Chemical vapour deposition, CVD

Mainly thin film application One of the most important methods for preparing thin films and coatings

- •Protection (corrosion, wear...)
- •Optical properties
- •Electronic properties
- Magnetic properties
- Decoration
- •New properties



"Chemical vapour deposition (CVD) is a process where one or more volatile precursors are transported via the vapour phase to the reaction chamber, where they decompose on a heated substrate"

Many materials may be deposited using CVD and related techniques. Metals, oxides, sulfides, nitrides, phosphides, arsenides, carbides, borides, silicides...

Example: Preparation of TiB₂, melting point 3325°C. May be deposited by CVD at 1000°C:

 $TiCl_4 + 2BCl_3 + 5H_2 \rightarrow TiB_2 + 10 HCl$

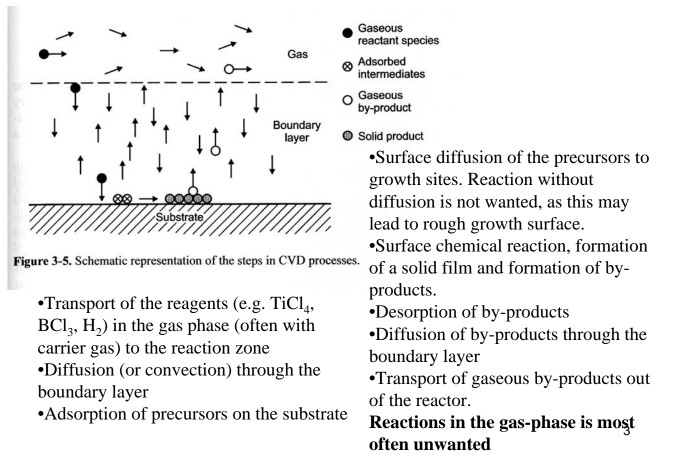
CVD was first used for hard coatings (cutting tools etc.)

Microelectronics, 3D-structures

Glass (SnO₂, TiN, SiO₂, TiO₂)

Solar cells, catalysis, membranes, waveguides, mirrors, "synthetic gold" (TiN_x)

Steps in the CVD process



The ideal precursor

- Liquid rather than solid or gaseous
- Good volatility
- Good thermal stability in the delivery system, during evaporation and transport
- Decompose cleanly and controllably on the substrate without incorporation
- Give stable by-products which are readily removed from the reaction zone
- Readily available in consistent quality and quantity at low cost
- Non-toxic and non-pyrophoric

Impossible to meet all criteria.

Industrially important precursors:

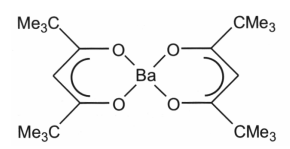
Hydrides: SiH_4 , AsH_3 ... Metal alkyls: Al^iBu_3 , $GaEt_3$ Metall halides: WF_6 , $TiCl_4$

Precursors

Volatility and gas phase reactivity Metal-organic and organometallic precursors are often used. Volatility is enhanced by minimizing intermolecular interactions (e.g. hydrogen bonds, dipole-dipole interaction and van der Waals interactions) Small molecules generally have higher vapour pressure. Oligomerization and aggregation should be minimized.

May use e.g. F-substitution

 β -diketonate ligands are often used in metal-containing precursors. Bidentate with good chelating properties. Metal diketonates are often monomeric and volatile.



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Figure 3-22. Barium dipivaloylmethanate, Ba(dpm)<sub>2</sub>.
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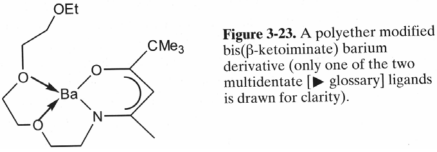
Oligomerization

Metal precursors are often under-coordinated. Oligomerization may occur in order to satisfy the metal coordination.

