



US010358715B2

(12) **United States Patent**  
**Kaufman-Osborn et al.**

(10) **Patent No.:** **US 10,358,715 B2**  
(45) **Date of Patent:** **Jul. 23, 2019**

(54) **INTEGRATED CLUSTER TOOL FOR SELECTIVE AREA DEPOSITION**

(58) **Field of Classification Search**

None  
See application file for complete search history.

(71) Applicant: **Applied Materials, Inc.**, Santa Clara, CA (US)

(56) **References Cited**

(72) Inventors: **Tobin Kaufman-Osborn**, Sunnyvale, CA (US); **Srinivas D. Nemani**, Sunnyvale, CA (US); **Ludovic Godet**, Sunnyvale, CA (US); **Qiwei Liang**, Fremont, CA (US); **Adib Khan**, Santa Clara, CA (US)

U.S. PATENT DOCUMENTS

6,200,893 B1 3/2001 Sneh  
6,391,785 B1 5/2002 Satta et al.  
(Continued)

(73) Assignee: **APPLIED MATERIALS, INC.**, Santa Clara, CA (US)

FOREIGN PATENT DOCUMENTS

KR 10-2010-0107228 10/2010  
WO 2011103062 A2 8/2011

(\*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.

OTHER PUBLICATIONS

T. M. Mayer, et al., Chemical Vapor Deposition of Fluoroalkylsilane Monolayer Films; for Adhesion Control in Microelectromechanical Systems, *Journal of Vacuum Science & Technology B*, 18 (5), Sep./Oct. 2000, 9 pp.

(21) Appl. No.: **15/173,356**

(Continued)

(22) Filed: **Jun. 3, 2016**

*Primary Examiner* — Jethro M. Pence

(65) **Prior Publication Data**

US 2017/0350004 A1 Dec. 7, 2017

(74) *Attorney, Agent, or Firm* — Patterson + Sheridan LLP

(51) **Int. Cl.**  
**C23C 16/04** (2006.01)  
**H01J 37/32** (2006.01)

(57) **ABSTRACT**

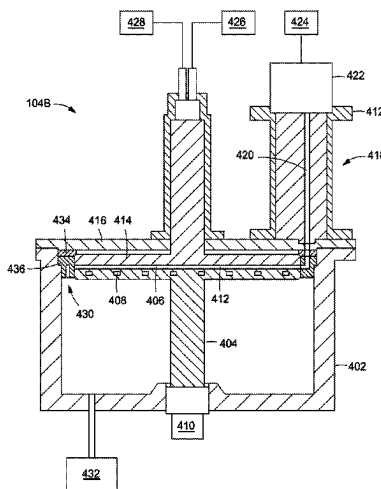
(Continued)

Embodiments described herein relate to apparatus and methods for processing a substrate. In one embodiment, a cluster tool apparatus is provided having a transfer chamber and a pre-clean chamber, a self-assembled monolayer (SAM) deposition chamber, an atomic layer deposition (ALD) chamber, and a post-processing chamber disposed about the transfer chamber. A substrate may be processed by the cluster tool and transferred between the pre-clean chamber, the SAM deposition chamber, the ALD chamber, and the post-processing chamber. Transfer of the substrate between each of the chambers may be facilitated by the transfer chamber which houses a transfer robot.

(52) **U.S. Cl.**  
CPC ..... **C23C 16/042** (2013.01); **C23C 16/02** (2013.01); **C23C 16/45502** (2013.01); **C23C 16/45544** (2013.01); **C23C 16/45561** (2013.01); **C23C 16/45582** (2013.01); **C23C 16/54** (2013.01); **C23C 16/56** (2013.01); **H01J 37/32357** (2013.01); **H01J 37/32724** (2013.01); **H01J 37/32733** (2013.01); **H01J 37/32899** (2013.01);

(Continued)

**17 Claims, 8 Drawing Sheets**



(51) **Int. Cl.**

**H01L 21/67** (2006.01)  
**H01L 21/677** (2006.01)  
**H01L 21/8234** (2006.01)  
**H01L 21/687** (2006.01)  
**C23C 16/02** (2006.01)  
**C23C 16/455** (2006.01)  
**C23C 16/54** (2006.01)  
**C23C 16/56** (2006.01)

(52) **U.S. Cl.**

CPC .... **H01L 21/6719** (2013.01); **H01L 21/67069**  
(2013.01); **H01L 21/67103** (2013.01); **H01L**  
**21/67109** (2013.01); **H01L 21/67167**  
(2013.01); **H01L 21/67196** (2013.01); **H01L**  
**21/67207** (2013.01); **H01L 21/67742**  
(2013.01); **H01L 21/67748** (2013.01); **H01L**  
**21/67754** (2013.01); **H01L 21/68785**  
(2013.01); **H01J 2237/327** (2013.01); **H01J**  
**2237/334** (2013.01); **H01L 21/823431**  
(2013.01)

## (56)

**References Cited**

## U.S. PATENT DOCUMENTS

7,071,022	B2	7/2006	Couillard et al.	
7,648,866	B2	1/2010	Jung et al.	
7,775,508	B2	8/2010	Choi et al.	
7,960,295	B2	6/2011	Park	
9,112,003	B2	8/2015	Haukka et al.	
9,502,289	B2	11/2016	Haukka et al.	
2008/0026146	A1	1/2008	Kobrin et al.	
2008/0044569	A1	2/2008	Myo et al.	
2009/0004850	A1*	1/2009	Ganguli .....	C23C 16/18 438/655
2009/0057266	A1	3/2009	Tuncel et al.	
2011/0124187	A1	5/2011	Afzali-Ardakani et al.	
2012/0178650	A1	7/2012	Carroll et al.	
2012/0238109	A1	9/2012	Hattori et al.	
2013/0193108	A1	8/2013	Zheng et al.	
2014/0357016	A1	12/2014	Fuse et al.	
2014/0370718	A1	12/2014	Chakrapani et al.	
2015/0090330	A1	4/2015	Sewell et al.	
2015/0147879	A1*	5/2015	Chatterjee .....	H01L 21/76834 438/653
2015/0179500	A1	6/2015	Boussie et al.	
2016/0056071	A1	2/2016	Draeger et al.	
2016/0104648	A1	4/2016	Park et al.	

## OTHER PUBLICATIONS

Elina Farm, et al., Self-Assembled Octadecyltrimethoxysilane Monolayers Enabling Selective-Area Atomic Layer Deposition of Iridium, Chemical Vapor Deposition, 12, 2006, 3 pp.  
Junsic Hong, et al., ALD Resist Formed by Vapor-Deposited Self-Assembled Monolayers, Langmuir 2007, 23, 6 pp.

Satu Ek, et al., Atomic Layer Deposition of a High-Density Aminopropylsiloxane Network on Silica through Sequential Reactions of ?-Aminopropyltrialkoxysilanes and Water, Langmuir 2003, 19, 9 pp.  
Hashemi et al, Self Correcting Process for High Quality Patterning by Atomic Layer Deposition, American Chemical Society, Jul. 16, 2015.  
Elina Farm, Selective-Area Atomic Layer Deposition, University of Helsinki, Nov. 2011.  
International Search Report and Written Opinion for International Application No. PCT/US2017/019982 (023712PCT) dated Jun. 26, 2017.  
SE Kirkwood et al., Mechanism for femtosecond laser pulse patterning of SAM on gold coated substrate, IOP Publishing Ltd 2007 p. 428-430.  
Ming Fang, et al., Area-Selective Atomic Layer Deposition: Conformal Coating, Subnanometer Thickness Control and Smart Positioning, ACS Nano 2015 Vo. 9. No. 9 pp. 8651-8654.  
Non-final Office Action in U.S. Appl. No. 10/731,656 dated May 11, 2004.  
European Office Action in application EP 00955875.0. dated May 8, 2004.  
JW Klaus et al, Atomically controlled growth of tungsten and tungsten nitride using sequential surface reactions; Applied Surface Sciences 162-163 (2000) 479-491.  
Mikko Ritala et al., Atomic layer epitaxy—a valuable tool for nanotechnology?; Nanotechnology 10 (1999) 19-24 printed in the UK.  
Per Martensson et al, Atomic Layer Epitaxy of Copper, J. Elect rochem. Soc., vol. 145, No. 8, Aug. 1998 The Electrochemical Society, Inc. pp. 2926-2931.  
Shin Yokoyama, et al., Atomic-layer selective deposition of silicon nitride on hydrogen-terminated Si surfaces, Applied Surface Science 130-132 1998. 352-356.  
Stephen M. Gates, Surface Chemistry in the Chemical Vapor Deposition of Electronic Materials, Chem. Rev. 1996, 96, pp. 1519-1532.  
A. Jain et al., Control of selectivity during chemical vapor deposition of copper from copper (I) compounds via silicon dioxide surface modification, Applied Physics Letters 61, pp. 2662-2664 (1992) American Institute of Physics.  
M. Leskela et al., ALD precursor chemistry: Evolution and future challenges, Article published online by EDP Sciences and available at <http://dx.doi.org/10.1051/jp4:19998106>.  
Jae-Sik Min et al., Metal-organic atomic-layer deposition of titanium-silicon-nitride films. Applied Physics Letters vol. 75, No. 11 Sep. 13, 1999.  
Mikko Ritala et al., Perfectly Conformal TiN and Al2O3 Films Deposited by Atomic Layer Deposition, Chemical Vapor Deposition 1999, 5, No. 1 Ó Wiley-VCH Verlag GmbH, D-69469 Weinheim, 1999.  
International Search Report and Written Opinion of related application PCT/US2017/024077 dated Jul. 11, 2017.  
Woo-Hee Kim, et al., A Process for Topographically-Selective Deposition on 3D Nanostructures by Ion Implantation, ACS Nano, 2016, 10 (4), 22 pp.

