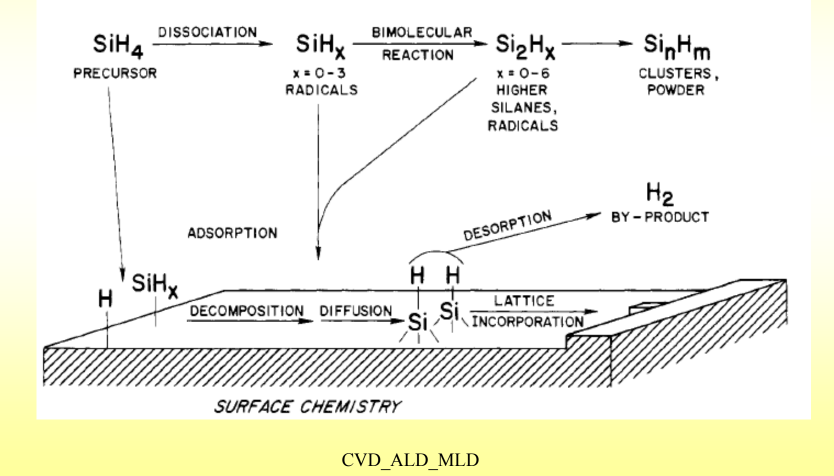
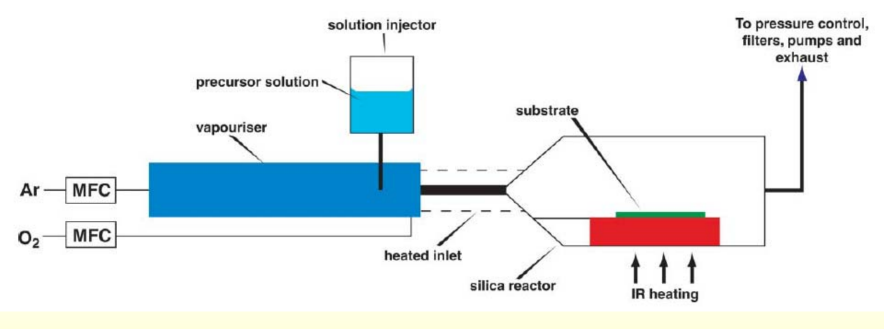


## Silicon CVD

GAS PHASE CHEMISTRY

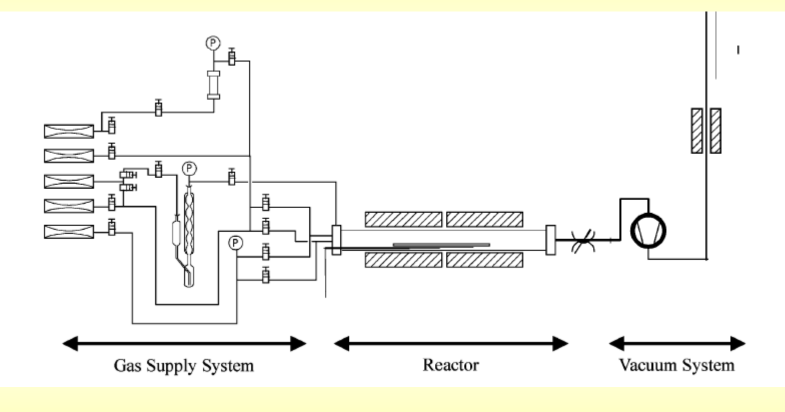


## **CVD Reactor**



Cold-wall reactor

## **CVD Reactor**



Hot-wall reactor

## **CVD Kinetics**

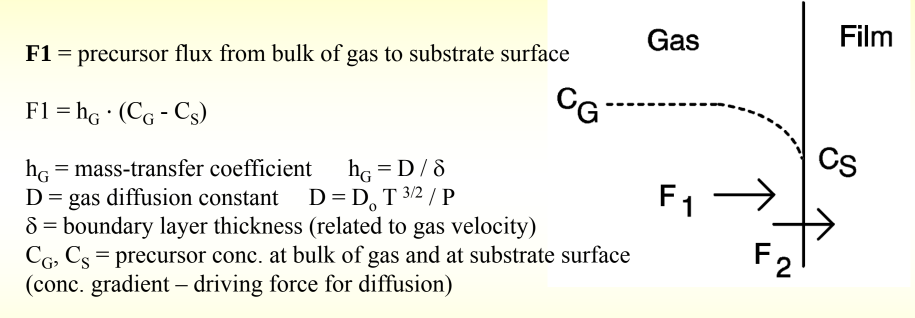
Deposition depends on the sequence of events:

(1) Diffusion of precursor to surface
 (2) Adsorption of precursor at surface
 (3) Chemical reaction at surface
 (4) Desorption of products from surface
 (5) Diffusion of products from surface

• The *slowest* event will be the rate-determining step

# **CVD Kinetics**

#### **Growth Rate Model**

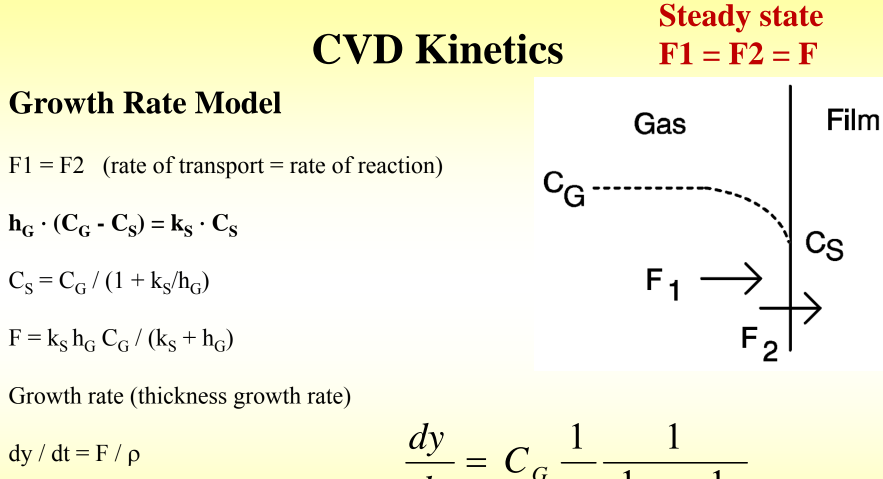


**Steady state** 

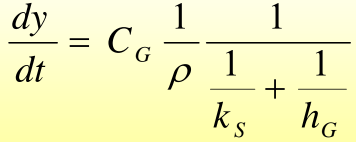
F1 = F2 = F

F2 = flux consumed in film-growth reaction (rate of chemical reaction)