This may be suppressed by:

•Sterically demanding ligands limits the accessibility of the metal center. These compound are often very reactive towards small molecules (O_2, H_2O) and the bulky ligands may reduce volatility

•Using multidentate, chelating, ligands. May satisfy the metal coordination requirements and prevent oligomerization. However, multidentate ligands may form bridges between molecules and dissociation and oligomerization may occur prior to reaction.



Problems also associated with carbon incorporation if the decomposition is not "clean" enough.

С

Multi element CVD

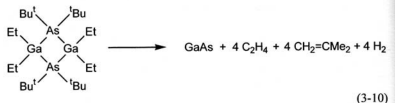
Two approaches:

Use two (or more) individual precursors. Must decompose individually on the substrate and react to give the desired product.

 $GaMe_3 + AsH_3 \rightarrow GaAs + 3CH_4$

Problems: Must have similar adsorption and decomposition properties and compatible volatilities. Stoichiometry difficulties.

Single source precursors contain all the elements in the desired stoichiometry. Bonds between the film-forming elements must be stronger than to the ligands in order to obtain clean decomposition. Single source precursor for GaAs:



Problems with single-source precursors are often related to increased molecular weight (low volatility) and to decomposition behaviour.

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Equipment

Substrate temperature 200-800°C, pressure 0.1 mbar – 1 bar Vapour pressure of precursor should be > 0.01 bar at 25° C

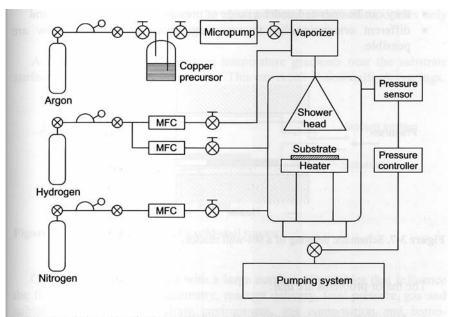


Figure 3-6. Schematic of a thermal CVD reactor for copper CVD from a Cu(II) precursor (see Eq. 3-19) (MFC = mass flow controller).

Hot wall reactor

Advantages:

- •Simple to operate
- •May accommodate several
- substrates
- •Uniform substrate temperature
- •May be operated at a range of
- temperatures and pressures
- •Different orientations of substrate

Disadvantages:

Deposition occur not only at the substrate, but also on e.g. reactor walls
The consumption of precursor is large and difficult to control. May result in feed-rate-limited deposition
Gas-phase reactions may occur

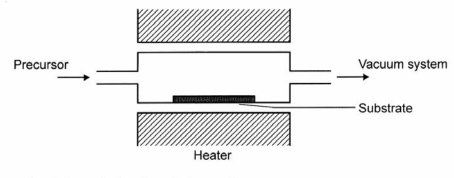


Figure 3-7. Schematic drawing of a hot-wall reactor.

Cold wall reactor

Substrate is at a higher temperature than the reactor walls. Often used in industry

Advantages:

- •Pressure and temperature can be controlled
- •Plasmas can be used
- •No deposition on reactor walls
- •Gas-phase reactions are suppressed
- •Higher deposition rates may be reached (higher precursor efficiency

Disadvantages:

•Steep temperature gradients near the substrate surface may lead to convection •Less flexibility on substrate orientation and usually only one substrate at a time.

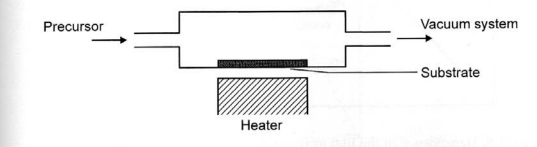


Figure 3-8. Schematic drawing of a cold-wall reactor.

