is selected from the group consisting of organic chemical compositions, inorganic chemical compositions, biochemical compositions, cells, micro-organisms, peptides, polypeptides, proteins, lipids, carbohydrates, drug candidate molecules, drug molecules, drug metabolites, combinatorial chemistry products, nucleic acids, and any combinations thereof.

58. The method of claim 52, wherein said apparatus further comprises one or more layers deposited on said substrate that upon irradiating said apparatus absorb and convert photons to energy sufficient to desorb and ionize said analyte.
59. The method of claim 58, wherein said one or more layers is selected from the group consisting of metals, semiconductors, insulators, ceramics, polymers, organic materials, inorganic materials, and any combinations thereof.

60. The method of claim 58, wherein said one or more deposited layers enhances the absorption of light by photons impedance matching, by acting as an anti-reflection coating, by increasing the optical path length, or by any combinations thereof.

61. The method of claim 52, wherein said substrate upon irradiation of said apparatus absorbs and converts photons to energy sufficient to desorb and ionize said analyte.

62. The method of claim 52, wherein said substrate is selected from the group consisting of semiconductors, glasses, plastics, polymers, metals, ceramics, insulators, organic materials, inorganic materials, and any combinations thereof.

63. The method of claim 52, wherein said deposited layer contacting said analyte is selected from the group consisting of silicon, silicon dioxide, germanium, germanium oxide, indium, gallium, cadmium, selenium, tellurium and alloys and compounds thereof, carbon, hydrogen, semiconductors, insulators, ceramics, metals, polymers, other inorganic material, organic material, and any combinations thereof.

64. The method of claim 52, wherein said layer is deposited by physical vapor deposition, chemical vapor deposition, liquid deposition, molecular beam epitaxy, plasma assisted chemical vapor deposition, sol-gels, nebulization, electroplating, tape casting, spin coating, assembly
from liquid chemical precursors, printing, self-assembly, and any combinations thereof.

65. The method of claim 58, wherein said layer is deposited by physical vapor deposition, chemical vapor deposition, liquid deposition, molecular beam epitaxy, plasma assisted chemical vapor deposition, sol-gels, nebulization, electroplating, tape casting, spin coating, assembly from liquid chemical precursors, printing, self-assembly, and any combinations thereof.

66. The method of claim 52, wherein said deposited layer of said apparatus contacting said analyte is a continuous film, a discontinuous film, or any combinations thereof.

67. The method of claim 52, wherein said deposited layer of said apparatus contacting said analyte is physically or chemically modified, surface functionalized, patterned, or any combinations thereof.

68. The method of claim 67, wherein said layer is chemically modified to control hydrophobicity or hydrophilicity.

69. The method of claim 67, wherein said layer is chemically modified to control the surface pH of said layer.

70. The method of claim 52, wherein said layer contacting said analyte is non-textured, micro-scale textured, nano-scale textured, or any combinations thereof.

71. The method of claim 70, wherein said layer is textured by prefabricating textured substrates, physical roughening, laser ablation, lithographic processes, textured film growth, or any combinations thereof.
72. The method of claim 52, wherein said analyte is in an amount greater than 1 attomole.

73. The method of claim 52, wherein the thickness of said deposited layer is essentially uniform.

74. The method of claim 73, wherein said thickness of said deposited layer is from 5 nm to 10 microns.

75. The method of claim 52, further comprising adding an [ionizing] enhancing agent to said analyte prior to irradiating said apparatus.

76. The method of claim 75, wherein said enhancing agent is ammonium citrate, HCl, TFA, salts, hydrated molecules, surfactants, chelating agents, detergents, acids, bases, and any combinations thereof.

77. The method of claim 52, wherein said apparatus further comprises a micro-fluidic apparatus, a nano-fluidic apparatus, or combinations thereof.

78. The method of claim 52, further comprising analyzing the masses of said one or more components of said ionized analyte by a device.

79. The method of claim 78, wherein analyzing the mass is by time of flight mass spectroscopy, quadrupole mass spectroscopy, ion trap device, or any combinations thereof.
80. An apparatus for determining the masses of one or more components of an analyte comprising a substrate, an analyte contacting the substrate, a source of radiation irradiating the substrate wherein illumination of the substrate causes the desorption and ionization of the analyte, a source of positive or negative voltage connected to the substrate that controls and directs the ionized analyte, and a spectrometer that analyzes the mass to charge ratio (m/z) of the ionized analyte components wherein the improvement comprises:

a substrate that is an apparatus comprising at least one layer for contacting an analyte deposited on a substrate material wherein said apparatus has optical properties to absorb and convert photons to energy sufficient to desorb and ionize said analyte.

81. An apparatus according to claim 80, wherein the analyte is substantially free of photon absorbing matrix.

82. An apparatus according to claim 80, wherein one or more of said layers of said apparatus is a continuous film, a discontinuous film, or any combinations thereof.

83. A method of improving the detection of an analyte by laser desorption mass spectrometry comprising the steps of:

providing a substrate having a fluorinated coated sample loading region;

providing an analyte dissolved in a first liquid as a sample; and
contacting the coated sample loading region with the sample wherein the sample does not spread on the coated sample loading regions to form a sample loaded substrate.

84. A device of claim 1 where the deposited layer is a composite material comprising an organic material and a photon adsorbing micro or nanoparticle.

85. A device of claim 84 where the organic material is polymer.

86. A device of claim 85 where the polymer is a halogenated material.

87. A device of claim 86 where the polymer is an acid.

88. A device of claim 85 wherein the polymer is a fluorinated/sulfur containing material.

89. A device of claim 84 where the photon adsorbing micro or nanoparticle is a semiconductor.

90. A device of claim 84 where the photon adsorbing micro or nanoparticle is a metal, organic, insulator or inorganic material.

91. A device of claim 84 where the photon adsorbing micro or nanoparticle is carbon.
FIG. 1
FIG. 2
FIG. 4
# INTERNATIONAL SEARCH REPORT

## A. CLASSIFICATION OF SUBJECT MATTER

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<td>US CL</td>
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According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S.: 250/288, 281, 282; 422/68.1, 81; 436/173, 174

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

NONE

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

NONE

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tr>
<td>X</td>
<td>US 5,719,060 A (HUTCHENS et al) 17 February 1998 (17.02.1998), col. 12, lines 41-55.</td>
<td>1-5, 7-33, 35-59, 61-91</td>
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<tr>
<td>Y</td>
<td>USR 5,600,351 A (COTTRELL et al) 09 November 1993 (09.11.1993), col. 1, line 65 - col. 2, line 19.</td>
<td>6, 34-60</td>
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<tr>
<td>Y</td>
<td>US 5,552,277 A (BOGART) 03 September 1996 (03.09.1996), col. 10, lines 60-67; col. 4, lines 10-15; col. 5, lines 5-10; col. 11, lines 9-24.</td>
<td>1, 28, 52, 80, 83</td>
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Further documents are listed in the continuation of Box C.

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<tr>
<td>Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231</td>
<td>KALIMAH FERNANDEZ</td>
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<td>Facsimile No. (703) 305-3230</td>
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</tr>
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Date of the actual completion of the international search: 08 July 2002 (08.07.2002)

Date of mailing of the international search report: 16 SEP 2002

Form PCT/ISA/210 (second sheet) (July 1998)