 $F2 = k_{S} \cdot C_{S}$ k<sub>S</sub> = surface-reaction rate constant: k<sub>S</sub> = A exp (-E<sub>a</sub>/kT) CVD ALD MLD



y = film thickness $\rho = atomic density of film$ 



## **Growth Rate**

$$\frac{dy}{dt} = C_G \frac{1}{\rho} \frac{1}{\frac{1}{k_S} + \frac{1}{h_G}}$$

Growth rate is determined by:

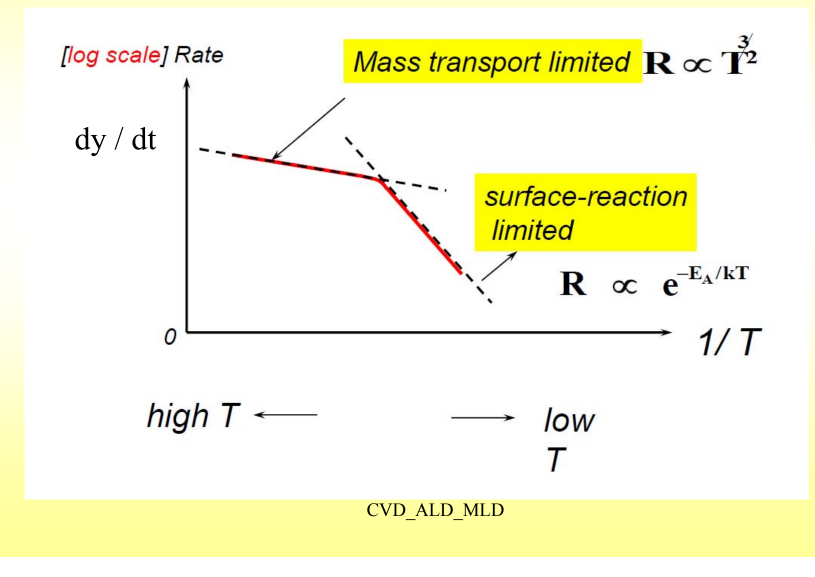
a) Concentration of a precursor in bulk of gas mixture

b) By the smaller of 
$$h_{\rm G}$$
 and  $k_{\rm S}$ 

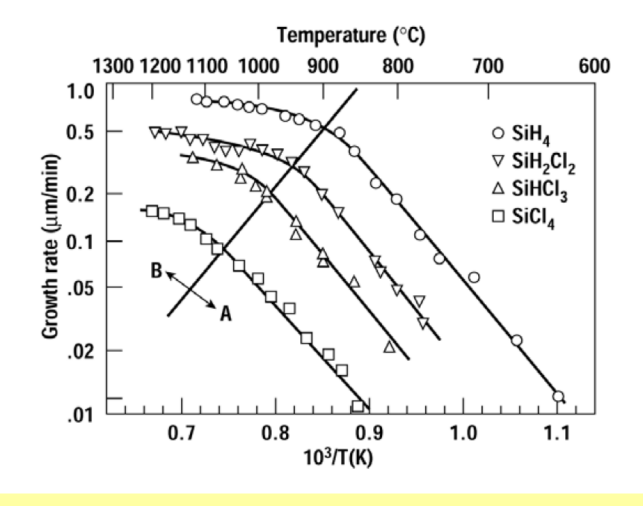
 $k_{\rm S} \ll h_{\rm G}$  = Surface reaction limited dy/dt ~ exp(-E<sub>a</sub>/kT)