\* cited by examiner

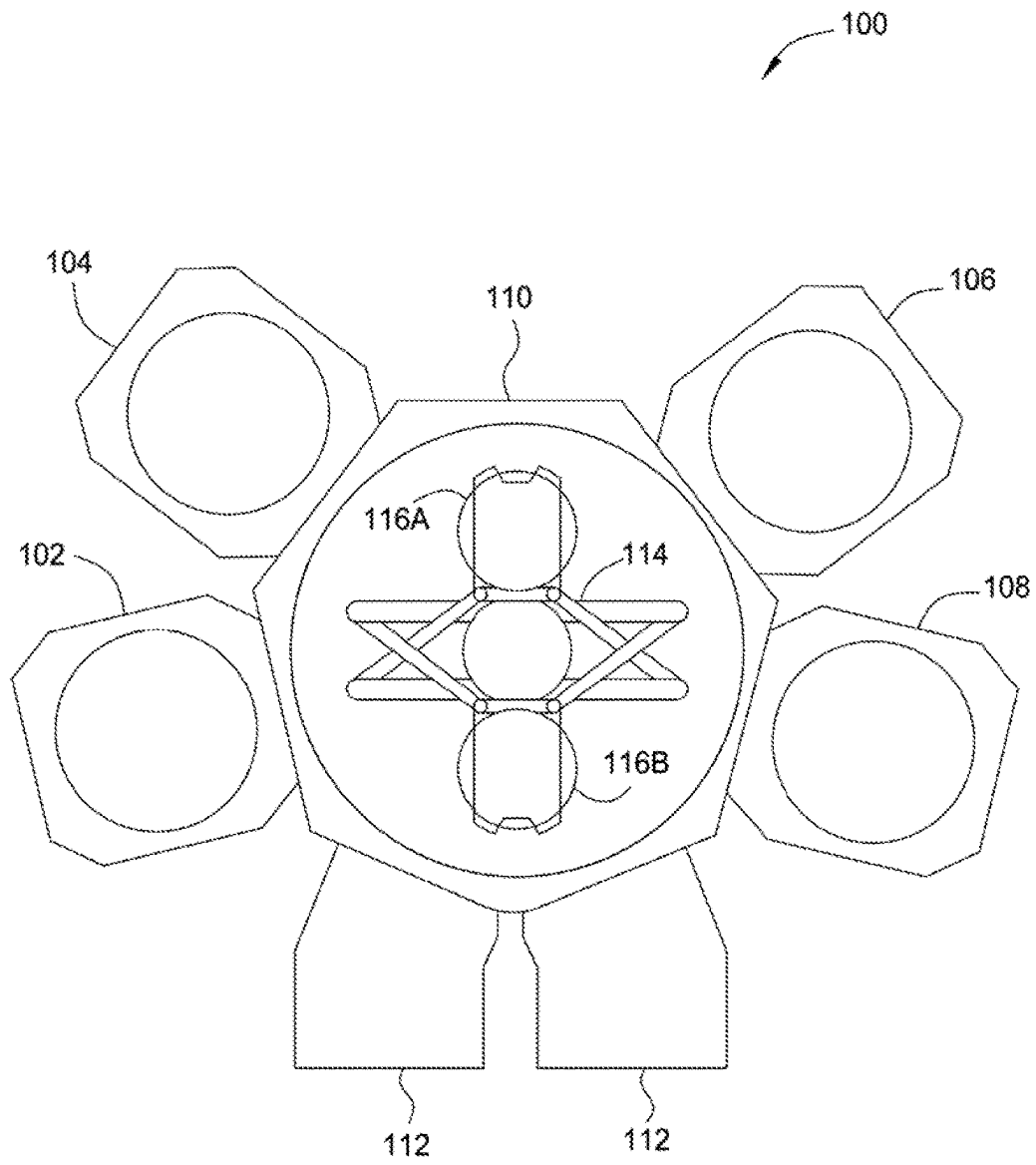


FIG. 1

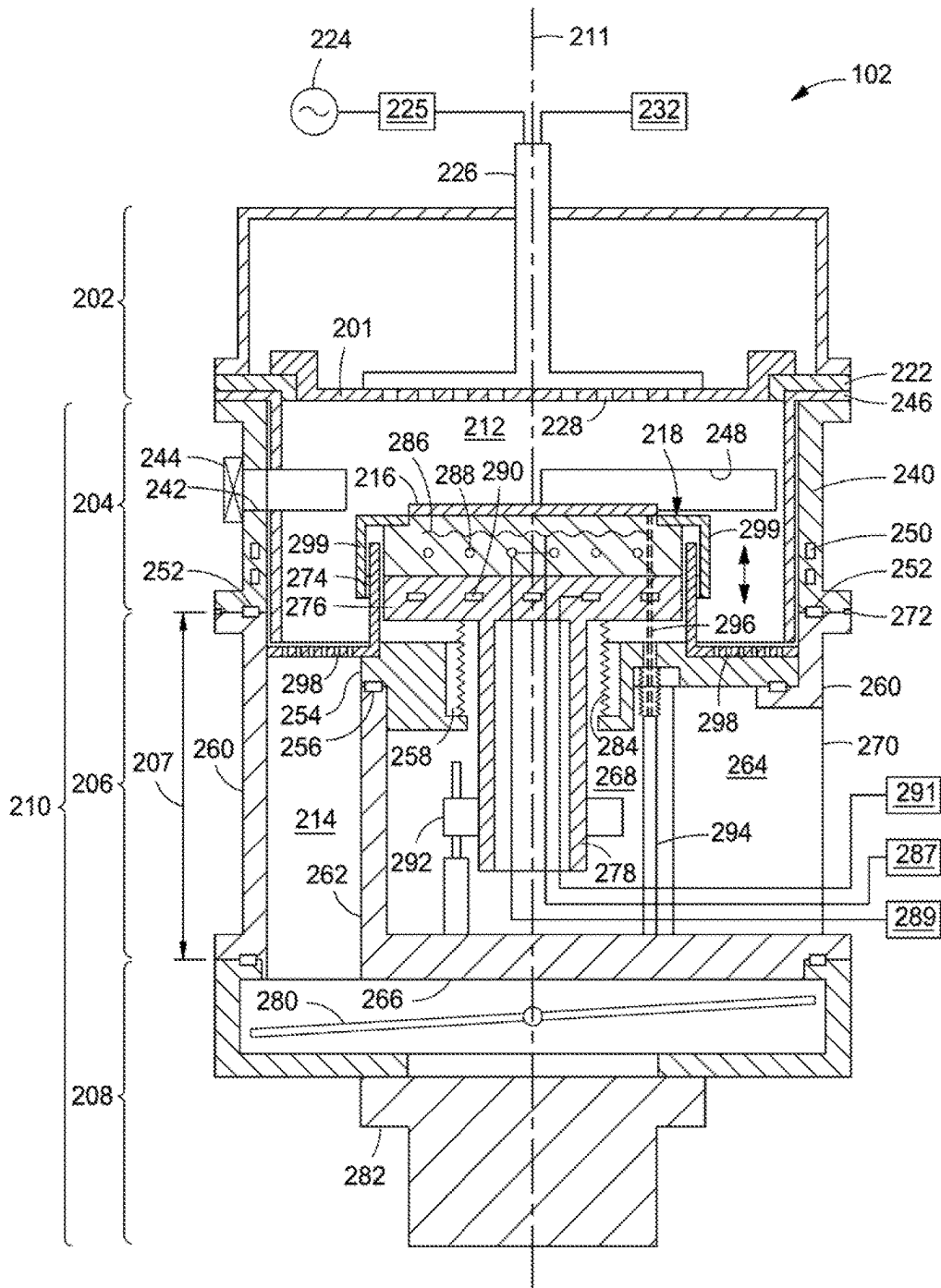


FIG. 2

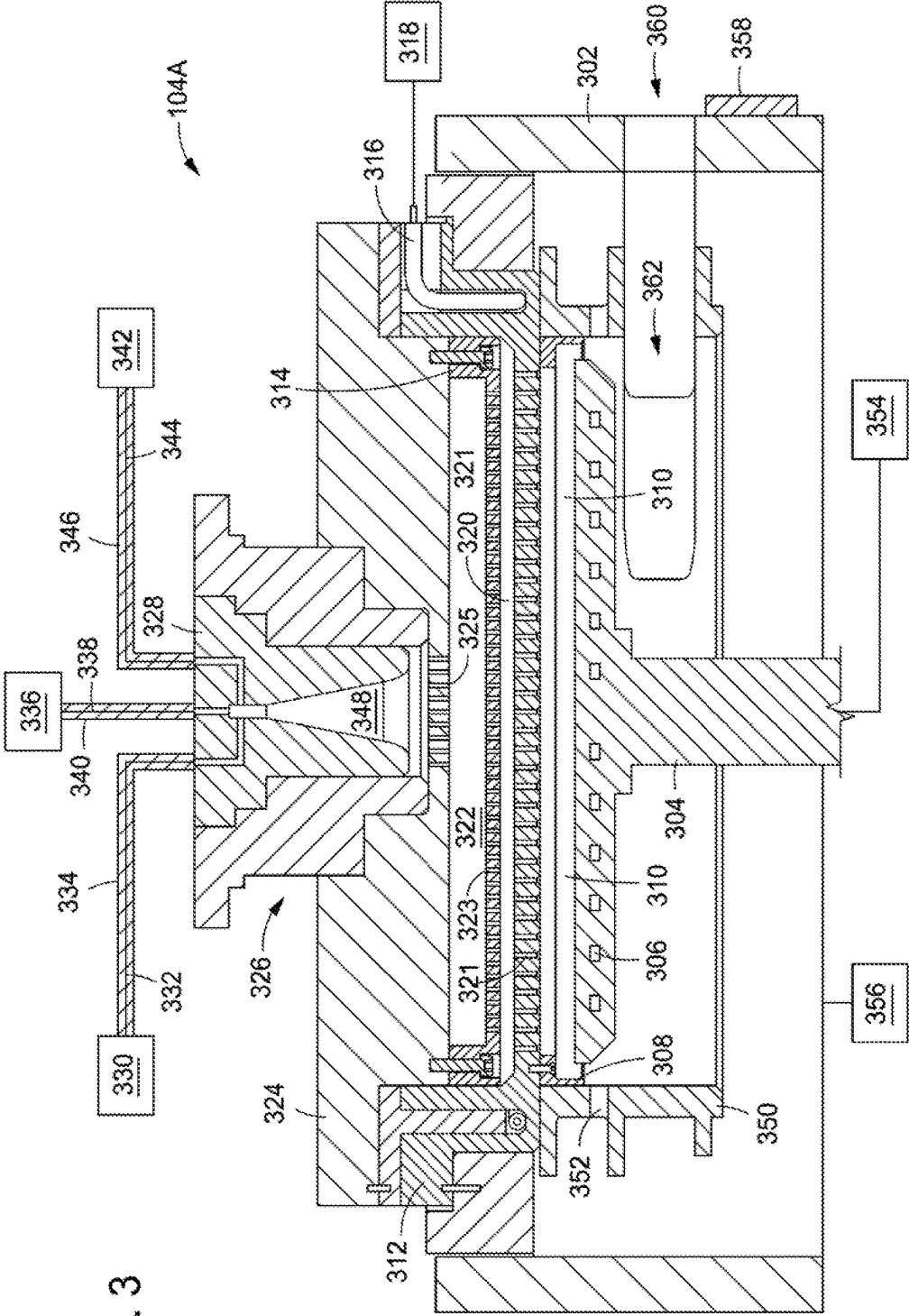


FIG. 3

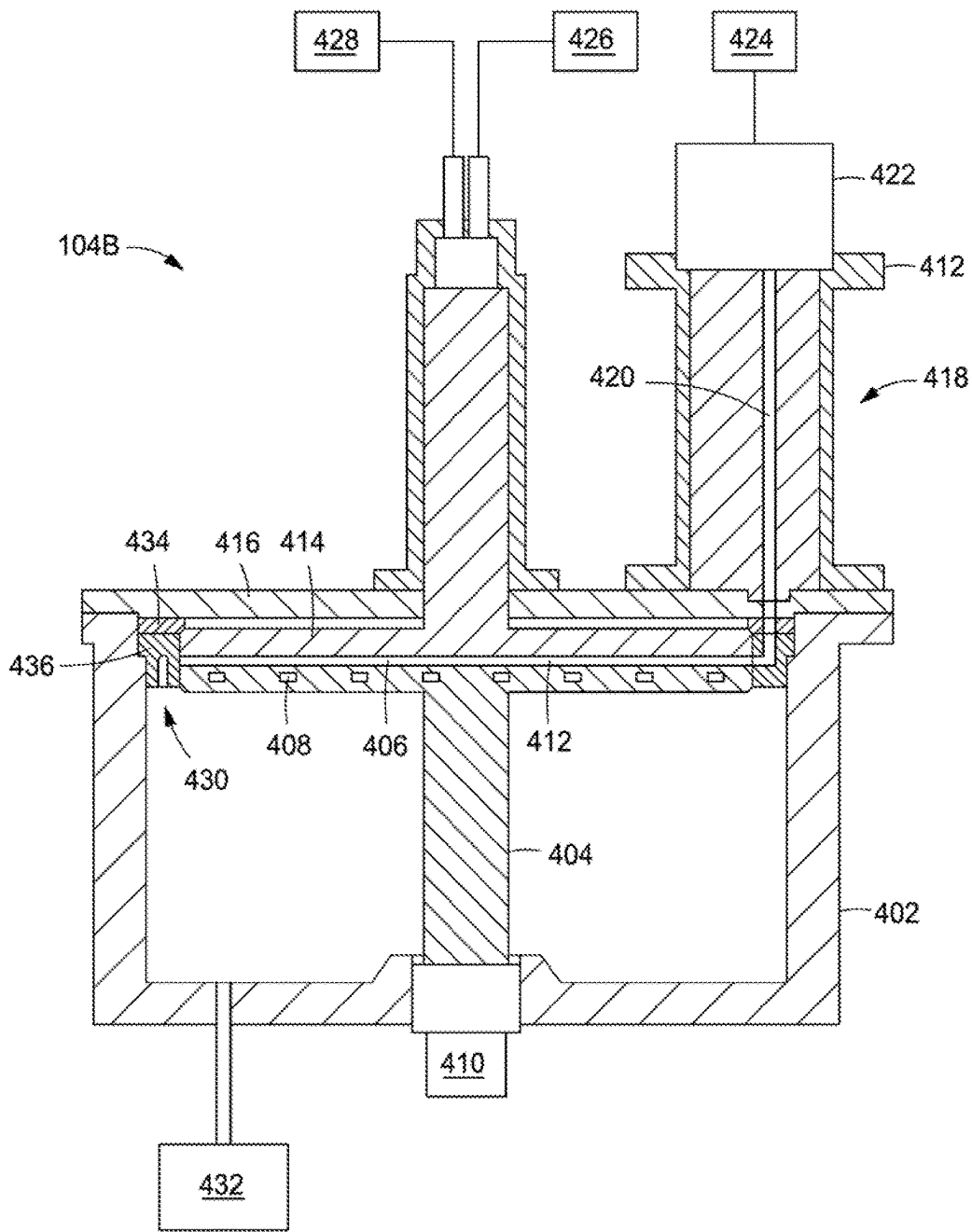


FIG. 4

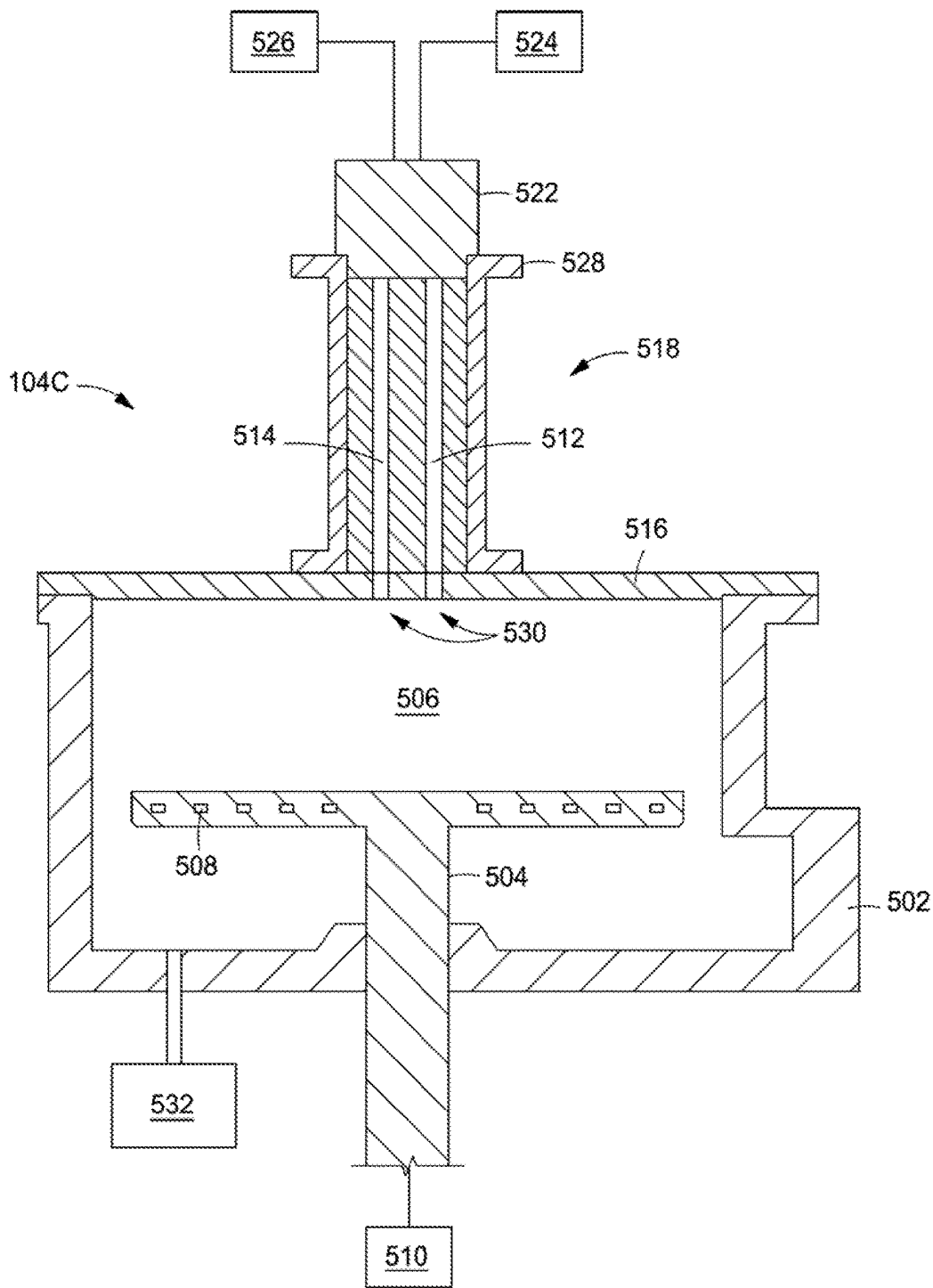


FIG. 5

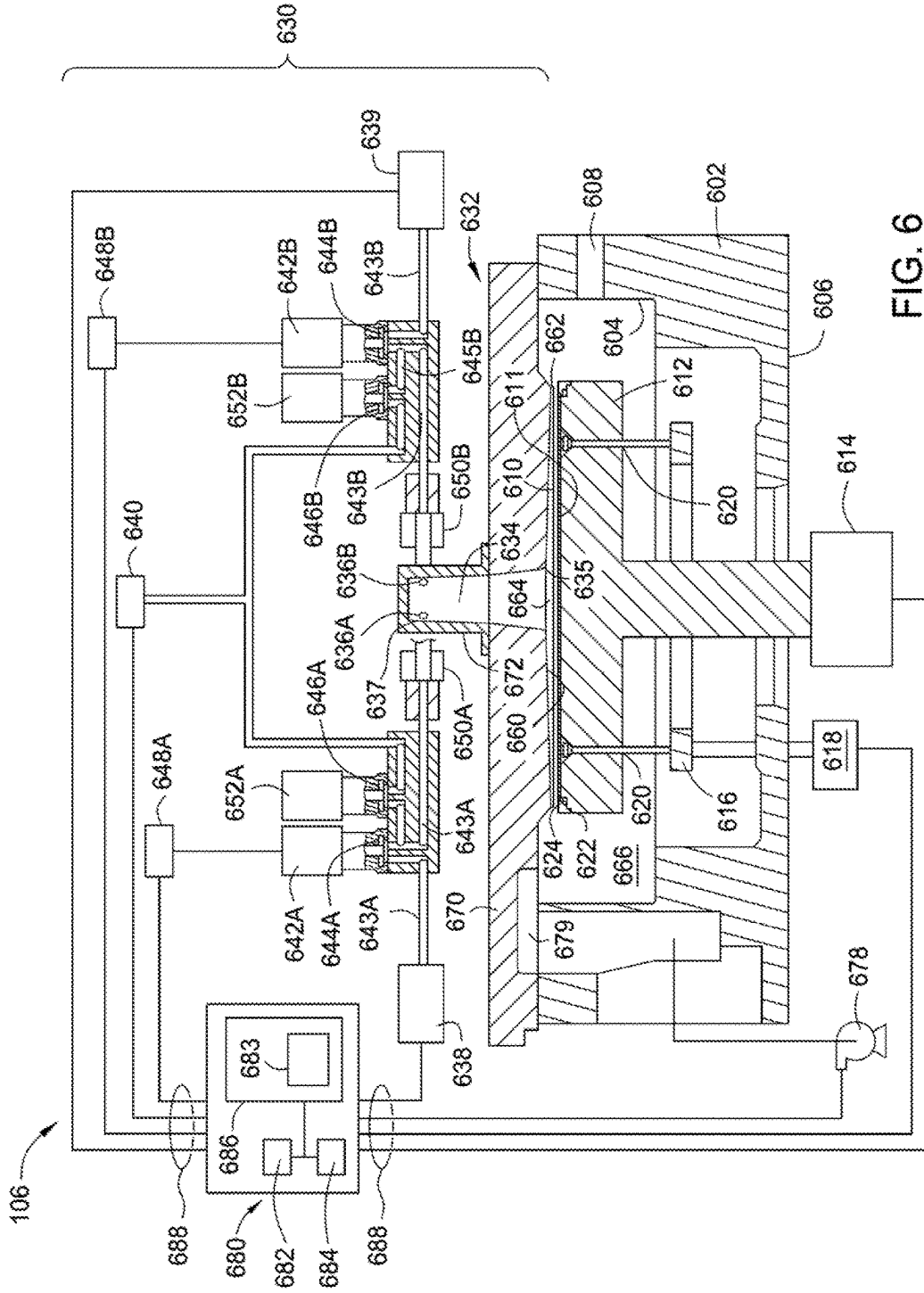


FIG. 6



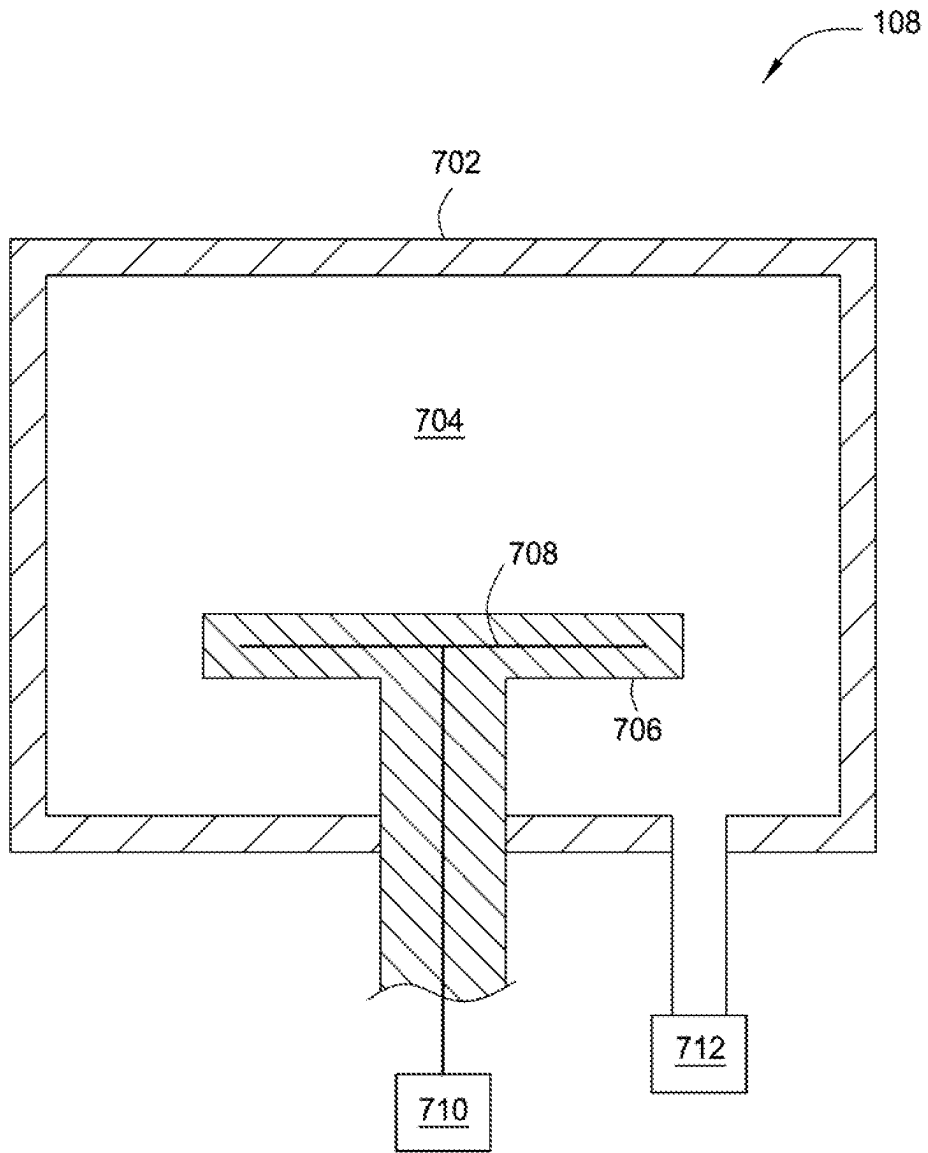


FIG. 7

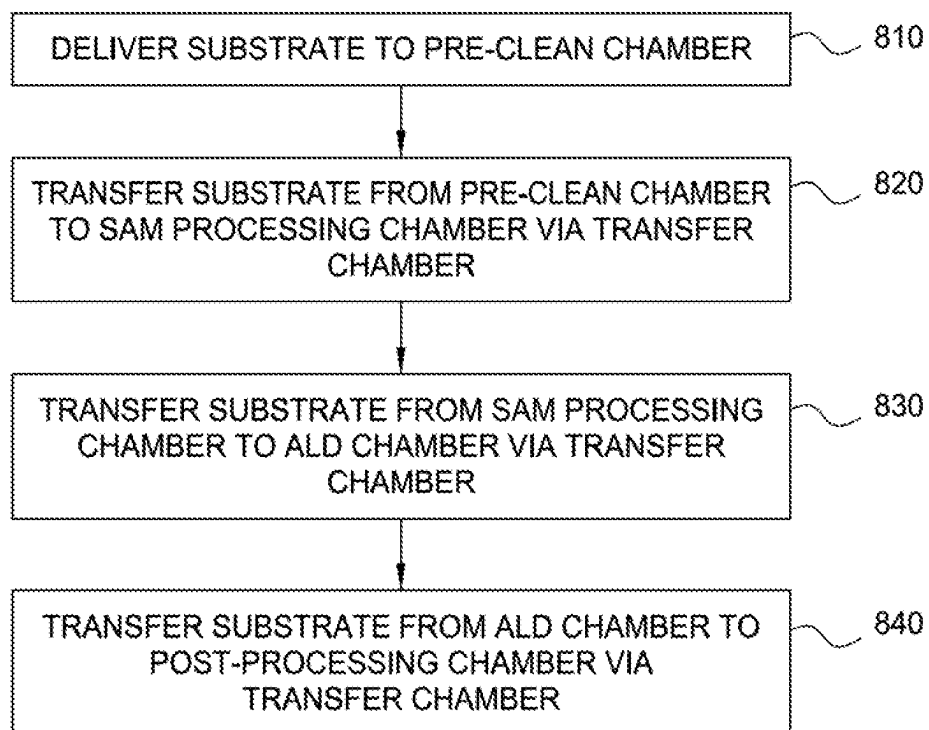


FIG. 8

1

# INTEGRATED CLUSTER TOOL FOR SELECTIVE AREA DEPOSITION

## BACKGROUND

### Field

Embodiments of the present disclosure generally relate to apparatus for processing substrates. More specifically, embodiments described herein relate to an integrated cluster tool for selective area deposition.

### Description of the Related Art

Reliably producing sub-half micron and smaller features is one of the key technology challenges for next generation very large scale integration (VLSI) and ultra large scale integration (ULSI) of semiconductor devices. However, as the limits of circuit technology are pushed, the shrinking dimensions of VLSI and ULSI technology have placed additional demands on processing capabilities.