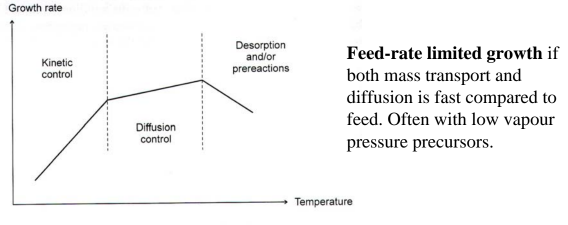
Growth rates

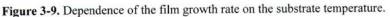
Many parameters influences the growth and coating properties For industrial use, a growth rate of 0.1μ m/min is necessary (for electronics) or higher (e.g. for glass coating)

Three regimes:

•Surface reaction (kinetically) limited.

- •Diffusion (or mass transport) limited.
- •Increased desorption rate of precursors.



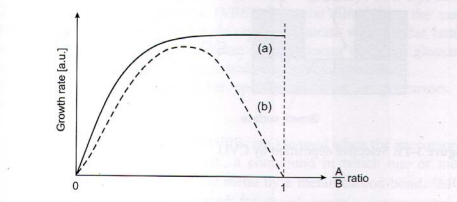


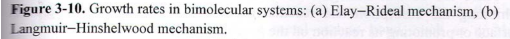
Bi-molecular systems

Reaction between A and B

•Elay-Rideal mechanism: Only A is adsorbed, and reacts directly with B in gas-phase. The growth rate shows a saturation with high coverage of the surface with A-molecules.

•Langmuir-Hinshelwood mechanism: Both A- and B-molecules are adsorbed and react on the surface. There is a maximum in growth rate when the surface is equally covered by the two types of molecules.





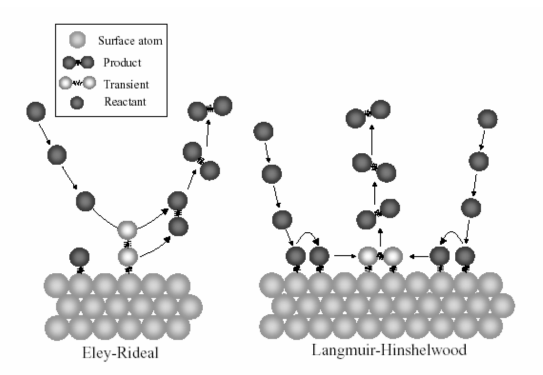


Figure 1.1: Graphic illustration of the Eley–Rideal (ER see left panel) and the Langmuir– Hinselwood (LH see right panel) mechanism. While in an ER–reaction an adsorbed species reacts by a direct collision with a gas–phase species, reactions proceeding via a LH mechanism reactants are equilibrated to the surface prior to reaction.

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Selective deposition

Especially important in microelectronics; patterning of the surface and 3D-growth

- •Reaction rate of precursor is limited on a non-growth surface. E.g. deposition of Cu from $(hfac)Cu(PMe_3)$ occur on Cu, Pt... but not on SiO₂.
- •Growth surface acts as co-reactant, and is selectively consumed. E.g. Si reacts with WF_6 or MoF₆, while reaction at SiO₂ or Si₃N₄ is slower.
- •A chemical reaction of a gaseous co-reactant occur on the growth surface. E.g. H_2 dissociation on a metal surface, but not on SiO₂ or metal oxide surfaces.
- •The rate is increased on part of the surface by irradiation.
- •Selective passivation by removal of reactive surface species. (E.g. -OH exch. -OSiMe₃) •Hindering nucleation (e.g. due to high surface energy)

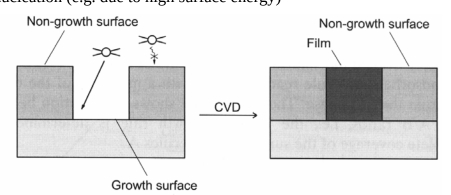


Figure 3-11. Selective deposition by CVD.