 $h_G << k_S$  = Mass transport limited dy/dt ~ T<sup>3/2</sup>

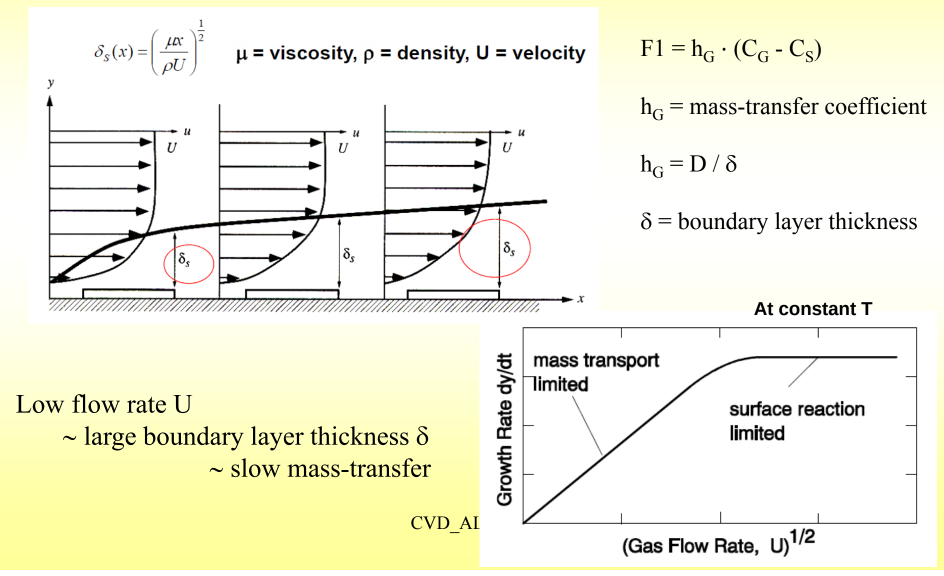
### **Deposition rate vs. Temperature**



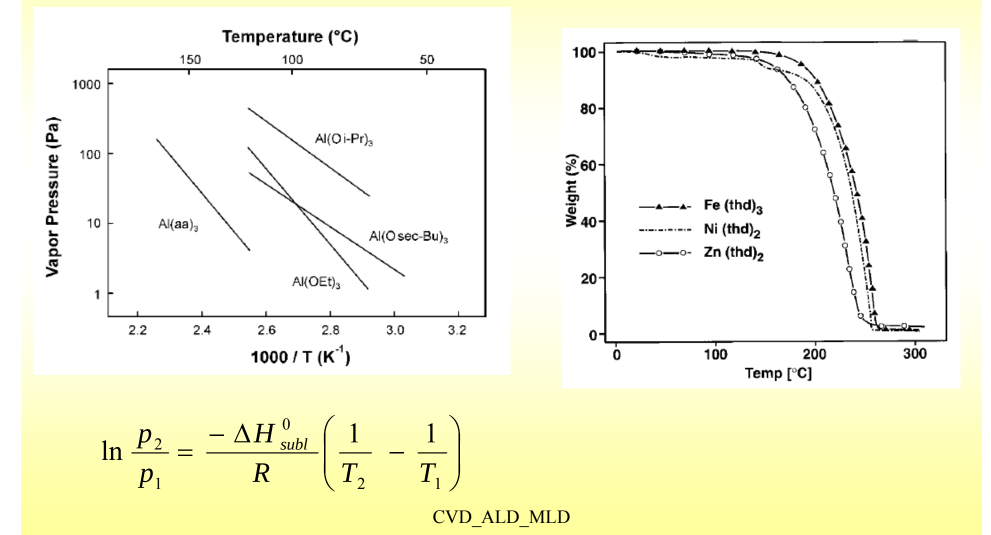
## **Deposition rate vs. Temperature**



## **Growth Rate Dependence on Flow Velocity**

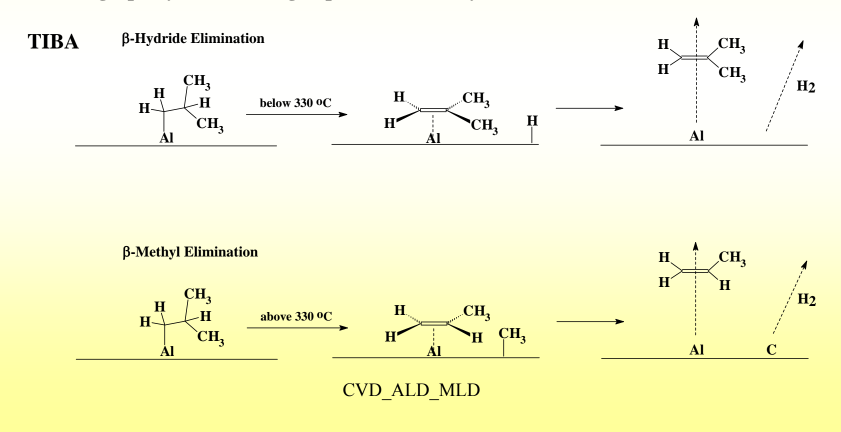


### **Precursor Volatility**



Aluminum

2.27 µΩcm, easily etched, Al dissolves in Si,
GaAs + Al → AlAs + Ga
Gas diffusion barriers, Al on polypropylene, food packaging = chip bags, party balloons, high optical reflectivity



Al deposits selectively on Al surfaces, not on SiO<sub>2</sub> Laser-induced nucleation 248 nm only surface adsorbates pyrolysed 193 nm gas phase reactions, loss of spatial selectivity control

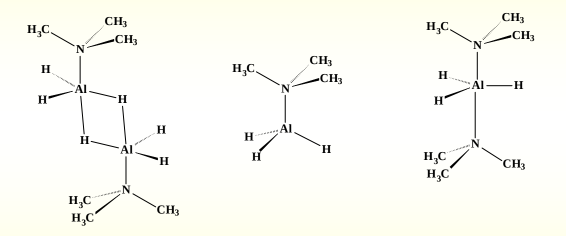
#### TMA

large carbon incorporation, Al<sub>4</sub>C<sub>3</sub>, RF plasma, laser

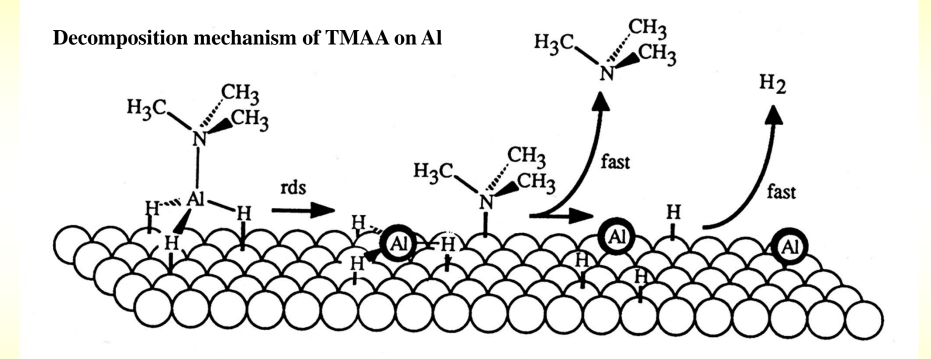
 $Al_2(CH_3)_6 \longrightarrow 1/2 Al_4C_3 + 9/2 CH_4$  under N<sub>2</sub>

 $Al_2(CH_3)_6 + 3H_2 \longrightarrow 2Al + 6CH_4$  under  $H_2$ 



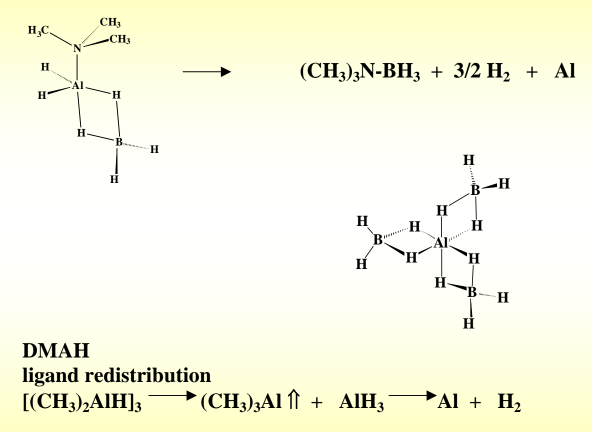


 $(CH_3)_3N-AlH_3 \longrightarrow Al + (CH_3)_3N + 3/2 H_2$  below 100 °C



 $(CH_3)_3N-AlH_3 \longrightarrow Al + (CH_3)_3N + 3/2 H_2$  below 100 °C

#### Aluminoboranes



at 280 °C, low carbon incorporation

Tungsten

5.6  $\mu\Omega$ cm, a high resistance to electromigration, the highest mp of all metals 3410 °C.

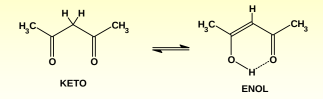
 $2 \operatorname{WF}_6 + 3 \operatorname{Si} \rightarrow 2 \operatorname{W} + 3 \operatorname{SiF}_4$ 