As circuit densities increase for next generation devices, the widths of interconnects, such as vias, trenches, contacts, gate structures and other features, as well as the dielectric materials therebetween, decrease to 45 nm and 32 nm dimensions and beyond. In order to enable the fabrication of next generation devices and structures, three dimensional (3D) stacking of features in semiconductor chips is often utilized. In particular, fin field effect transistors (FinFETs) are often utilized to form three dimensional (3D) structures in semiconductor chips. By arranging transistors in three dimensions instead of conventional two dimensions, multiple transistors may be placed in the integrated circuits (ICs) very close to each other. As circuit densities and stacking increase, the ability to selectively deposit subsequent materials on previously deposited materials gains importance.

Self-assembled monolayers (SAMs) may be utilized as a masking material to improve subsequent material deposition selectivity. SAMs are generally surface chemistry dependent and can be formed preferentially on various materials. However, current apparatus for depositing SAMs often suffers from slow deposition rates which undesirably decreases throughput. In addition, SAM apparatus integration with pre and post processing apparatus which provides suitable throughput is lacking in the industry.

Thus, what is needed in the art are improved substrate processing apparatus and methods.

### SUMMARY

In one embodiment, a substrate processing apparatus is provided. The apparatus includes a platform having a centrally disposed transfer chamber and a first process chamber coupled to the transfer chamber. The first process chamber is configured to perform an etching process. A second process chamber is coupled to the transfer chamber and the second process chamber is configured to perform a self-assembled monolayer treatment process. A third process chamber is coupled to the transfer chamber and the third process chamber is configured to perform an atomic layer deposition process. A fourth process chamber is coupled to the transfer chamber and the fourth process chamber is configured to perform an anneal process.

In another embodiment, a substrate processing apparatus is provided. The apparatus includes a platform having a centrally disposed transfer chamber and a first process chamber coupled to the transfer chamber. The first process

2

chamber is configured to perform an etching process. A second process chamber is coupled to the transfer chamber adjacent the first process chamber and the second process chamber is configured to perform a self-assembled monolayer treatment process. A third process chamber is coupled to the transfer chamber adjacent the second process chamber and the third process chamber is configured to perform an atomic layer deposition process. A fourth process chamber is coupled to the transfer chamber adjacent the third process chamber and the fourth process chamber is configured to perform an anneal process.

In yet another embodiment, a substrate processing method is provided. The method includes transferring a substrate to a first process chamber and performing an etching process on the substrate in the first process chamber. The substrate is transferred from the first process chamber to a second process chamber via a transfer chamber and a self-assembled monolayer treatment process is performed on the substrate in the second process chamber. The substrate is transferred from the second process chamber to a third process chamber via the transfer chamber and an atomic layer deposition process is performed on the substrate in the third process chamber. The substrate is transferred from the third process chamber to a fourth process chamber via the transfer chamber and an anneal process is performed on the substrate in the fourth process chamber.

### BRIEF DESCRIPTION OF THE DRAWINGS

So that the manner in which the above recited features of the present disclosure can be understood in detail, a more particular description of the disclosure, briefly summarized above, may be had by reference to embodiments, some of which are illustrated in the appended drawings. It is to be noted, however, that the appended drawings illustrate only exemplary embodiments and are therefore not to be considered limiting of its scope, may admit to other equally effective embodiments.

FIG. 1 illustrates a schematic, plan view of a cluster tool apparatus according to one embodiment described herein.

FIG. 2 illustrates a schematic, cross-sectional view of a process chamber according to one embodiment described herein.

FIG. 3 illustrates a schematic, cross-sectional view of a process chamber according to one embodiment described herein.

FIG. 4 illustrates a schematic, cross-sectional view of a process chamber according to one embodiment described herein.

FIG. 5 illustrates a schematic, cross-sectional view of a process chamber according to one embodiment described herein.

FIG. 6 illustrates a schematic, cross-sectional view of a process chamber according to one embodiment described herein.

FIG. 7 illustrates a schematic, cross-sectional view of a process chamber according to one embodiment described herein.

FIG. 8 illustrates operations of a method for processing a substrate according to one embodiment described herein.

To facilitate understanding, identical reference numerals have been used, where possible, to designate identical elements that are common to the figures. It is contemplated that elements and features of one embodiment may be beneficially incorporated in other embodiments without further recitation.

Embodiments described herein relate to apparatus and methods for processing a substrate. In one embodiment, a cluster tool apparatus is provided having a transfer chamber and a pre-clean chamber, a self-assembled monolayer (SAM) deposition chamber, an atomic layer deposition (ALD) chamber, and a post-processing chamber disposed about the transfer chamber. A substrate may be processed by the cluster tool and transferred between the pre-clean chamber, the SAM deposition chamber, the ALD chamber, and the post-processing chamber. Transfer of the substrate between each of the chambers may be facilitated by the transfer chamber which houses a transfer robot.

As utilized herein, "self-assembled monolayer" (SAM) generally refers to a layer of molecules that are attached (e.g., by a chemical bond) to a surface and that have adopted a preferred orientation with respect to that surface and even with respect to each other. The SAM typically includes an organized layer of amphiphilic molecules in which one end of the molecule, the "head group" shows a specific, reversible affinity for a substrate. Selection of the head group will depend on the application of the SAM, with the type of SAM compounds based on the substrate utilized. Generally, the head group is connected to an alkyl chain in which a tail or "terminal end" can be functionalized, for example, to vary wetting and interfacial properties. The molecules that form the SAM will selectively attach to one material over another material (e.g., metal vs. dielectric) and if of sufficient density, can successfully operation subsequent deposition allowing for selective deposition on materials not coated with the SAM.

FIG. 1 illustrates a schematic, plan view of a cluster tool apparatus **100** according to one embodiment described herein. Examples of suitable apparatus which may be utilized in accordance with the embodiments described herein include the CENTURA® and ENDURA® platforms, both of which are available from Applied Materials, Inc., Santa Clara, Calif. It is contemplated that other suitably configured apparatus from other manufacturers may also be advantageously utilized in accordance with the embodiments described herein. In addition, the PRODUCER® platform, also available from Applied Materials, Inc., Santa Clara, Calif., having dual-chamber capability may be advantageously employed according to the embodiments described herein. In embodiments utilizing the PRODUCER® platform, load lock chambers with bake or anneal capability may be employed as post processing chambers in accordance with the embodiments described hereinafter.

Generally, the apparatus **100** includes a plurality of process chambers **102**, **104**, **106**, **108**, a transfer chamber **110**, and load lock chambers **112**. Each of the process chambers **102**, **104**, **106**, **108** are coupled to the transfer chamber **110**. In one embodiment, the process chamber **104** is disposed adjacent the process chamber **102**. In one embodiment, the process chamber **106** is disposed adjacent the process chamber **104**. In one embodiment, the process chamber **108** is disposed adjacent the process chamber **106**. Each process chamber represents, and may be used for, a different stage or phase of substrate processing. In one embodiment, the process chamber **102** is a pre-clean chamber. The process chamber **102** prepares surfaces of a substrate being processed for subsequent processing. In various examples, the process chamber **102** may remove substrate defects which result from air exposure, remove native oxide layers, and/or remove sacrificial layers disposed on a surface of the substrate to be treated by SAM or ALD processing. In another

example, the process chamber **102** is utilized for substrate surface functionalization. In this example, surface terminal groups may be modified to enable, assist, or prevent the formation of a SAM on the substrate, depending upon the desired implementation.

Specific examples of surface treatment which may be performed by the process chamber **102** include metal oxide removal via plasma treatment, surface hydroxyl functionalization using H<sub>2</sub>/O<sub>2</sub> plasma treatment or water vapor exposure, and/or oxide removal using a SICONI® process. The SICONI® process is available from Applied Materials, Inc., Santa Clara, Calif. It is contemplated that other similar treatment processes may be utilized in accordance with the embodiments described herein. More specifically, the process chamber **102** is utilized to enable selective area SAM adsorption. For example, an octadecyltrichlorosilane (ODTCS) SAM may bond to a dielectric or metal oxide material preferentially to a metal or Si—H terminated surface, assuming desirable conditions are present. The process chamber **102** is utilized to remove the metal oxide or native oxide to form an exposed metal surface of Si—H terminated surface which generally prohibits SAM adsorption.

In another embodiment, the process chamber **102** may be utilized as a post-process chamber. The process chamber **102**, which is described in greater detail with regard to FIG. 2, includes a heated pedestal which is utilized to perform a bake or anneal process. The bake or anneal process may be utilized after the substrate has been processed by other chambers of the apparatus **100**.

In one embodiment, the process chamber **104** is a SAM deposition chamber. The process chamber **104** is configured to enable vapor phase SAM molecules to selectively adsorb to one material of a substrate preferentially to another material of the substrate. The selective adsorption is generally controlled by the reactivity of the SAM molecule headgroup and the surface termination characteristics/functional groups disposed on the substrate surface. For example, a substrate having exposed SiO<sub>2</sub> and Cu materials which are exposed to the same SAM treatment process will result in the SAM molecules selective to metals bonding to the Cu preferentially and substantially no adsorption on the SiO<sub>2</sub> material. The resulting SAM material has a high water contact angle (i.e. greater than about 105°) which indicates the formation of a dense SAM. Thus, the process chamber **104** is utilized to perform SAM deposition processes.

In one embodiment, the process chamber **106** is an ALD chamber. The process chamber **106** is configured to enable deposition on surfaces of the substrate not covered by the SAM materials. For example, ALD materials generally do not form on surfaces which have a water contact angle greater than about 105°, such as greater than about 110°. Accordingly, the ALD process may be selectively deposited on a desired material of the substrate by utilizing the SAM material to improve the selectivity of deposition.

In one embodiment, the process chamber **108** is a post-process chamber, such as an anneal or bake chamber. In another embodiment, the process chamber **108** is a plasma processing chamber. In one example, the process chamber **108** enables SAM material removal from the substrate by thermal desorption or exposure to a plasma, such as an oxygen and/or hydrogen containing plasma. Alternatively or in combination, the process chamber **108** may be utilized for ALD clean up processes. For example, for an imperfect SAM treatment, some ALD nucleation may occur on defect sites of the substrate which are not covered by SAM materials. The process chamber **108** may be utilized to perform an etch process to etch exposed ALD materials

which results in removal of the ALD films grown on the defect sites while only removing a small amount of material from the intended ALD growth location.

The transfer chamber 110, which enables transfer of the substrate between the process chambers 102, 104, 106, 108, houses a transfer robot 114 therein. The transfer robot 114 may be a single blade robot or a dual blade robot as illustrated. The dual blade robot 114 has a pair of substrate transport blades 116A, 116B attached to distal ends of a pair of extendable arms. The blades 116A, 116B are used to support and carry individual substrates between the chambers 102, 104, 106, 108. The transfer chamber 110 is also maintained under vacuum or an otherwise reduced oxygen environment. Air exposure of the substrate between SAM treatment and ALD treatment is potentially detrimental to the effectiveness of the SAM material for ALD blocking and transferring the substrate between the process chamber 104 and the process chamber 106 in-situ provides for improved processing performance, such as higher deposition selectivity. In addition, it may be desirable to perform cyclic SAM and ALD processes, thus, the transfer chamber enables efficient transfer of substrates between the process chambers 104, 106 while also improving the processing performance by preventing exposure of the substrate to an ambient air environment.

FIG. 2 illustrates a schematic, cross-sectional view of the process chamber 102 according to one embodiment described herein. The process chamber 102 may be a plasma etch chamber, a plasma treatment chamber, a bake chamber, an anneal chamber, or other suitable vacuum processing chamber.

The process chamber 102 may be assembled from multiple modules. The modular design enables the process chamber 102 to meet various process requirements. As shown in FIG. 2, the process chamber 102 includes a source module 202, a process module 204, a flow module 206, and an exhaust module 208. The source module 202, the process module 204 and the flow module 206 collectively enclose a process region 212. The process module 204, the flow module 206, and the exhaust module 208 may be considered collectively as a chamber module 210 in certain embodiments.

During operation, a substrate 216 may be positioned on a substrate support assembly 218 and exposed to a process environment, such as plasma generated in the process region 212. Exemplary processes which may be performed in the process chamber 102 include etching, plasma treatment processes, and baking or annealing processes. Vacuum may be maintained in the process region 212 by suction from the exhaust module 208 through evacuation channels 214 defined by the flow module 206.

The process region 212 and the evacuation channels 214 are substantially symmetrically about a central axis 211 to provide symmetrical electrical current, gas flow, and thermal flow to establish uniform process conditions.

In one embodiment, the source module 202 is a capacitively coupled plasma source configured to generate one or more plasmas, at least one of which may be considered a remote plasma and one of which may be considered a direct plasma. The source module 202 may include a plate stack 201, which may function as an electrode (i.e. anode), isolated from and supported by the process module 204 by an isolator 222. The plate stack 201 may include various showerheads, diffusers, and screen/blocker plates arranged in a stacked orientation. The plate stack 201 may be connected to a gas source 232 through a gas inlet tube 226. The plate stack 201 and the gas inlet tube 226 may be all

fabricated from a radio frequency (RF) conductive material, such as aluminum or stainless steel. The plate stack 201 may be coupled to a RF power source 224 via the conductive gas inlet tube 226. An RF matching network 225 may also be coupled to the RF power source 224. The conductive gas inlet tube 226 may be coaxial with the central axis 211 of the process chamber 102 so that both RF power and processing gases are symmetrically provided.