CVD-related techniques

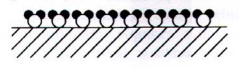
CVD	chemical vapor deposition		
PVD	physical vapor deposition		
VPE	vapor phase epitaxy		
OMCVD	organometallic CVD		
OMVPE	organometallic VPE		
MOCVD	metal-organic CVD		
MOVPE	metal-organic VPE		
LPCVD	low-pressure CVD		
PECVD	plasma-enhanced CVD		
PACVD	plasma-assisted CVD		
LCVD	laser-induced CVD		
RPCVD	remote-plasma CVD		
CVI	chemical vapor infiltration		
ALD	atomic-layer deposition		
CBE	chemical-beam epitaxy		
MOMBE	metal-organic molecular-beam epitaxy		

Table 3-2. Survey	of frequently used	d acronyms in CV	/D and related methods.
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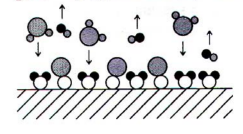
15

ALE, ALD

ZnCl₂ (ad)



 $\text{ZnCl}_2(\text{ad}) + \text{H}_2\text{S}(\text{g}) \longrightarrow \text{ZnS}(\text{ad}) + 2 \text{ HCl}(\text{g})$



ZnS (ad)

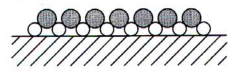
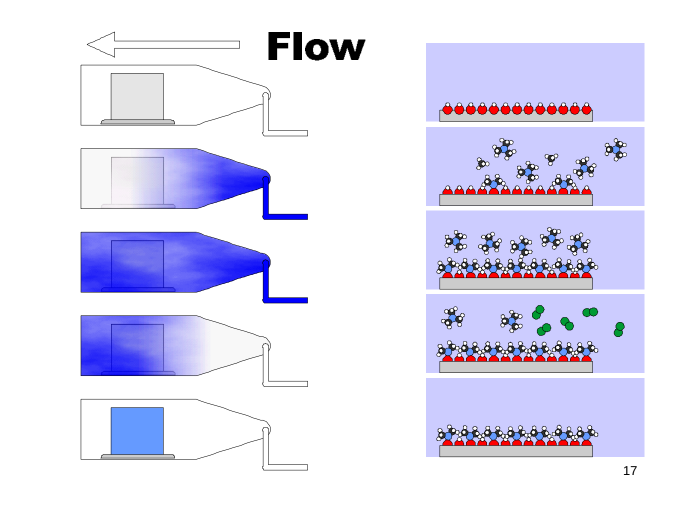
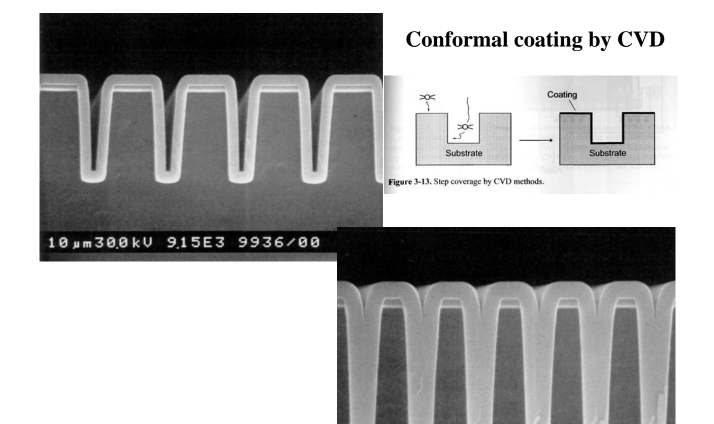


Figure 3-12. ALD reaction cycle leading to the formation of a ZnS thin film.





1µm300kV 148E4 9942/00 AL20

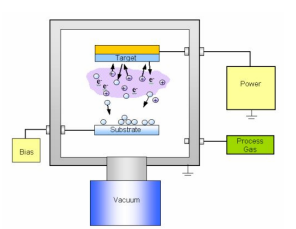
Non-CVD processes for gas phase deposition

PVD, Physical vapour deposition

Evaporation/sputtering of a target material onto a substrate.