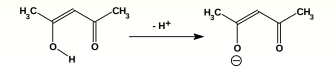
 $WF_6 + 3H_2 \rightarrow W + 6HF$ 

 $WF_6 + 3/2 SiH_4 \rightarrow W + 3 H_2 + 3/2 SiF_4$ 

 $W(CO)_6 \rightarrow W + 6 CO$ 

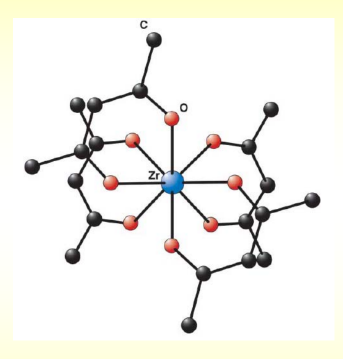
### **Diketonate Ligands**



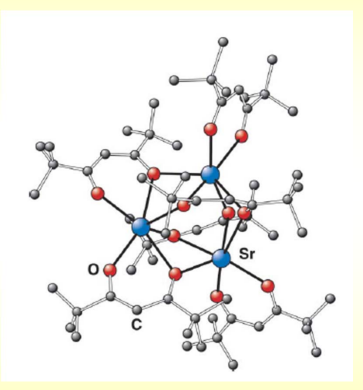


R <sub>1</sub>	R <sub>2</sub>	Name	Abbreviation
CH3	CH <sub>3</sub>	Pentane-2,4-dionate (acetylacetonate)	acac
CH3	CF3	1,1,1-trifluoropentane-2,4-dionate (trifluoroacetylacetonate)	tfac
CF3	$CF_3$	1,1,1,5,5,5-hexafluoropentane-2,4-dionate (hexafluoroacetylacetonate)	hfac
CH3	C(CH <sub>3</sub> ) <sub>3</sub>	1,1-dimethylhexane-3,5-dionate	dhd
C(CH <sub>3</sub> ) <sub>3</sub>	C(CH <sub>3</sub> ) <sub>3</sub>	2,2,6,6-tetramethylheptane-3,5-dionate	thd
CH3	CH <sub>2</sub> CH(CH <sub>3</sub> ) <sub>2</sub>	6-methylheptane-2,4-dionate	mhd
C(CH <sub>3</sub> ) <sub>3</sub>	CH <sub>2</sub> CH(CH <sub>3</sub> ) <sub>2</sub>	2,2,7-trimethyloctane-3,5-dionate	tmod
C <sub>6</sub> H <sub>5</sub>	C <sub>6</sub> H <sub>5</sub>	1,3-diphenylpropane-1,3-dionate (dibenyzoylmethanate)	dbm

## **Diketonate Precursors**



Mononuclear



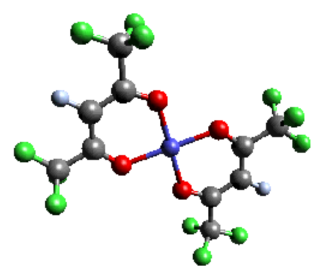
Polynuclear

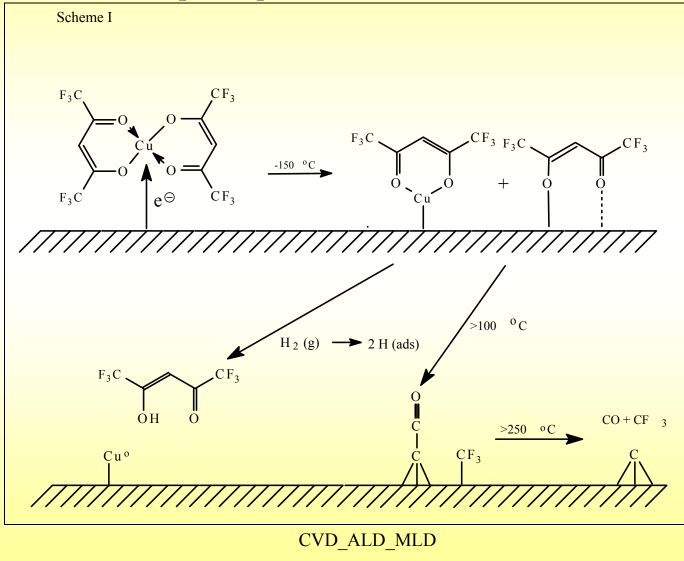
#### **Copper(II)** hexafluoroacetylacetonate

excellent volatility (a vapor pressure of 0.06 Torr at r. t.), low decomposition temperature, stability in air, low toxicity, commercial availability

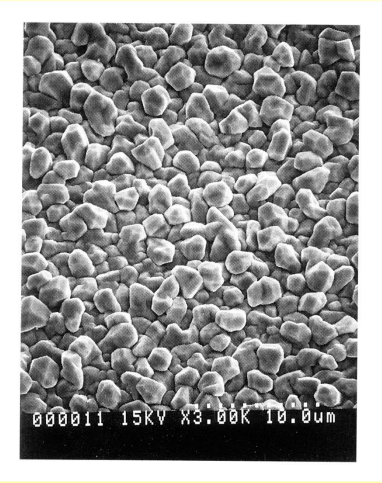
deposition on metal surfaces (Cu, Ag, Ta) the first step, which can already occur at -150 °C, a dissociation of the precursor molecules on the surface (Scheme I).

An electron transfer from a metal substrate to the single occupied HOMO which has an anti-bonding character with respect to copper  $d_{xy}$  and oxygen p orbitals weakens the Cu-O bonds and facilitates their fission.

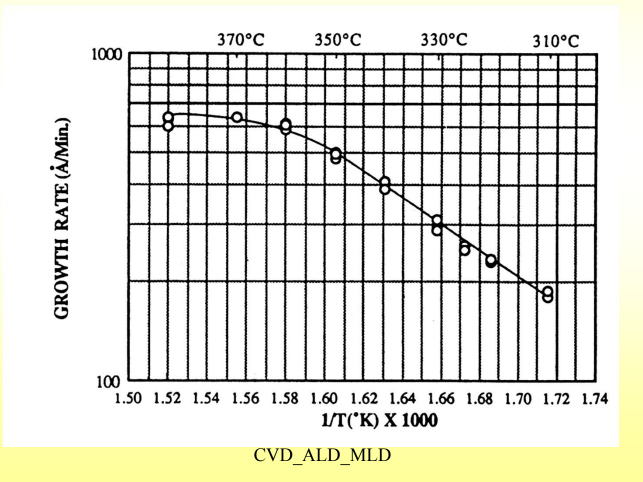




# SEM of Cu film, coarse grain, high resistivity



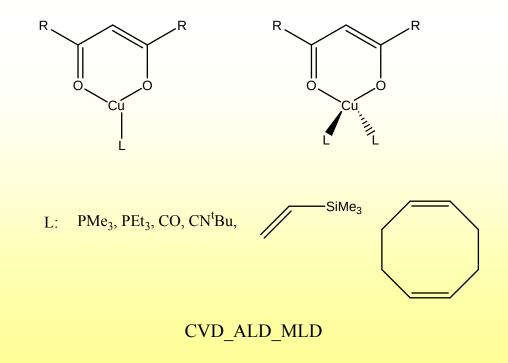
Growth rate of Cu films deposited from Cu(hfacac)<sub>2</sub> with 10 torr of H<sub>2</sub>