Even though a capacitive plasma source is described above, the source module 202 may be any suitable gas/plasma source according to a process requirement. For example, the source module 202 may be an inductive coupled plasma source, a remote plasma source, or a microwave plasma source.

The process module 204 is coupled to the source module 202. The process module 204 may include a chamber body 240 enclosing the process region 212. The chamber body 240 may be fabricated from a conductive material resistive to processing environments, such as aluminum or stainless steel. The substrate support assembly 218 may be centrally disposed within the chamber body 240 and positioned to support the substrate 216 in the process region 212 symmetrically about the central axis 211.

A slit valve opening 242 is formed through the chamber body 240 to allow passages of the substrate 216. A slit valve 244 is disposed outside the chamber body 240 to selectively open and close the slit valve opening 242.

In one embodiment, an upper liner assembly 246 is disposed within an upper portion of the chamber body 240 shielding the chamber body 240 from the process environment. The upper liner assembly 246 includes an opening 248 corresponding to the slit valve opening 242 formed in the chamber body 240. In one embodiment, the upper liner assembly 246 includes two or more openings 248 formed symmetrically about the central axis 211 to compensate for the asymmetry of the chamber body 240 caused by the slit valve opening 242, thus creating symmetry in the process region 212 within the process chamber 102. For example, the upper liner assembly 246 is a cylindrical wall having three identical openings 248 formed 120 degrees apart from one another. The upper liner assembly 246 may be constructed from a conductive, process compatible material, such as aluminum, stainless steel, and/or yttria (e.g., yttria coated aluminum).

In one embodiment, cooling channels 250 are formed in the chamber body 240 to provide temperature control to the chamber body 240 and the upper liner assembly 246 to enhance the thermal symmetry within the process chamber 102 and symmetry of the plasma provided in the process region 212.

The flow module 206 is attached to the process module 204. The flow module 206 provides flow paths between the process region 212 defined in the process module 204 and the exhaust module 208. The flow module 206 also provides an interface between the substrate support assembly 218 and the atmospheric environment exterior to the process chamber 102.

The flow module 206 has a height 207. The height 207 is selected according to amount of vertical movement or degree of variable volume dictated by the process requirements. Accordingly, when building a process chamber for a particular process, a flow module with a suitable height may be selected to meet the process requirements. The flow module may be swapped for another flow module with a different height when configuring the process chamber for a different process.

The flow module 206 includes an outer wall 260, an inner wall 262, two or more pairs of radial walls 264 connecting between the inner wall 262 and the outer wall 260, and a bottom wall 266 attached to the inner wall 262 and the two or more pairs of radial walls 264. The outer wall 260 includes two or more through holes 270 formed between each pair of radial walls 264. A chassis 254 is sealably disposed over the inner wall 262 and the two or more pairs of radial walls 264. The through holes 270 connect an atmosphere volume 268 defined by the inner wall 262 with the exterior environment, thus accommodating utility connections, such as electrical connections, gas connections, and cooling fluid connections. The chassis 254 includes a central opening 258 for receiving the substrate support assembly 218.

The outer wall 260 of the flow module 206 is shaped to match the chamber body 240 of the process module 204. In one embodiment, the outer wall 260 includes a flange which corresponds to a flange on the chamber body 240. A plurality of bolts may be used to secure the flanges to couple the flow module 206 to the process module 204. In one embodiment, a seal 252 is disposed between the flange of the chamber body 240 and the flange of the outer wall 260 to form a vacuum seal therebetween. The seal 252 may be an o-ring or other type of seal. In one embodiment, a RF grounding gasket 272 is disposed between the flow module 206 and the process module 204 to provide a solid contact therebetween for a uniform and symmetric RF ground return path.

The inner wall 262, bottom wall 266, radial walls 264 and the chassis 254 divide the volume inside the outer wall 260 into the evacuation channels 214 and the atmosphere volume 268. The evacuation channels 214 connect with the process region 212 of the process module 204. A seal 256 is disposed in the grooves 254b, 264b to provide vacuum seal between the evacuation channels 214 and the atmosphere volume 268. The seal 256 may be an o-ring or other type of seal, such as an elastomeric seal.

The outer wall 260 and the inner wall 262 may be cylindrical walls concentrically arranged. When assembled, a central axis of the outer wall 260 and the inner wall 262 coincides with the central axis 211 of the process chamber 102. The two or more pairs of radial walls 264 are arranged between the inner wall 262 and the outer wall 260 to divide the space therebetween into the evacuation channels 214 and through holes 270. In one embodiment, the two or more pairs of radial walls 264 are arranged so that the evacuation channels 214 are symmetrical about the central axis 211. The flow module 206 includes three pairs of radial walls 264 disposed at 120 degrees apart from one another forming three evacuation channels 214 symmetrical to the central axis 211. The symmetrical arrangement of the evacuation channels 214 promotes symmetrical removal of gases from the process region 212, resulting in symmetrical flow of gases across the substrate 216. In addition, the symmetrical positioning of the evacuation channels 214 and radial walls 264 promotes symmetry in the thermal distribution and electrical conductance in the process chamber 102.

The exhaust module 208 includes a symmetric flow valve 280 and a vacuum pump 282 attached to the symmetric flow valve 280. The vacuum pump 282 may be a symmetric turbo molecular pump in certain embodiments. The symmetric flow valve 280 connects to the evacuation channels 214 to provide symmetric and uniform flow in the process chamber 102.

The substrate support assembly 218 is positioned along the central axis 211 to position the substrate 216 symmetrically about the central axis 211. The substrate support

assembly 218 is supported by the chassis 254. The substrate support assembly 218 includes a support plate 274, a base plate 276 that are disposed in the process region 212, and a hollow shaft 278 disposed through the central opening 258 of the chassis 254. A bellows 284 connects between the base plate 276 and the chassis 254 and surrounds the hollow shaft 278. The bellows 284 allows the substrate support assembly 218 to move vertically along the central axis 211 and provides vacuum seal between the atmosphere volume 268 in the flow module 206 and the process region 212 in the process module 204.

The support plate 274 may be an electrostatic chuck (ESC) having a chucking electrode 286. The chucking electrode 286 may be a monopolar 4-zone ESC, a bipolar 4-zone ESC, or a high temperature ESC. It is contemplated that the bipolar ESC may not require exposure to plasma for RF grounding during de-chucking of the substrate 216. The 4-zone ESCs may utilize four concentric heating zones (i.e. a heating element 288) to modulate center to edge heating profiles of substrates disposed on the chucking electrode 286 during processing to provide for improved temperature uniformity. The high temperature ESC may be suitable for utilization with temperatures up to about 600° C. In one embodiment, temperatures generated by the chucking electrode 286 may be suitable for sublimation of byproducts of substrate layers etched to maintain high material selectivity.

The support plate 274 may also include the heating element 288 for heating the substrate 216 during processing. The base plate 276 includes cooling channels 290 formed therein. The chucking electrode 286 may be connected to a bias power source 287 through the hollow shaft 278, the atmosphere volume 268, and one of the through holes 270. The heating element 288 is connected to a heating power source 289 through the hollow shaft 278, the atmosphere volume 268, and one of the through holes 270. The cooling channels 290 are connected to a cooling fluid source 291 through the hollow shaft 278, the atmosphere volume 268, and one of the through holes 170. In one embodiment, the support plate 274 is configured to heat the substrate 216 to a temperature greater than the vaporization temperature of a SAM material disposed on the substrate 216.

In one embodiment, an actuator assembly 292 is coupled to the hollow shaft 278 to move the substrate support assembly 218 vertically. The actuator assembly 292 may enable the substrate support assembly 218 to move within the process region 212 and change a processing position of the substrate 216. For example, the actuator assembly 292 positions the substrate support assembly 218 a distance of between about 0.5 inches and about 6 inches from the plate stack 201. The reduced distance between the plate stack 201 and the substrate support assembly 218, when compared to conventional plasma processing apparatus, provides an increased capacitively coupled plasma (CCP) generation window in a low pressure regime, such as less than 20 mTorr, for example about 1 mTorr. Accordingly, a gap between the RF electrode (i.e. the plate stack 201) and RF ground may be increased to enable low pressure CCP discharge by reducing the breakdown voltage of the CCP. In embodiments generating a CCP, the substrate support assembly 218 may also function as an electrode (i.e. cathode) or ground via the grounding gasket 272, depending upon the desired implementation. The actuator assembly 292 is disposed in the atmosphere volume 268. Lift pin actuators 294 are disposed in the atmosphere volume 268 to move lift pins 296.

A plasma screen 298 is disposed between the process region 212 and the evacuation channels 214 to confine the

plasma within the process region 212. A substrate support liner 299 is disposed around the substrate support assembly 218 to shield the substrate support assembly 218 from the process chemistry.

During operation, one or more processing gases from the gas source 232 may enter the process region 212 through the plate stack 201. Suitable process gases for utilization in a low pressure regime include H<sub>2</sub>, He, Ar, O<sub>2</sub>, NF<sub>3</sub>, NH<sub>3</sub>, N<sub>2</sub>, N<sub>2</sub>O, H<sub>2</sub>O, SiF<sub>4</sub>, SiH<sub>4</sub>, SiCl<sub>4</sub>, and various fluorocarbon precursors. An RF power may be applied between the plate stack 201 and the substrate support assembly 218 to ignite and maintain a direct plasma (ion generation) in the process region 212. The RF power may also be applied to the plate stack 201 via the gas inlet tube 226 and a remote plasma (radical generation) may be generated within the plate stack 201. Synchronized or step RF pulsing may be utilized to control the ion energy and radical density. A zero DC bias may be utilized and may provide for reduced ion bombardment which may enable reduced particle generation. Multiple RF frequencies (i.e. 400 kHz, 2 MHz, 13.56 MHz, 27 MHz, 40 MHz, 60 MHz) may be utilized to modulate plasma density and ion energy. Tailored waveform and phase angle modulation may be utilized to control ion energy distribution. It is contemplated that the RF power source 224 and RF matching network 225 may be configured to operation at powers as low as about 5 W while maintaining stable plasma discharge.

The substrate 216 disposed on the substrate support assembly 218 may be processed by both the direct plasma and the remote plasma. For example, the direct plasma generated in the process region 212 is utilized to modify materials on the surface of the substrate 216 by exposing the material to ions. Radicals generated from the remote plasma in the plate stack 201 are extracted from the remote plasma to remove the modified materials with high selectivity to a non-modified material on the substrate 216. Accordingly, a highly selective material removal process may be achieved.

The one or more processing gases utilized to form the plasmas may be continuously supplied to the process region 212 or the plate stack 201 and the vacuum pump 282 operates through the symmetric flow valve 280 and the flow module 206 to generate a symmetric and uniform plasma profile adjacent the substrate 216. By defining the process region 212 and the evacuation channels 214 in separate modules, embodiments of the present disclosure provide uniform and symmetric process environment with simplified chamber structure, thus, reducing fabrication costs and enabling damage free plasma etching processes with high degrees of material selectivity.

FIG. 3 illustrates a schematic, cross-sectional view of the process chamber 104 according to one embodiment described herein. Embodiments of FIG. 3 illustrate one implementation of a process chamber 104A. The process chamber 104A includes a chamber body 302 which defines a process volume 310. A substrate support 304 is disposed in the process volume 310 and a showerhead 312 is disposed opposite the substrate support 304. A pumping liner 350 is coupled to the chamber body 302 and is disposed radially outward of the substrate support 304. A lid plate 324 is coupled to the showerhead 312 and supported by the chamber body 302. A backing plate 314 is disposed between the showerhead 312 and the lid plate 324. An injection assembly 326 is coupled to the lid plate 324 and the injection assembly may be in fluid communication with the process volume 310.

The chamber body 302 may be manufactured from a material suitable to withstand temperatures of up to about

300° C. For example, the chamber body 302 may be formed from aluminum, alloys thereof, stainless steel, and other suitable metallic materials. A slit valve opening 360 is formed in the chamber body 302 to allow for ingress and egress of a substrate to and from the process volume 310. A slit valve door 358 is coupled to the chamber body 302 and may be moveable to seal and unseal the slit valve opening 360. In one embodiment, the slit valve door 358 is formed from the same materials as the chamber body 302. Alternatively, the slit valve door 358 may be formed from materials different from the chamber body 302.

The substrate support 304 may be moveably disposed within the process volume 310. As illustrated, the substrate support 304 is disposed in an elevated processing position. The substrate support 304 may be lowered such that a substrate supporting surface of the substrate support 304 is coplanar with or below the slit valve opening 360 to allow for positioning of a substrate on the substrate support 304. The substrate support may be formed from a material suitable for operation at elevated processing temperatures and may be a metallic material, a ceramic material, or combinations thereof. For example, the pedestal may be formed from aluminum, aluminum alloys, stainless steel, or ceramic materials, such as aluminum oxide or aluminum nitride.