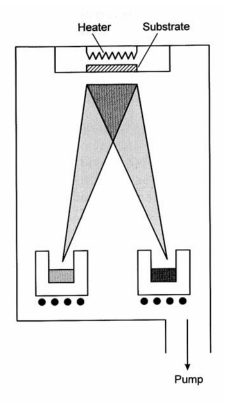
Evaporation of a solid
Transport of the gaseous species to the substrate
Condensing gaseous species on the substrate, followed by nucleation and growth

Thermal evaporation Electron or laser beam heats the surface Heating a crucible by induction or resistive heating



Sputtering: Atoms are removed from the target by ion-bombardment (glow discharge or plasma). Ballistic transport pf atoms from the target to the substrate.

Advantages: E.g. low substrate temperature (allows deposition e.g. on polymers) Disadvantages: E.g. shadowing effect.



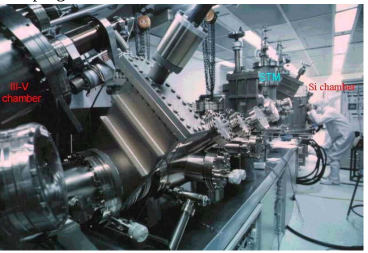
MBE, Molecular beam epitaxy

Evaporation of elemental sources. Independently at a controlled rate.

Molecular beams intercept at the substrate surface.

UHV conditions (10⁻¹⁰ bar), low growth rates. Line-of-sight.

High purity, complex layer structures, good control of doping.



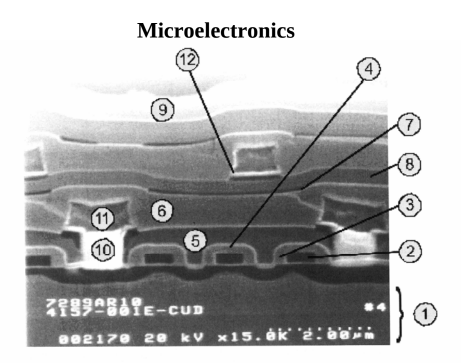


Figure 3-15. A cross-section through a multi-level structure of an electronic device. 1: silicon substrate; 2: gate, 3: spacer (CVD oxide); 4: covering oxide; 5: borate-phosphate-silicate glass (PACVD). 6: silica (PACVD from $Si(OEt)_4$); 7: glass layer (spin-on). 8: phosphate-doped silica (plasma CVD from $Si(OEt)_4$); 9: oxynitride/nitride passivation layer (PACVD); 10: tungsten plug (thermal CVD); 11: AlCu (PVD); 12: TiN barrier (reactive PVD).

21

Metal CVD

Protective films, reflective or conducting coating, electrodes, microelectronics

Commercially PVD methods are often used for metallic films. However Al, Cu and W are often deposited by CVD methods.

Aluminium:

•Metallized polymer films in food packaging (gas diffusion barrier), Reflective layers (mirrors, CDs...)

•Interconnects in microelectronics.

Common precursor for CVD: tributylaluminium: AlⁱBu₃,

 β -hydride elimination >50°C, suppressed in the gas phase by adding isobutene

Deposition at 2-300°C (hot wall reactor) (β -hydrogen elimination) At temperatures > 330C: b-methyl elimination \rightarrow carbon incorporation

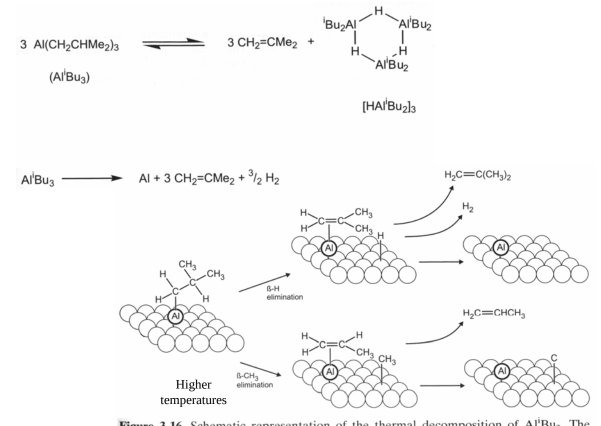


Figure 3-16. Schematic representation of the thermal decomposition of AlⁱBu₃. The upper pathway (at low temperatures) results in the clean deposition. At higher temperatures (lower pathway) carbon impurities may be incorporated into the film. 23