Cu(I) precursors

**Disproportionation to Cu(0) and Cu(II)** 

2 Cu(diketonate) $L_n \rightarrow Cu + Cu(diketonate)_2 + n L$ 



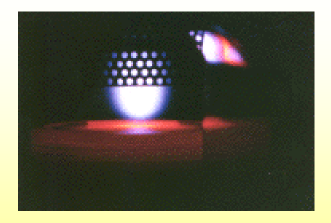
**Diamond films** 

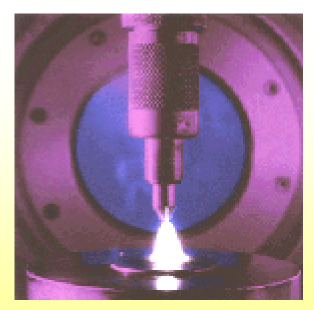
activating gas-phase carbon-containing precursor molecules:

•thermal (e.g. hot filament)

•plasma (D.C., R.F., or microwave)

•combustion flame (oxyacetylene or plasma torches)





**Experimental conditions:** 

temperature 1000-1400 K

the precursor gas diluted in an excess of hydrogen (typical  $CH_4$  mixing ratio ~1-2vol%)

**Deposited films are polycrystalline** 

Film quality:

•the ratio of sp<sup>3</sup> (diamond) to sp<sup>2</sup>-bonded (graphite) carbon

•the composition (e.g. C-C versus C-H bond content)

•the crystallinity

Combustion methods: high rates (100-1000 µm/hr), small, localised areas, poor quality films. Hot filament and plasma methods: slower growth rates (0.1-10 µm/hr), high quality films.

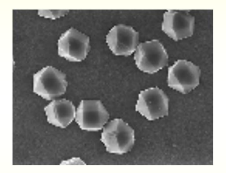
Hydrogen atoms generated by activation (thermally or via electron bombardment) H-atoms play a number of crucial roles in the CVD process:

H abstraction reactions with hydrocarbons, highly reactive radicals: CH<sub>3</sub> (stable hydrocarbon molecules do not react to cause diamond growth) radicals diffuse to the substrate surface and form C-C bonds to propagate the diamond lattice.

H-atoms terminate the 'dangling' carbon bonds on the growing diamond surface, prevent cross-linking and reconstructing to a graphite-like surface.

Atomic hydrogen etches both diamond and graphite but, under typical CVD conditions, the rate of diamond growth exceeds its etch rate whilst for graphite the converse is true. This is the basis for the preferential deposition of diamond rather than graphite.

Diamond initially nucleates as individual microcrystals, which then grow larger until they coalesce into a continuous film





Enhanced nucleation by ion bombardment: damage the surface - more nucleation sites implant ions into the lattice

form a carbide interlayer - glue, promotes diamond growth, aids adhesion

Substrates: metals, alloys, and pure elements:

Little or no C Solubility or Reaction: Cu, Sn, Pb, Ag, and Au, Ge, sapphire, diamond, graphite

C Diffusion: Pt, Pd, Rh, Fe, Ni, and Ti

the substrate acts as a carbon sink, deposited carbon dissolves into the metal surface, large amounts of C transported into the bulk,

a temporary decrease in the surface C concentration, delaying the onset of nucleation

**Carbide Formation:** Ti, Zr, Hf, V, Nb, Ta, Cr, Mo, W, Fe, Co, Ni, Y, Al B, Si, SiO<sub>2</sub>, quartz, Si<sub>3</sub>N<sub>4</sub> also form carbide layers. SiC, WC, and TiC

**Applications of diamond films:** 

Thermal management - a heat sink for laser diodes, microwave integrated circuits active devices mounted on diamond can be packed more tightly without overheating

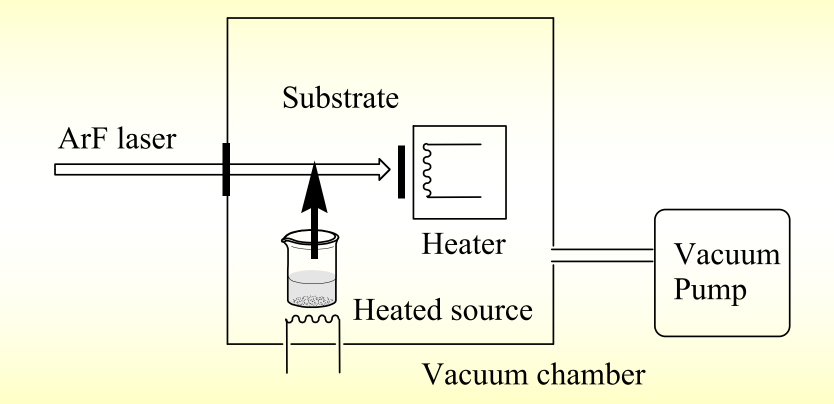
**Cutting tools** - an abrasive, a coating on cutting tool inserts CVD diamond-coated tools have a longer life, cut faster and provide a better finish than conventional WC tool bits

Wear Resistant Coatings -protect mechanical parts, reduce lubrication gearboxes, engines, and transmissions

**Optics - protective coatings for infrared optics in harsh environments,** ZnS, ZnSe, Ge: excellent IR transmission but brittle the flatness of the surface, roughness causes attenuation and scattering of the IR signal

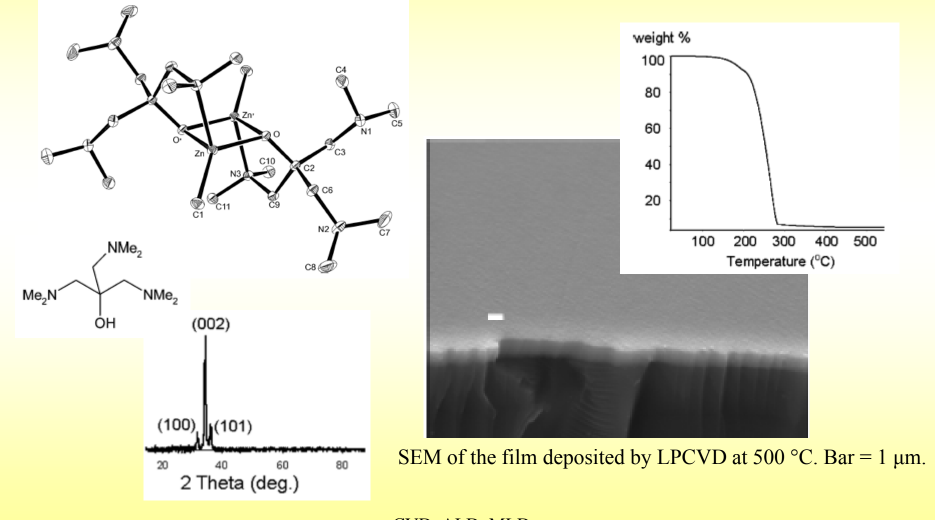
**Electronic devices** - doping, an insulator into a semiconductor *p*-doping: B<sub>2</sub>H<sub>6</sub> incorporates B into the lattice doping with atoms larger than C very difficult, *n*-dopants such as P or As, cannot be used for diamond, alternative dopants, such as Li