The substrate support 304 has a heating member 306 disposed therein and the heating member 306 is coupled to a power source 354. The power source 354 may also provide power for raising and lowering the substrate support 304 within the process volume 310. The heating member 306 is a resistive heater or the like and may be disposed within the substrate support 304 in any desirable orientation. For example, the heating member 306 is formed in the substrate support 304 in a spiral orientation or other suitable orientation configured to heat the substrate support uniformly, such as a tortured path orientation. In one embodiment, the heating member 306 is configured to heat the substrate support 304 to a temperature between about 100° C. and about 300° C.

The pumping liner 350 is sized to surround the substrate support 304 and the process volume 310. Similar to the substrate support 304, the pumping liner 350 may be formed from metallic materials, ceramic materials, or combinations thereof. For example, the pedestal may be formed from aluminum, aluminum alloys, stainless steel, or ceramic materials, such as aluminum oxide or aluminum nitride. The pumping liner 350 has an opening 362 formed therein to allow for ingress and egress of a substrate into the process volume 310. The opening 362 is positioned substantially co-planar with the slit valve opening 360. A plurality of holes 352 are formed along an inner diameter of the pumping liner 350. The plurality of holes 352 provide for evacuation of gases and other materials from the process volume 310 to an exhaust 356. Accordingly, the process volume 310 is in fluid communication with the exhaust 356 via the holes 352 of the pumping liner 350.

The showerhead 312, which is disposed opposite the substrate support 304, is coupled directly or indirectly to and supported by the chamber body 302. The showerhead 312 may be formed from materials similar to the materials utilized for the substrate support 304 and the pumping liner 350. The showerhead 312 has a plurality of first passages 321 formed therein which extend from the process volume 310 to a first plenum 320 formed between the showerhead 312 and the backing plate 314. The first passages 321 enable fluid communication and the transfer of vapor from the first plenum 320 to the process volume 310.

A showerhead liner **308** is also disposed in the process volume **310**. The showerhead liner **308** may be formed from the same or similar materials as the showerhead **312** and the showerhead liner may be coupled to the showerhead **312**. In one embodiment, the showerhead liner **308** is a ring-like body. The showerhead liner **308** has an inner diameter which is substantially similar to an outer diameter of the substrate support **304**. The inner diameter of the showerhead liner **308** may also be sized such that the innermost surface of the showerhead liner **308** is radially outward of the first passages **321** in order to not interfere with vapor delivery to the process volume **310**. The showerhead liner **308** occupies physical space within the process volume **310** and reduces the volume of the process volume **310**, thereby reducing the amount of SAM precursor needed to form SAM molecules on a substrate. Thus, efficiency of a SAM formation process may be increased.

The showerhead **312** also has a heater **316** disposed therein. The heater **316** is a resistive heater of the like and may be disposed within the showerhead **312** radially outward of the first passages **321**. In one embodiment, the heater **316** is disposed within the showerhead **312** in a circumferential orientation which substantially surrounds the first passages **321**. The heater **316** may be coupled to a power source **318** to enable resistive heating of the showerhead **312**. In one embodiment, the showerhead **312** is configured to be heated to a temperature between about 150° C. and about 250° C.

The backing plate **314**, which is disposed between the showerhead and the lid plate **324**, and partially defines the first plenum **320**, has a second plurality of passages **323** disposed therein. A second plenum **322** is formed between the backing plate **314** and the lid plate **324**. The passages **323** enable the second plenum **322** to be in fluid communication with the first plenum **320**. A third plurality of passages **325** is formed in the lid plate **324** between the second plenum **322** and the injection assembly **326**.

The injection assembly **326** is configured to deliver vaporized materials to the process volume **310**. In operation, vaporized materials, such as SAM precursors and/or co-reactant precursors, are delivered from the injection assembly **326** through the third plurality of passages **325** to the second plenum **322**. The vaporized materials travel through the second plurality of passages **323** of the backing plate **314** to the first plenum **320** and through the first plurality of passages **321** of the showerhead **312** to the process volume **310**. After processing of a substrate, the vaporized materials and other effluent are removed from the process volume **310** through the holes **352** of the pumping liner **350** via the exhaust **356**.

The injection assembly **326** includes a housing **327** coupled to the lid plate **324** and an injector **328** coupled to the housing **327**. The injector **328** is disposed within the housing **327** and the injector **328** includes a third plenum **348**. In one embodiment, the third plenum **348** is funnel shaped. The shape of the third plenum **348** may be configured to facilitate and encourage mixing of vaporized materials prior to delivery to the process volume **310**. Although the third plenum **348** is illustrated as a funnel shape, other shapes which facilitate mixing of vaporized materials are contemplated.

A first ampoule **330** is coupled to the injection assembly **326** via a first conduit **332**. More specifically, the first ampoule **330** is in fluid communication with the third plenum **348** of the injector **328** via the first conduit **332**. The first conduit **332** extends from the first ampoule **330** to the third plenum **348**. A first heater jacket **334** surrounds the first

conduit **332** on portions of the first conduit **332** disposed beyond the injector **328**. In one embodiment, the first heater jacket **334** is resistively heated to maintain a temperature of the first conduit **332** between about 50° C. and about 250° C.

The first ampoule **330** is configured to vaporize and deliver SAM precursors to the process volume **310**. Suitable examples of SAM precursors include phosphonic acid materials, such as butylphosphonic acid, hexylphosphonic acid, octylphosphonic acid, dodecylphosphonic acid, and octadecylphosphonic acid. In another embodiment, the SAM precursors may be thiol materials, such as dodecanethiol and octadecanethiol. In another embodiment, the SAM precursors may be silylamine materials, such as tris(dimethylamino)octylsilane and tris(dimethylamino)octadecylsilane. In another embodiment, the SAM precursors may be chlorosilane materials, such as dodecyltrichlorosilane and octadecyltrichlorosilane. In another embodiment, the SAM precursors may be oxysilane materials, such as octadecylthioethoxy-silane and octadecyltrimethylsilane. In another embodiment, the SAM precursors may have a fluorinated R group, such as (1,1,2,2-perfluorodecyl)trichlorosilane, trichloro(1,1,2,2-perfluorooctyl)silane, (tridecafluoro-1,1,2,2-tetrahydrooctyl)trichlorosilane, (tridecafluoro-1,1,2,2-tetrahydro-octyl)triethoxysilane, (tridecafluoro-1,1,2,2-tetrahydrooctyl)methyldichlorosilane, (tridecafluoro-1,1,2,2-tetrahydrooctyl)dimethylchlorosilane, and (heptadecafluoro-1,1,2,2-tetrahydrodecyl)trichlorosilane, among others.

A second ampoule **336** is coupled to the injection assembly **326** via a second conduit **338**. More specifically, the second ampoule **336** is in fluid communication with the third plenum **348** of the injector **328** via the second conduit **338**. The second conduit **338** extends from the second ampoule **336** to the third plenum **348**. A second heater jacket **340** surrounds the second conduit **338** on portions of the second conduit **338** disposed beyond the injector **328**. In one embodiment, the second heater jacket **340** is resistively heated to maintain a temperature of the second conduit **338** between about 50° C. and about 250° C.

The second ampoule **336** is configured to vaporize and deliver co-reactant precursors to the process volume **310**. Suitable examples of co-reactant precursors include hydroxyl moiety materials, such as ambient air, water solution or vapor, hydrogen peroxide solution or vapor, organic alcohol solutions or vapors, such as methanol, isopropanol, ethanol, and diols, among others. Hydrogen gas and oxygen gas may also be utilized in combination to form hydroxyl moieties. It is contemplated that other non-hydroxyl moiety precursors may also be utilized in accordance with the embodiments described herein. Non-hydroxyl moiety precursors may include nitrogen gas, (di)isocyanates, hydrogen sulfide, and ammonia, among others.

In one embodiment, a cleaning gas source **342** is coupled to the injection assembly **326** via a third conduit **344**. More specifically, the cleaning gas source **342** is in fluid communication with the third plenum **348** of the injector **328** via the third conduit **344**. The third conduit **344** extends from the cleaning gas source **342** to the third plenum **348**. A third heater jacket **346** may optionally surround the third conduit **344** on portions of the third conduit **344** disposed beyond the injector **328**. In one embodiment, the third heater jacket **346** is resistively heated to maintain a temperature of the third conduit **344** between about 50° C. and about 250° C. Gases provided by the cleaning gas source **342** may include



chlorine containing materials, fluorine containing materials, and other materials suitable for cleaning components of the process chamber 104A.

In another embodiment, the cleaning gas source 342 is a remote plasma source. In this embodiment, the remote plasma source may energize cleaning gas to generate radicals and/or ions and the deliver the plasma products to the process volume 310. In one embodiment, the remote plasma source is optional.

In another embodiment, the cleaning gas source 342 is a carrier gas source. A carrier gas may be utilized to facilitate delivery of vapor phase SAM precursors and the carrier gas, depending on the process volume 310, may be delivered at a flow rate suitable to facilitate transport of the SAM precursors from the third plenum 348, through the third passages 325, through the second plenum 322 and the second passages 323, and through the first plenum 320 and the first passages 321 to the process volume 310. Suitable carrier gases include gases, such as noble gases or the like, that are generally inert under SAM adsorption conditions that facilitate delivery of the SAM molecules to surfaces of the substrate.

The heated showerhead 312 and the heated substrate support 304 may heat the process volume 310 to a temperature of between about 50° and about 250° C. The ampoules 330, 336 and conduits 332, 338 may be heated to similar temperatures. The showerhead liner 308, backing plate 314, lid plate 324, and injection assembly 326 may also be conductively heated by the showerhead 312. The temperature of the flow path along which the SAM precursors travel is maintained at an elevated temperature to prevent condensation of the vaporized SAM precursors on the various apparatus. The process volume 310 may also be maintained at a pressure of less than about 600 Torr which may also facilitate maintenance of the vapor state of the SAM precursor and co-reactant precursor.

In an operational embodiment, the SAM precursor is continually flowed through the process volume 310 from the first ampoule 330 to the exhaust 356. In this embodiment, the pressure of the process volume 310 is maintained in an isobaric state. In another embodiment, the SAM precursor fills the process volume 110 and be maintained in the process volume 110 for a period of time prior to being exhausted from the process volume 310. In another embodiment, co-reactant precursors are continually flowed to the process volume 310 or provided in a discontinuous manner, such as pulsing. In another embodiment, the SAM precursor and the co-reactant precursor are provided to the process volume 310 in an alternating manner, either continually or statically.

FIG. 4 illustrates a schematic, cross-sectional view of the process chamber 104 according to another embodiment described herein. Embodiments of FIG. 4 illustrate one implementation of a process chamber 104B. The process chamber 104B includes a chamber body 402 which defines a process volume 406. A substrate support 404 is disposed within the process volume 406 and a heater 414 is disposed in the process volume 406 opposite the substrate support 404. A lid plate 416 is coupled to the chamber body 402 and a vapor generation assembly 418 is coupled to the lid plate 416.

The chamber body 402 is formed from the same or similar materials as the chamber body 302. Similarly, the substrate support 404 may be formed from the same or similar materials as the substrate support 304. The substrate support 404 includes a heating member 408 disposed therein. The heating member 408 is coupled to a power source 410 and

be configured to heat the substrate support 404 to temperatures between about 100° C. and about 500° C.

The heater 414, which is disposed opposite the substrate support 404, may further define the process volume 406 between the heater 414 and the substrate support 404. The heater 414 is coupled to a power source 428 and is configured to heat the heater 414 to temperatures between about 100° C. and about 500° C. A temperature of the process volume 406 may be maintained during processing at a temperature of between about 50° C. and about 500° C., such as between about 100° C. and about 250° C. A gas source 426 is also coupled to the heater 414 and the gas source 426 is in fluid communication with the process volume 406. In one embodiment, the gas source 426 is configured to deliver a co-reactant precursor to the process volume 406. Alternatively, the gas source 426 is configured to deliver a purge gas, a carrier gas, or a cleaning gas to the process volume 406, depending on the desired implementation.

The vapor generation assembly 418, such as a vaporizer, a direct liquid injection vaporizer or the like, is coupled to the lid plate 416. The vapor generation assembly 418 is coupled to the lid plate 416 radially outward of the process volume 406. The position of the vapor generation assembly 418, and injection location of vapor to the process volume 406, provides for a cross-flow type exposure of a substrate to SAM precursors. The vapor generation assembly 418 includes a vaporizer 422 and an injector 420 extending from the vaporizer 422. The vaporizer 422 is coupled to a SAM precursor source 524 and receives SAM precursors in liquid form for vaporization. The vaporizer 422 is maintained at a temperature of between about 100° C. and about 500° C. to vaporize the SAM precursor and the temperature of the vaporizer 422 is determine, at least in part, by the vapor pressure of the SAM precursor.

Vaporized SAM precursor exits the vaporizer 422 and travels through the injector 420. The injector 420 extends from the vaporizer 422 through the vapor generation assembly 418, which is maintained at an elevated temperature by a heater jacket 412 to maintain the SAM precursor in a vapor state, and to the lid plate 416.