Growth on Si and Al, but not on silica, SiO_2 On silica, two butyl groups are easily lost, but elimination of the third is inhibited

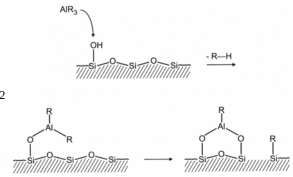


Figure 3-17. Inhibition of Al deposition from AlR₃ on a silica surface.

Tertiary amine adducts of AlH_3 may be used as precursors.

Easy cleaving of Al-N bond, and desorption of trimethylamine

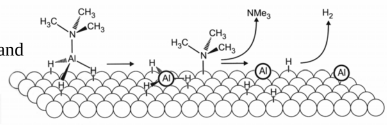


Figure 3-18. Thermal decomposition mechanism of Me₃N-AlH₃ on Al surfaces.

W-deposition

Important areas of use:

•Wear protection (cutting and grinding) and corrosion protection

•Metallization in integrated circuits. High resistance to electromigration, low thermal expansion and does not react with most semiconductor materials. Low adhesion to $SiO_2...$

Tungsten hexafluoride (WF₆) Thermal decomposition >750C, reducing agents are used in CVD. (H_2 , Si, SiH₄)

 $2WF_6 + 3Si \rightarrow (<400 \text{ °C}) \rightarrow 2W + 3SiF_4$ $WF_6 + 3Si \rightarrow (>400 \text{ °C}) \rightarrow W + 3SiF_2$

Reacts fast, but reaction stops because Si must diffuse through W Si is added in the gas stream:

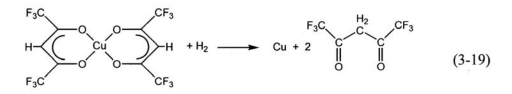
 $2WF_6 + 3SiH_4 \rightarrow (250 \text{ °C}) \rightarrow 2W + 6H_2 + 3SiF_4$ $4WF_6 + 3SiH_4 \rightarrow (>600 \text{ °C}) \rightarrow 4W + 12HF + 3SiF_4$

Cu CVD

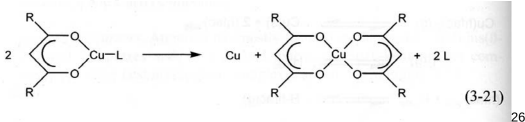
Cu would be a good substitute for Al in integrated circuits. Good conductance, less electroimigration, but higher diffusion rates in Si and SiO_2 .

Halides are not volatile enough. Metalorganic Cu compounds are used.

Cu(II) precursors are reduced:



Cu(I) precursors disproportionate:



Diamond CVD

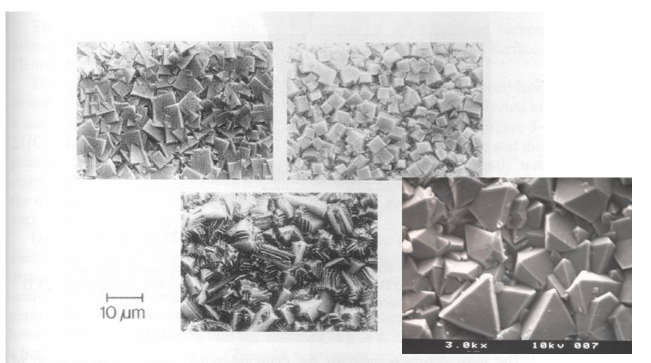
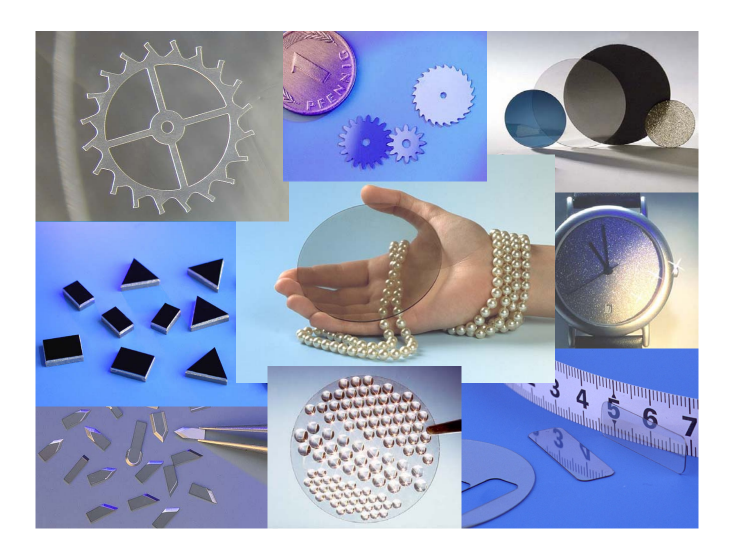


Figure 3-20. Diamond films (on a SIALON surface) deposited by microwave activation of 1.7 % methane in hydrogen. The different morphologies originate from different gas pressures and different microwave powers.



Diamond CVD

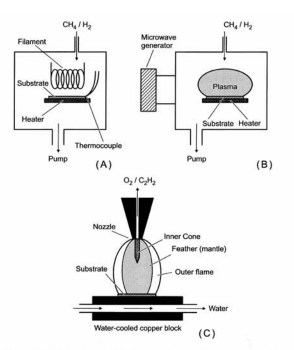
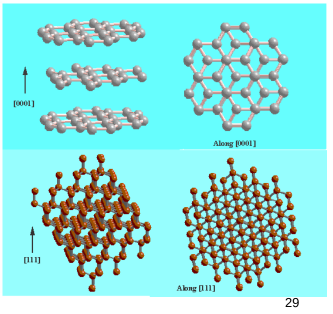


Figure 3-21. Schematic diagrams of the three most commonly used apparatus for diamond deposition. (A) Hot filament reactor. (B) Microwave plasma-enhanced CVD reactor. (C) Oxyacetylene torch.

Diamond is metastable at low pressure

Growth is possible due to H• present

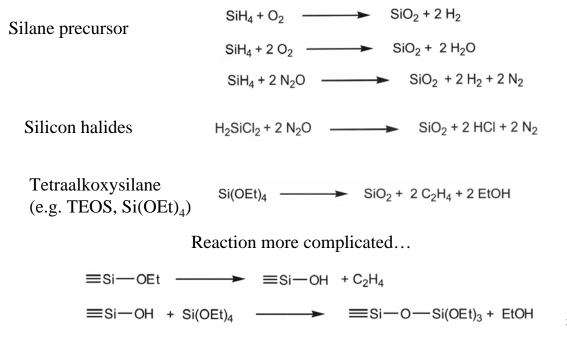
- Suppression of nucleation of graphite through selective etching
- Diamond surface is stabilized by H



CVD of metal oxides

Oxides may be formed by using oxidizing gasses as O_2 , N_2O , H_2O , H_2O_2 . Si O_2 in microelectronics:

Gate insulating layers, surface passivation, planarization, packaging



YBCO, yttrium barium copper oxide

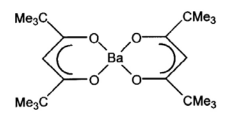


Figure 3-22. Barium dipivaloyImethanate, Ba(dpm)2.