### **Laser-Enhaced CVD**



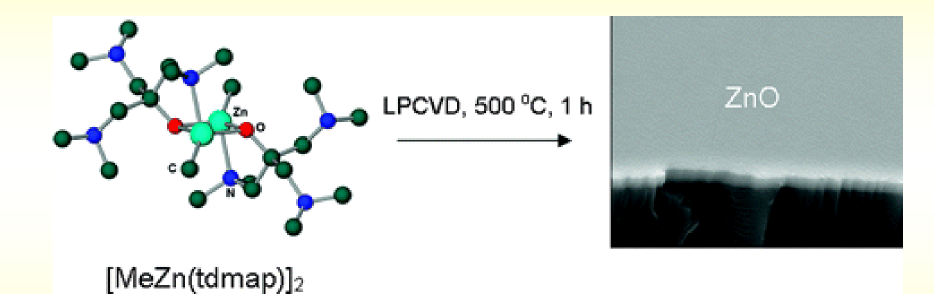
 $Si(O_2CCH_3)_4 \rightarrow SiO_2 + 2 O(OCCH_3)_2$ 

### LPCVD of ZnO from Aminoalcoholates

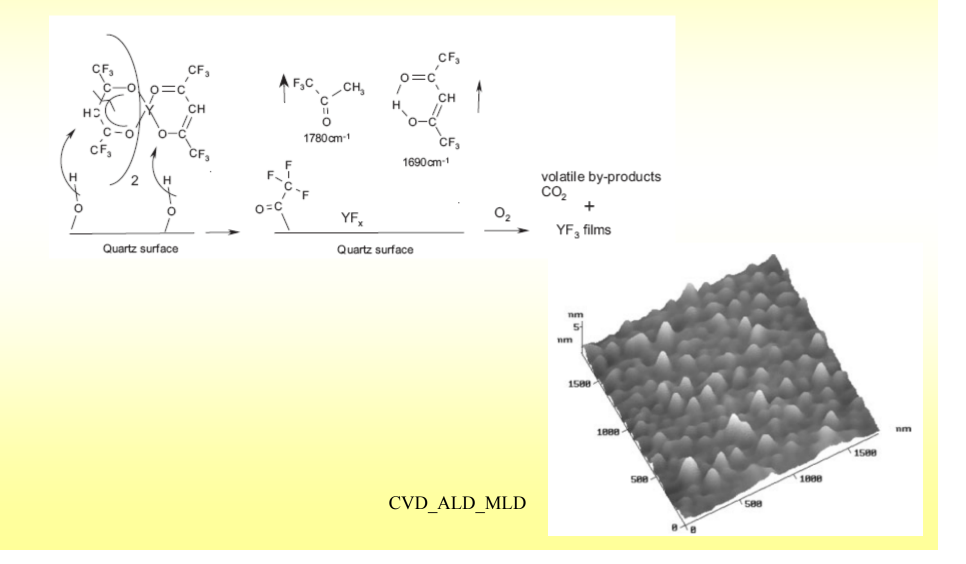


Hexagonal ZnO PDF 79-0208 CVD\_ALD\_MLD

### LPCVD of ZnO from Aminoalcoholates



### **CVD of YF<sub>3</sub> from hfacac Complex**

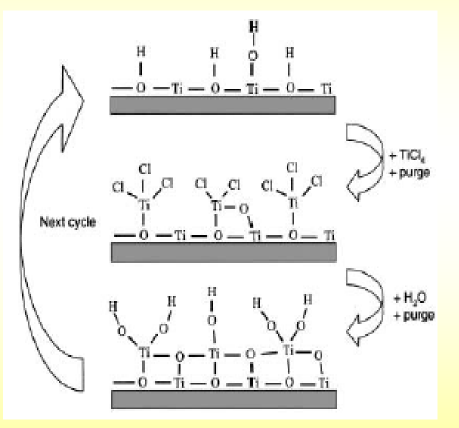


## **ALD Atomic Layer Deposition**

Special modification of CVD Method for the deposition of thin films Film growth by cyclic process

4 steps:

1/ exposition by1st precursor
 2/ cleaning of the reaction chamber
 3/ exposition by 2nd precursor
 4/ cleaning of the reaction chamber



# **ALD Atomic Layer Deposition**

Cycle repetitions until desired film thickness is reached

1 cycle: 0.5 s – several sec. thickness 0.1- 3 Å

Self-Limiting Growth Mechanism High reactivity Formation of a monolayer

**Control of film thickness and composition** 

**Deposition on large surface area** 

## **ALD vs. CVD Comparison**

**ALD Carried out at room temperature** 

**Control over number of deposited layers = film thickness** 

**Reactor walls inactive – no reactive layer** 

**Separate loading of reactive precursors** 

**Self-limiting growth** 

Precursor transport to the reaction zone does not have to be highly uniform (as in CVD)

**Solid precursors** 

## **ALD vs. CVD Comparison**

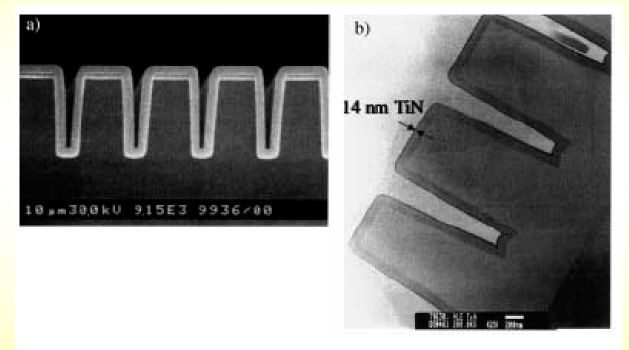


Figure 2. Cross-sectional SEM images for a 300-nm  $Al_2O_3$  film (a) and a 14-nm TiN film (b) deposited on a patterned silicon substrate.