A manifold 436 is coupled to the chamber body 402 radially outward of the substrate support 404 and the heater 414. The manifold 436 is formed from the same of similar materials as the substrate support 404 and the heater 414. The manifold 436 is sized to circumscribe the process volume 406 such that an inner diameter of the manifold 436 is larger than an outer diameter of the substrate support 404 and an outer diameter of the heater 414. Vapor may flow through the manifold 436 from the injector 420 to an outlet 430 disposed opposite the injector 420. An exhaust 432 is also coupled to and in fluid communication with the process volume 406. More specifically, the exhaust 432 is in fluid communication with the process volume 406 via the outlet 430. Accordingly, process volume effluent may be evacuated from the process volume 406 through the outlet 430 to the exhaust 432.

A thermal insulator 434 is coupled to the lid plate 416 radially outward of the heater 414. The thermal insulator 434 is sized similarly to the manifold 436 and may be disposed between the manifold 436 and the lid plate 416. The thermal insulator 434 may also be coupled to or in contact with the chamber body 402. The thermal insulator 434 is formed from a thermally insulative material, such as a ceramic material or the like, configured to reduce or prevent the conduction of heat from the substrate support 404, the heater 414, and the manifold 436 from to the lid plate 416. In one

15

embodiment, the thermal insulator **434** is optional. In this embodiment, an air gap functions as a thermal break between the lid plate **416** and the substrate support **404**, the heater **414**, and the manifold **436**.

FIG. 5 illustrates a schematic, cross-sectional view of the process chamber **104** according to one embodiment described herein. Embodiments of FIG. 5 illustrate one implementation of a process chamber **104C**. The process chamber **104C** includes a chamber body **502** which defines a process volume **506**. A substrate support **504** is disposed within the process volume **506** and a lid plate **516** is coupled to the chamber body **502** opposite the substrate support **504**. A vapor generation assembly **518** is coupled to the lid plate **516**.

The chamber body **502** is formed from the same or similar materials as the chamber body **402**. Similarly, the substrate support **504** is formed from the same or similar materials as the substrate support **404**. The substrate support **504** includes a heating member **508** disposed therein. The heating member **508** is coupled to a power source **510** and is configured to heat the substrate support **504** to temperatures between about 100° C. and about 500° C.

The vapor generation assembly **518**, such as a vaporizer, a direct liquid injection vaporizer or the like, is coupled to the lid plate **516** adjacent a center of the process volume **506**. The position of the vapor generation assembly **518**, and injection location of vapor to the process volume **506**, provides for a top-down type exposure of a substrate to SAM precursors. The vapor generation assembly **518** includes a vaporizer **522** and one or more injectors **512**, **514** extending from the vaporizer **522**. The vaporizer **522** is coupled to a SAM precursor source **524** and receives SAM precursors in liquid form for vaporization. The vaporizer **522** is maintained at a temperature of between about 100° C. and about 500° C. to vaporize the SAM precursor and the temperature of the vaporizer **522** may be determined, at least in part, by the vapor pressure of the SAM precursor.

Vaporized SAM precursors exit the vaporizer **522** and travel through one or both of the injectors **512**, **514**. The injectors **512**, **514** extend from the vaporizer **522** through the vapor generation assembly **518**, which is maintained at an elevated temperature by a heater jacket **528** to maintain the SAM precursor in a vapor state, and to the lid plate **516**. In one embodiment, SAM precursor from the source **524** is introduced to the process volume via the injector **512** through an outlet **530**. A gas source **526** is also in fluid communication with the process volume **506**. The gas source **526** introduces a liquid or gas to the vapor generation assembly **518** and the generated vapor is introduced to the process volume **506** via the injector **514** and the outlet **530**. In one embodiment, the gas source **526** provides a co-reactant precursor. In another embodiment, the gas source **526** provides a purge gas, a carrier gas, or a cleaning gas, depending upon the desired implementation.

The process volume **506** is also in fluid communication with an exhaust **532**. Accordingly, process volume effluent may be evacuated from the process volume **506** via the exhaust **532**. Both process chambers **104B** and **104C** may be maintained at a pressure of less than about 600 Torr. Processes performed in the process chambers **104B**, **104C** may be isobaric or non-isobaric. Similarly, the processes performed in the process chambers **104B**, **104C** may be isothermal or non-isothermal.

FIG. 6 illustrates a schematic, cross-sectional view of the process chamber **106** according to one embodiment described herein. The process chamber **106** includes a gas delivery system **630** adapted for ALD or sequential layer

16

deposition. The process chamber **106** contains a chamber body **602** having sidewalls **604** and a bottom **606**. A slit valve **608** in provides ingress and egress of a substrate **610**, to and from the process chamber **106**.

A substrate support **612** supports the substrate **610** on a substrate receiving surface **611**. The substrate support **612** is mounted to a lift motor **614** to raise and lower the substrate support **612** and the substrate **610** disposed thereon. A lift plate **616** connected to a lift motor **618** is mounted in the process chamber **106** and raises and lowers lift pins **620** movably disposed through substrate support **612**. The lift pins **620** raise and lower the substrate **610** over the surface of substrate support **612**. The substrate support **612** may include a vacuum chuck (not shown), an electrostatic chuck (not shown), or a clamp ring (not shown) for securing the substrate **610** to the substrate support **612** during processing.

The substrate support **612** may be heated to heat the substrate **610** disposed thereon. For example, the substrate support **612** may be heated using an embedded heating element, such as a resistive heater (not shown), or may be heated using radiant heat, such as heating lamps (not shown) disposed above the substrate support **612**. A purge ring **622** may be disposed on the substrate support **612** to define a purge channel **624** which provides a purge gas to a peripheral portion of the substrate **610** to prevent deposition thereon.

A gas delivery system **630** is disposed at an upper portion of the chamber body **602** to provide a gas, such as a process gas and/or a purge gas, to the process chamber **106**. A vacuum system **678** is in communication with a pumping channel **679** to evacuate any desired gases from the process chamber **106** and to help maintain a desired pressure or a desired pressure range inside a pumping zone **666** of the process chamber **106**.

In one embodiment, the gas delivery system **630** contains a chamber lid assembly **632**. The chamber lid assembly **632** includes an expanding channel **634** extending from a central portion of chamber lid assembly **632** and a lower surface **660** extending from the expanding channel **634** to a peripheral portion of the chamber lid assembly **632**. A lower surface **660** is sized and shaped to substantially cover the substrate **610** disposed on the substrate support **612**. The expanding channel **634** has gas inlets **636a**, **636b** to provide gas flows from two similar pairs of valves **642a/652a**, **642b/652b**, which may be provided together and/or separately.

In one configuration, valve **642a** and valve **642b** are coupled to separate reactant gas sources but may also be coupled to the same purge gas source. For example, valve **642a** is coupled to reactant gas source **638** and valve **642b** is coupled to reactant gas source **639**, and both valves **642a**, **642b** are coupled to purge gas source **640**. Each valve **642a**, **642b** includes a delivery line **643a**, **643b** having a valve seat assembly **644a**, **644b** and each of valves **652a**, **652b** includes a purge line **645a**, **645b** having a valve seat assembly **646a**, **646b**. Delivery line **643a**, **643b** is in fluid communication with the reactant gas source **638**, **639** and is in fluid communication with gas inlet **636a**, **636b** of the expanding channel **634**. The valve seat assembly **644a**, **644b** of delivery line **643a**, **643b** controls the flow of the reactant gas from the reactant gas source **638**, **639** to the expanding channel **634**. Purge line **645a**, **645b** is in fluid communication with purge gas source **640** and intersects delivery line **643a**, **643b** downstream of valve seat assembly **644a**, **644b** of delivery line **643a**, **643b**. Valve seat assembly **646a**, **646b** of purge line **645a**, **645b** controls the flow of the purge gas from purge gas source **640** to the expanding channel **634**. If a carrier gas is used to deliver reactant gases from the

reactant gas source **638**, **639**, preferably the same gas is used as a carrier gas and a purge gas (i.e., an argon gas used as a carrier gas and a purge gas).

Each valve seat assembly **644a**, **644b**, **646a**, **646b** may contain a diaphragm (not shown) and a valve seat (not shown). The diaphragm may be biased open or closed and may be actuated closed or open respectively. The diaphragms may be pneumatically actuated or may be electrically actuated. Pneumatically actuated valves include pneumatically actuated valves available from Fujikin, Inc. and Veriflo Division, Parker Hannifin, Corp. Electrically actuated valves include electrically actuated valves available from Fujikin, Inc. For example, an ALD valve that may be used is the Fujikin Model No. FPR-UDDFAT-21-6.35-PI-ASN or the Fujikin Model No. FPR-NHDT-21-6.35-PA-AYT. Programmable logic controllers **648a**, **648b** may be coupled to valves **642a**, **642b** to control actuation of the diaphragms of the valve seat assemblies **644a**, **644b**, **646a**, **646b** of valves **642a**, **642b**. Pneumatically actuated valves may provide pulses of gases in time periods as low as about 0.020 seconds. Electrically actuated valves may provide pulses of gases in time periods as low as about 0.005 seconds. An electrically actuated valve typically requires the use of a driver coupled between the valve and the programmable logic controller.

Each valve **642a**, **642b** may be a zero dead volume valve to enable flushing of a reactant gas from delivery line **643a**, **643b** when valve seat assembly **644a**, **644b** is closed. For example, purge line **645a**, **645b** may be positioned adjacent valve seat assembly **644a**, **644b** of delivery line **643a**, **643b**. When valve seat assembly **644a**, **644b** is closed, purge line **645a**, **645b** may provide a purge gas to flush delivery line **643a**, **643b**. In the embodiment shown, purge line **645a**, **645b** is positioned slightly spaced from the valve seat assembly **644a**, **644b** of delivery line **643a**, **643b** so that a purge gas is not directly delivered into valve seat assembly **644a**, **644b** when open. A zero dead volume valve as used herein is defined as a valve which has negligible dead volume (i.e., not necessary zero dead volume).

Each valve pair **642a/652a**, **642b/652b** may be adapted to provide a combined gas flow and/or separate gas flows of the reactant gas and the purge gas. In reference to valve pair **642a/652a**, one example of a combined gas flow of the reactant gas and the purge gas includes a continuous flow of a purge gas from purge gas source **640** through purge line **645a** and pulses of a reactant gas from reactant gas source **638** through delivery line **643a**. The continuous flow of the purge gas may be provided by leaving the diaphragm of valve seat assembly **646a** of the purge line **645a** open. The pulses of the reactant gas from reactant gas source **638** may be provided by opening and closing the diaphragm of valve seat assembly **644a** of delivery line **643a**. In reference to valve pair **642a/652a**, one example of separate gas flows of the reactant gas and the purge gas includes pulses of a purge gas from purge gas source **640** through purge line **645a** and pulses of a reactant gas from reactant gas source **638** through delivery line **643a**. The pulses of the purge gas may be provided by opening and closing the diaphragm of valve seat assembly **646a** of purge line **645a**. The pulses of the reactant gas from reactant gas source **638** may be provided by opening and closing the diaphragm of valve seat assembly **644a** of delivery line **643a**.

Delivery lines **643a**, **643b** of valves **642a**, **642b** may be coupled to gas inlets **636a**, **636b** through gas conduits **650a**, **650b**. Gas conduits **650a**, **650b** may be integrated or may be separate from valves **642a**, **642b**. In one aspect, valves **642a**, **642b** are coupled in close proximity to expanding channel

**634** to reduce any unnecessary volume of delivery line **643a**, **643b** and gas conduits **650a**, **650b** between valves **642a**, **642b** and gas inlets **636a**, **636b**.

The expanding channel **634** contains a channel which has an inner diameter which increases from an upper portion **637** to a lower portion **635** of expanding channel **634** adjacent lower surface **660** of chamber lid assembly **632**. In one embodiment, expanding channel **234** may be shaped as a truncated cone (including shapes resembling a truncated cone). Whether a gas is provided toward the walls of expanding channel **634** or directly downward towards substrate **610**, the velocity of the gas flow decreases as the gas flow travels through expanding channel **634** due to the expansion of the gas. The reduction of the velocity of the gas flow helps reduce the likelihood the gas flow will blow off reactants adsorbed on the surface of substrate **610**.

Not wishing to be bound by theory, it is believed that the diameter of expanding channel **634**, which is gradually increasing from upper portion **637** to lower portion **635** of expanding channel **634**, allows less of an adiabatic expansion of a gas through expanding channel **634** which helps to control the temperature of the gas. For instance, a sudden adiabatic expansion of a gas delivered through gas inlet **636a**, **636b** into expanding channel **634** may result in a drop in the temperature of the gas which may cause condensation of the gas and formation of droplets. On the other hand, a gradually expanding channel **634** is believed to provide less of an adiabatic expansion of a gas. Therefore, more heat may be transferred to or from the gas, and, thus, the temperature of the gas may be more easily controlled by controlling the surrounding temperature of the gas (i.e., controlling the temperature of chamber lid assembly **632**). The gradually expanding channel **634** may contain one or more tapered inner surfaces, such as a tapered straight surface, a concave surface, a convex surface, or combinations thereof or may contain sections of one or more tapered inner surfaces (i.e., a portion tapered and a portion non-tapered).