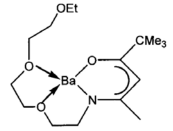


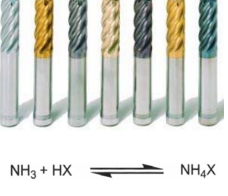
Figure 3-23. A polyether modified bis(B-ketoiminate) barium derivative (only one of the two multidentate [) glossary] ligands is drawn for clarity).

Metal nitrides Nitrides are often hard compounds, and are used for protection and hard coatings Nitrogen sources are N₂, NH₃, N₂H₄ (hydrazine)

3 H₂SiCl₂ + 4 NH₃ ----- Si₃N₄ + 6 HCl + 6 H₂

2 TiCl₄ + 4 H₂ + N₂ ----- 2 TiN + 8 HCl

6 TiCl₄ + 8 NH₃ - 6 TiN + 24 HCl + N₂



Suppressed at high temperatures

Transamination reactions between metal dialkylamides and ammonia

$$\equiv M - NR_2 + NH_3 \longrightarrow \equiv M - NH_2 + HNR_2$$

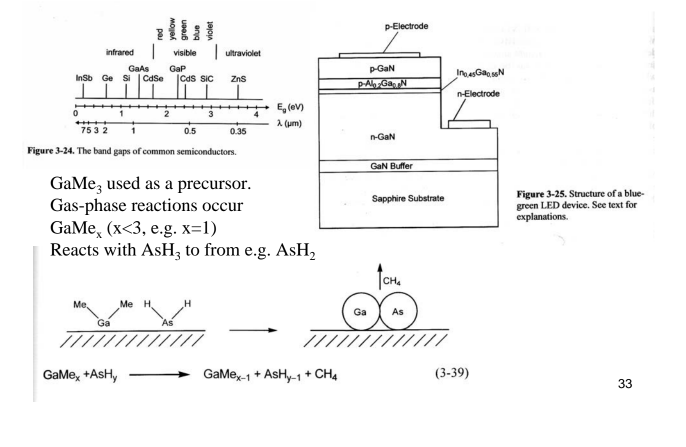
$$3 \equiv M - NH_2 \longrightarrow \qquad \stackrel{M - N - M}{I} + 2 NH_3$$

$$3 = Si(NMe_2)_4 + 4 NH_3 \longrightarrow \qquad Si_3N_4 + 12 HNMe_2$$

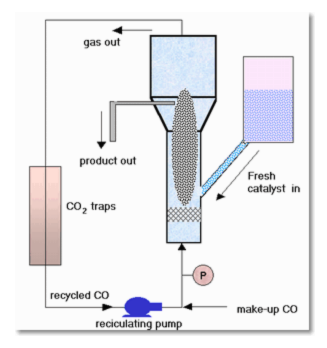
$$6 = Ti(NR_2)_4 + 8 NH_3 \longrightarrow \qquad 6 = TiN + 24 HNR_2 + N_2$$

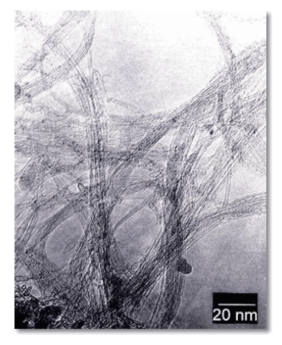
Semiconductors

III-V (GaN, GaAs, InP...) and II-VI (CdS, ZnSe...) semiconductors

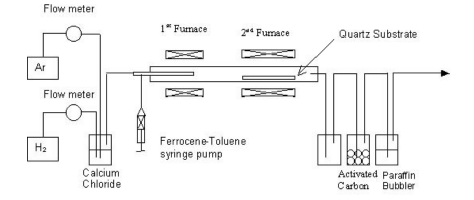


Carbon nanotubes by CVD





35



CVD production of aligned single walled carbon nanotubes (SWCNT)