# **Precursor Properties**

**Selection of suitable combination of precursors** 

**Molecular size influences film thickness** 

Gases, volatile liquids, solids with high vapor pressure

**Typical precursors:** 

<u>Metallic</u> - halogenides (chlorides), alkyls, alkoxides, organometallics (cyclopentadienyl complexes), alkyl amides

<u>Nonmetallic</u> - water, hydrogen peroxide, ozone, hydrides, ammonia, hydrazine, amines

## **Precursor Properties**

Thermally stable

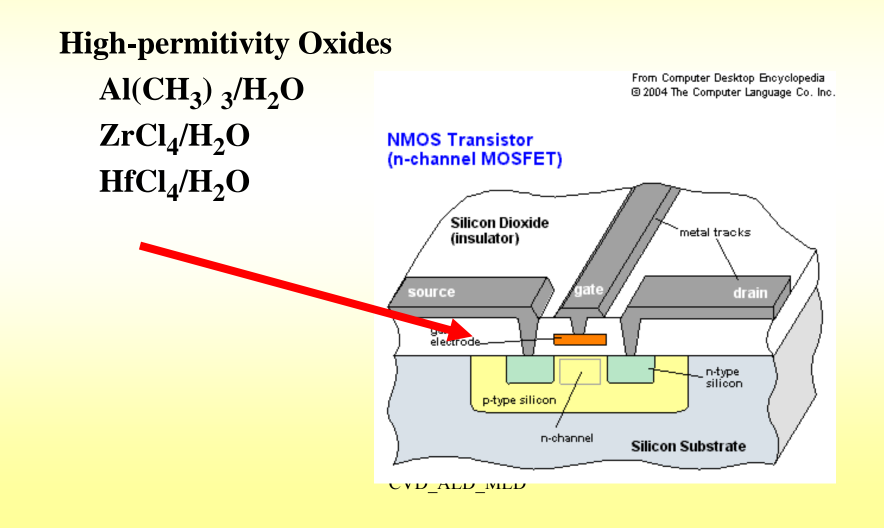
Must react with surface centers (hydroxyl groups on oxide surface)

Thermodynamics

**Kinetics** 

**Mechanisms** 

## **Examples of ALD**



## **Examples of ALD**

DRAM capacitors (Ba,Sr)TiO<sub>3</sub> – Sr and Ba cyclopentadienyl compounds and water as precursors

Nitrides of transition metals TiN - TiCl<sub>4</sub> and NH<sub>3</sub> TaN - TaCl<sub>5</sub>/Zn/NH<sub>3</sub> WN - WF<sub>6</sub> and NH<sub>3</sub> WC<sub>x</sub>N<sub>y</sub>

## **Examples of ALD**

Metallic films Difficult by ALD: metal surface has no reaction sites, low reactivity with reducing agents

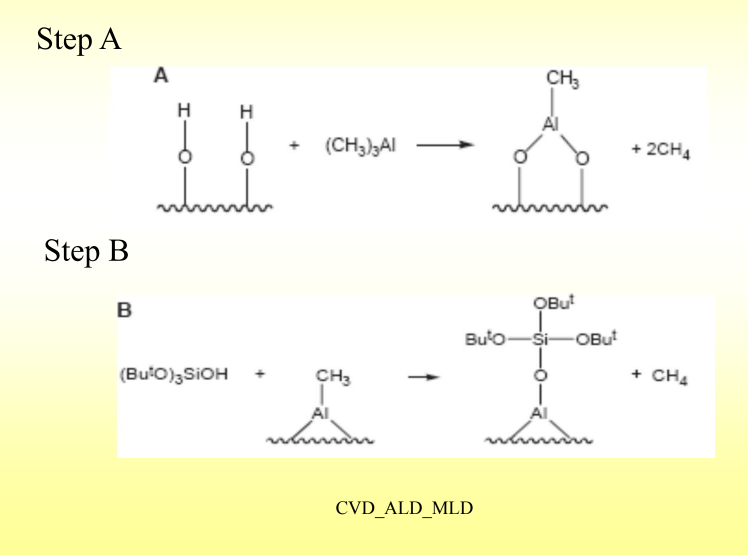
W - WF<sub>6</sub> and  $Si_2H_6$ 

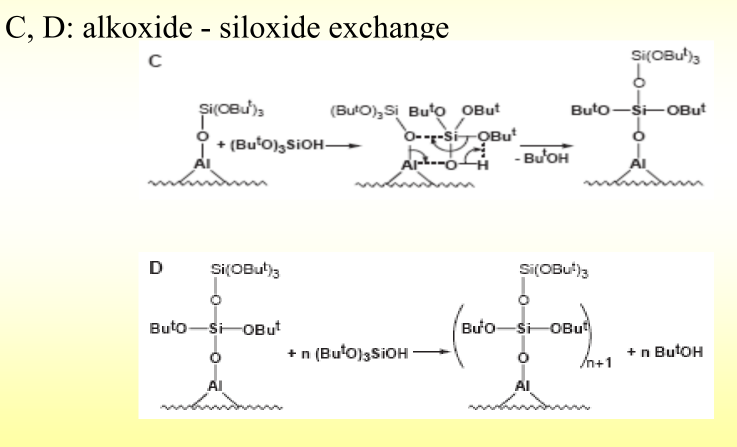
**Ru, Pt - organometallic precursors and oxygen** applies to all precious metals capable of catalytic dissociation of O<sub>2</sub>

- Ni, Cu metal oxide reduction by hydrogen radicals formed in plasma
- Al direct reduction of AlMe<sub>3</sub> by H radicals from plasma CVD\_ALD\_MLD

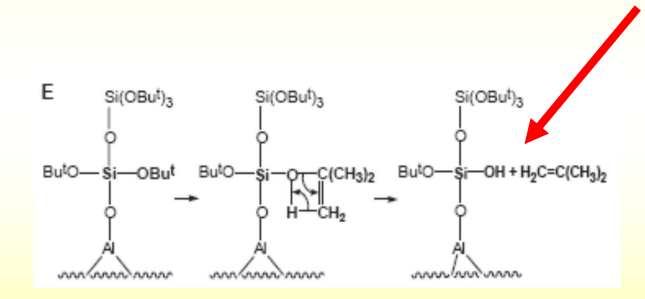
Precursors: trimethylalane, tris(tert-butoxy)silanol Deposition of amorphous SiO<sub>2</sub> and nanolaminates of Al<sub>2</sub>O<sub>3</sub> 32 monolayers in 1 cycle

Applications: microelectronics optical filters protective layers (against diffusion, oxidation, corrosion)

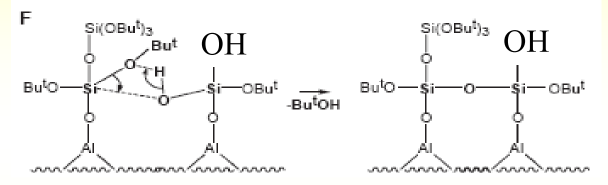




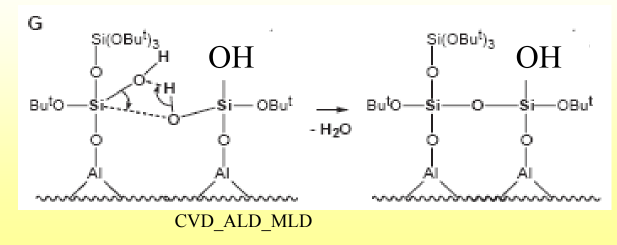
E: elimination of isobutene = formation of -OH