In one embodiment, gas inlets **636a**, **636b** are located adjacent upper portion **637** of expanding channel **634**. In other embodiments, one or more gas inlets **636a**, **636b** may be located along the length of expanding channel **634** between upper portion **637** and lower portion **635**.

At least a portion of lower surface **660** of chamber lid assembly **632** may be tapered from expanding channel **634** to a peripheral portion of chamber lid assembly **632** to help provide an improved velocity profile of a gas flow from expanding channel **634** across the surface of substrate **610** (i.e., from the center of the substrate to the edge of the substrate). Lower surface **660** may contain one or more tapered surfaces, such as a straight surface, a concave surface, a convex surface, or combinations thereof. In one embodiment, lower surface **660** is tapered in the shape of a funnel.

A choke **662** may be located at a peripheral portion of chamber lid assembly **632** adjacent the periphery of substrate **610**. Choke **662**, when chamber lid assembly **632** is assembled to form a processing zone around substrate **610**, contains any member restricting the flow of gas therethrough at an area adjacent the periphery of substrate **610**. In one embodiment, the spacing between choke **662** and substrate support **612** is between about 0.04 inches and about 2.0 inches, and preferably between 0.04 inches and about 0.2 inches. The spacing may vary depending on the gases being delivered and the process conditions during deposition. Choke **662** helps provide a more uniform pressure distribution within the volume or reaction zone **664** defined between

chamber lid assembly **632** and substrate **610** by isolating reaction zone **664** from the non-uniform pressure distribution of pumping zone **666**.

In one aspect, since reaction zone **664** is isolated from pumping zone **666**, a reactant gas or purge gas needs only adequately fill reaction zone **664** to ensure sufficient exposure of substrate **610** to the reactant gas or purge gas. Process chamber **106** sequentially introduces reactants to the surface of substrate **610** to provide absorption of alternating thin layers of the reactants onto the surface of substrate **610**. As a consequence, atomic layer deposition does not require a flow of a reactant which reaches the surface of substrate **610** simultaneously. Instead, a flow of a reactant needs to be provided in an amount which is sufficient to adsorb a thin layer of the reactant on the surface of substrate **610**.

Chamber lid assembly **632** may include cooling elements and/or heating elements depending on the particular gas being delivered therethrough. Controlling the temperature of chamber lid assembly **632** may be used to prevent gas decomposition, deposition, or condensation on chamber lid assembly **632**. For example, water channels (not shown) may be formed in chamber lid assembly **632** to cool chamber lid assembly **632**. In another example, heating elements (not shown) may be embedded or may surround components of chamber lid assembly **632** to heat chamber lid assembly **632**. In one embodiment, components of chamber lid assembly **632** may be individually heated or cooled. For example, chamber lid assembly **632** may contain lid plate **670** and lid cap **672** in which lid plate **670** and lid cap **672** form expanding channel **634**. Lid cap **672** may be maintained at one temperature range and lid plate **670** may be maintained at another temperature range. For example, lid cap **672** may be heated by being wrapped in heater tape or by using another heating device to prevent condensation of reactant gases and lid plate **670** may be maintained at ambient temperature. In another example, lid cap **672** may be heated and lid plate **670** may be cooled with water channels formed therethrough to prevent thermal decomposition of reactant gases on lid plate **670**.

Chamber lid assembly **632** contains components that may be made of stainless steel, aluminum, nickel-plated aluminum, nickel, or other suitable materials compatible with the processing to be performed. In one embodiment, lid cap **672** contains aluminum or stainless steel and lid plate **670** contains aluminum.

Control unit **280**, such as a programmed personal computer, work station computer, or the like, may be coupled to the process chamber **106** to control processing conditions. For example, control unit **680** may be configured to control flow of various process gases and purge gases from gas sources **638**, **639**, and **640** through valves **642a**, **642b** during different stages of a substrate process sequence. Illustratively, the control unit **680** contains central processing unit (CPU) **682**, support circuitry **684**, and memory **686** containing associated control software **683**.

The control unit **680** may be one of any form of general purpose computer processor that can be used in an industrial setting for controlling various chambers and sub-processors. The CPU **682** may use any suitable memory **686**, such as random access memory, read only memory, floppy disk drive, hard disk, or any other form of digital storage, local or remote. Various support circuits may be coupled to the CPU **682** for supporting process chamber **106**. The control unit **680** may be coupled to another controller that is located adjacent individual chamber components, such as programmable logic controllers **648a**, **648b** of valves **642a**, **642b**. Bi-directional communications between the control unit **680**

and various other components of process chamber **106** are handled through numerous signal cables collectively referred to as signal buses **688**. In addition to control of process gases and purge gases from gas sources **638**, **639**, **640** and from programmable logic controllers **648a**, **648b** of valves **642a**, **642b**, control unit **680** may be configured to be responsible for automated control of other activities used in wafer processing—such as wafer transport, temperature control, chamber evacuation, among other activities.

In operation, the substrate **610** is delivered to the process chamber **106** through the slit valve **608** by a robot (not shown), such as the transfer robot **114** (FIG. 1). Substrate **610** is positioned on substrate support **612** through cooperation of the lift pins **620** and the robot. Substrate support **612** raises substrate **610** into close opposition to lower surface **660** of chamber lid assembly **632**. A first gas flow may be injected into expanding channel **634** of process chamber **106** by valve **642a** together or separately (i.e., pulses) with a second gas flow injected into process chamber **106** by valve **642b**. The first gas flow may contain a continuous flow of a purge gas from purge gas source **640** and pulses of a reactant gas from reactant gas source **638** or may contain pulses of a reactant gas from reactant gas source **638** and pulses of a purge gas from purge gas source **640**. The second gas flow may contain a continuous flow of a purge gas from purge gas source **640** and pulses of a reactant gas from reactant gas source **639** or may contain pulses of a reactant gas from reactant gas source **639** and pulses of a purge gas from purge gas source **640**. The gas flow travels through expanding channel **634** as a pattern of vortex flow which provides a sweeping action across the inner surface of expanding channel **634**. The pattern of vortex flow dissipates to a downwardly flow toward the surface of substrate **610**. The velocity of the gas flow reduces as it travels through expanding channel **634**. The gas flow then travels across the surface of substrate **610** and across lower surface **660** of chamber lid assembly **632**. Lower surface **660** of chamber lid assembly **632**, which is downwardly sloping, helps reduce the variation of the velocity of the gas flow across the surface of substrate **610**. The gas flow then travels by choke **662** and into pumping zone **666** of process chamber **106**. Excess gas, by-products, etc. flow into the pumping channel **679** and are then exhausted from process chamber **106** by vacuum system **678**. In one aspect, the gas flow proceeds through expanding channel **634** and between the surface of substrate **610** and lower surface **660** of chamber lid assembly **632** in a laminar manner which aids in uniform exposure of a reactant gas to the surface of substrate **610** and efficient purging of inner surfaces of chamber lid assembly **632**.

FIG. 7 illustrates a schematic, cross-sectional view of the process chamber **108** according to one embodiment described herein. In one embodiment, the process chamber **108** is similar to the process chamber **102**. In this embodiment, the process chamber **108** may be utilized for thermal post-ALD treatment, plasma post-ALD treatment, or a combination thereof. For embodiments utilizing a plasma post-ALD treatment, a plasma etch process may be performed to remove ALD material from undesired locations on the substrate. It is contemplated that the plasma etch process may be utilized if an imperfect SAM treatment resulted in undesired areas of the substrate remaining exposed during ALD treatment.

In another embodiment, the process chamber **108** is different from the process chamber **102**. A different chamber is illustrated in FIG. 7 having a chamber body **702** defining a process volume **704**. A substrate support **706** is disposed within the process volume **704** and a heater **708**, such as a

21

resistive heater, is disposed within the substrate support **706**. The heater **708** is coupled to a power source **710** and the heater is configured to heat a substrate disposed on the substrate support **706** to a temperature greater than a vaporization temperature of SAM materials disposed on the substrate. Thus, the process chamber **108** volatilizes SAM materials from the substrate and the volatilized materials are exhausted from the process volume **704** via an exhaust.

FIG. **8** illustrates operations of a method **800** for processing a substrate according to one embodiment described herein. At operation **810**, a substrate is delivered to a pre-clean chamber, such as process chamber **102**. A pre-clean process is performed on the substrate and the substrate is transferred from the pre-clean chamber to a SAM processing chamber, such as process chamber **104**, via a transfer chamber (i.e. transfer chamber **110**) at operation **820**.

A SAM material deposition process is performed on the substrate and at operation **830**, the substrate is transferred from the SAM processing chamber to an ALD chamber, such as process chamber **106**, via the transfer chamber. An ALD process is performed on the substrate and the substrate is transferred from the ALD chamber to a post-processing chamber, such as process chamber **108**, via the transfer chamber at operation **840**. Post-processing is performed on the substrate and the substrate may be transferred from the post-processing chamber for subsequent processing.

In summation, the integrated cluster tool apparatus provides for improved in-situ processing and increased throughput when utilizing SAM treatment processes in combination with subsequent deposition processes, such as ALD treatments. Additional process capability, such as pre and post processing of substrates, provides various advantages, such as improved material deposition selectivity, in accordance with embodiments described herein.

While the foregoing is directed to embodiments of the present disclosure, other and further embodiments of the disclosure may be devised without departing from the basic scope thereof, and the scope thereof is determined by the claims that follow.

What is claimed is:

1. A substrate processing apparatus, comprising:
  - a cluster tool having a centrally disposed transfer chamber;
  - a first process chamber coupled to the transfer chamber, the first process chamber configured to perform a surface modification process;
  - a second process chamber coupled to the transfer chamber, the second process chamber configured to perform a self-assembled monolayer treatment process, the second process chamber comprising:
    - a chamber body defining a process volume;
    - a substrate support disposed in the process volume;
    - a lid plate coupled to the chamber body;
    - a heater disposed in the process volume;
    - an annular manifold coupled to the chamber body radially outward of the substrate support and the heater;
    - an injection assembly in fluid communication with the annular manifold, the injection assembly configured to provide a cross-flow type exposure of self-assembled monolayer precursors within the process volume along a top surface of the substrate support; and
    - a vapor generation assembly in fluid communication with the injection assembly;

22

a third process chamber coupled to the transfer chamber, the third process chamber configured to perform an atomic layer deposition process; and

a fourth process chamber coupled to the transfer chamber, the fourth process chamber configured to perform an anneal process.

2. The apparatus of claim 1, wherein the transfer chamber has a transfer robot disposed therein.

3. The apparatus of claim 2, wherein the transfer robot is a single blade robot.

4. The apparatus of claim 2, wherein the transfer robot is a dual blade robot.

5. The apparatus of claim 1, wherein the first process chamber is configured to generate a plasma therein to perform the surface modification process.

6. The apparatus of claim 1, wherein the first process chamber has a heated pedestal disposed therein.

7. The apparatus of claim 6, wherein the heated pedestal is configured to heat a substrate to a temperature greater than a vaporization temperature of a self-assembled monolayer material.

8. The apparatus of claim 1, wherein the first process chamber and the fourth process chamber are identical.

9. The apparatus of claim 1, wherein the second process chamber is operably maintained at a temperature greater than a vaporization temperature of a self-assembled monolayer material.

10. The apparatus of claim 1, wherein the third process chamber is configured to perform a cyclic atomic layer deposition process.

11. The apparatus of claim 1, wherein the fourth process chamber has a heated pedestal disposed therein.

12. The apparatus of claim 11, wherein the heated pedestal is configured to heat a substrate to a temperature greater than a vaporization temperature of a self-assembled monolayer material.

13. The apparatus of claim 1, further comprising: one or more load lock chambers coupled to the transfer chamber.

14. The apparatus of claim 13, wherein the load lock chambers are configured to perform an anneal process.

15. A substrate processing apparatus, comprising: a cluster tool having a centrally disposed transfer chamber;

a first process chamber coupled to the transfer chamber, the first process chamber configured to perform a surface modification process;

a second process chamber coupled to the transfer chamber adjacent the first process chamber, the second process chamber configured to perform a self-assembled monolayer treatment process, the second process chamber comprising:

- a chamber body defining a process volume;
- a substrate support disposed in the process volume;
- a lid plate coupled to the chamber body;
- a heater disposed in the process volume;

an annular manifold coupled to the chamber body radially outward of the substrate support and the heater;

an injection assembly in fluid communication with the annular manifold, the injection assembly configured to provide a cross-flow type exposure of self-assembled monolayer precursors within the process volume along a top surface of the substrate support; and

a vapor generation assembly in fluid communication with the injection assembly;

a third process chamber coupled to the transfer chamber adjacent the second process chamber, the third process chamber configured to perform an atomic layer deposition process; and

a fourth process chamber coupled to the transfer chamber adjacent the third process chamber, the fourth process chamber configured to perform an anneal process.

16. The apparatus of claim 15, where in the first process chamber and the fourth process chamber each have a heated pedestal disposed therein.

17. The apparatus of claim 16, wherein the first process chamber and the fourth process chamber are identical.

\* \* \* \* \*