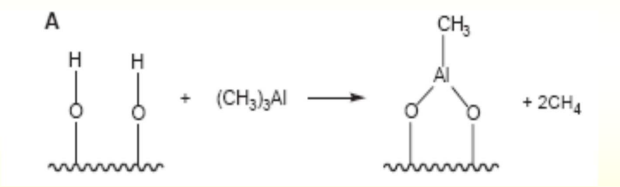
#### F: elimination of butanol = condensation

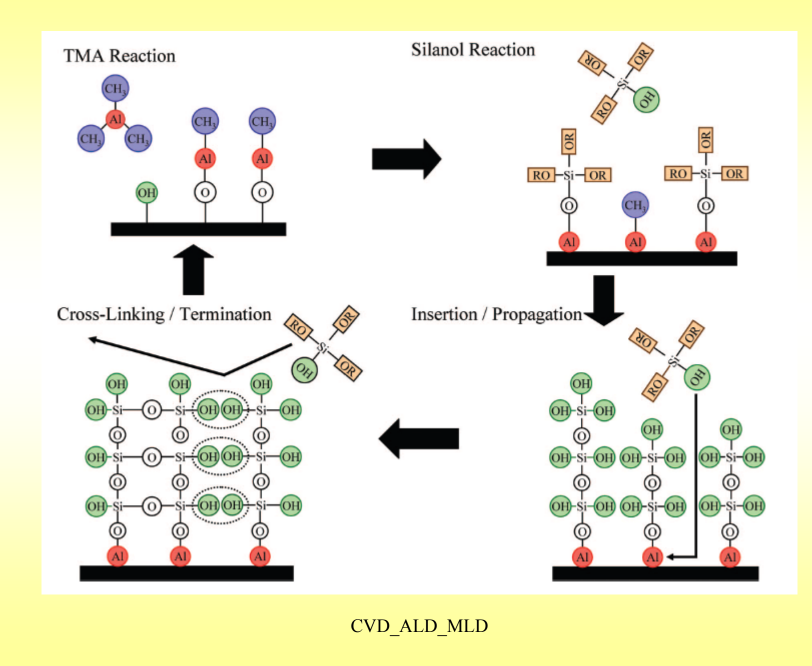


G: elimination of water = condensation

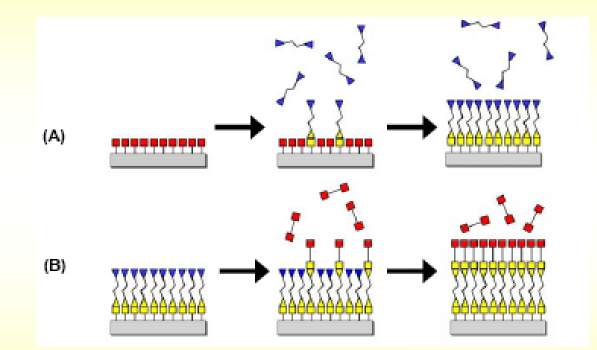


#### Repeat Step A



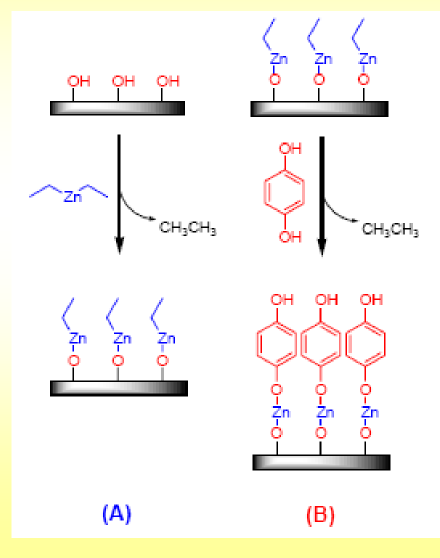


## **MLD - Molecular Layer Deposition**

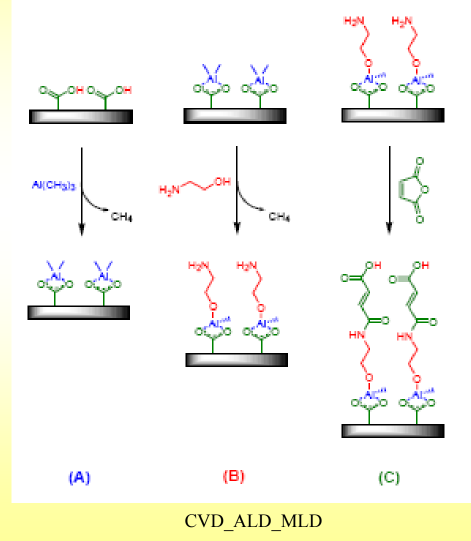


Sequential, self-limiting reactions A and B for MLD growth using two homobifunctional reactants

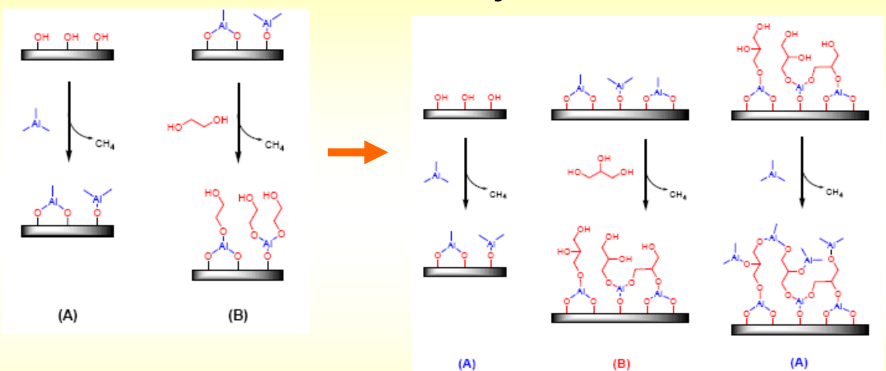
# **AB MLD**



# **ABC MLD**

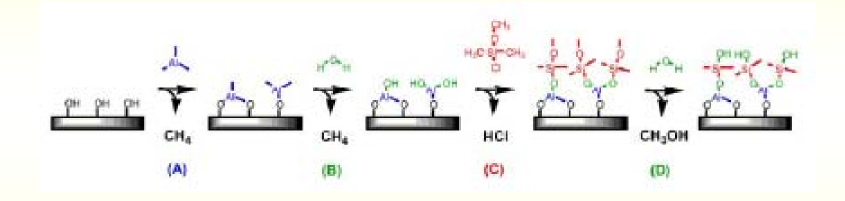


#### **Diols vs. Polyols**



homobifunctional precursors can react twice with the AlCH<sub>3</sub>\* surface species, double reactions lead to a loss of reactive surface sites and decreasing growth rate

# ABCD MLD growth of an alumina-siloxane



## **AB Lewis Acid-Lewis Base Reactions**

