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## 2.9 Advanced Wet Chemical Deposition Processes

(A) Electrochemical Deposition (ECD)

also: electroplating (EP) or electrodeposition or galvanic deposition

- (B) Electroless Deposition (ELD)
- 2.9.1 Repetition of Fundamentals on ECD
- 2.9.2 Application of Copper ECD in Damascene Interconnect Systems
- 2.9.3 Equipment for ECD in Microelectronics Front End Processing
- 2.9.4 Basics of Electroless Deposition
- 2.9.5 Electroless Deposition of Copper Diffusion Barriers



#### **Basics of ECD**

- If piece of metal immersed into solution containing its own ions a electrochemical double layer is formed: Helmholtz double layer
- Metal dissolves and ions from solution are deposited on metal
- Equilibrium potential  $E_{eq}$  is established given by NERNST-Equation:

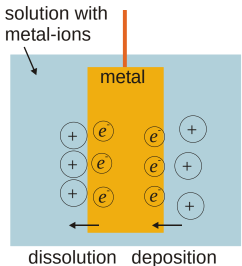
$$E_{eq} = E_0 + \frac{R \cdot T}{z \cdot F} \cdot \ln \frac{a_{Me^{z+}}}{a_{Me}}$$

 $E_0$ Standard electrode potential

T Temperature

F Faraday constant

- R Molecular gas constant
- Valency of ions Ζ
- Activity of metal/ions а resp.





## **Concept of electrochemical deposition**

- Two electrodes immersed into electrolyte and external voltage is applied
- Disturbance of equilibrium conditions at double layer
- Reactions that takes place correspond to a redox reaction
- Anodic Reaction: oxidation of metal dissolution into metal ions

$$Me \rightarrow Me^{z+} + z \cdot e^{-}$$

• Cathodic Reaction: reduction of metal ions - deposition of metal

$$Me^{z^+} + z \cdot e^- \rightarrow Me$$

• Resulting potential:

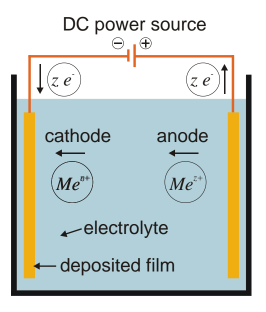
$$E = E_c - E_a + I \cdot R_S$$

I Current

*R*<sub>S</sub> Resistance of electrolyte

 $E_{o}$ ,  $E_{a}$  Potential at anode and cathode resp., established

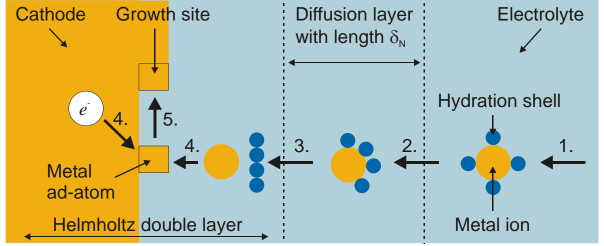
different from  $E_{\rm eq}$  due to overpotentials





#### Reaction mechanisms at cathode

- 1. Hydrated metal ions (ions with hydration shell) reach the cathode surface due to transportation mechanisms and enter the diffusion layer
- 2. Ions pass diffusion layer due to concentration gradient and hydration shell aligns with the field strength that increases towards the cathode surface
- 3. Ions enter Helmholtz double layer and hydration shell is stripped off due to high field strength in this layer
- 4. Reduction of ions with electrons from the cathode to form metal atoms that adsorb at the metal surface (ad-atoms)
- 5. Diffusion of ad-atoms to a energetic favorable growth sites and integration into the metal crystal lattice



Reaction mechanisms at cathode [1]



#### **Overvoltage**

- Potentials of electrodes  $E_a$ ,  $E_c$  deviate from the equilibrium potential  $E_{eq}$  by a overvoltage  $\eta$
- This overvoltage is caused by the inhibition of each single reaction step
- The overall overvoltage is:

$$\eta = \eta_{tr} + \eta_{diff} + \eta_{re} + \eta_{crys} + \eta_{res}$$

$\eta_{tr}$	charge transfer ov	$\eta_{diff}$	diffusion ov	$\eta_{\sf re}$	reaction ov
$\eta_{crvs}$	crystallization ov	$\eta_{res}$	resistance ov		

#### Deposition rate *r*

• Derived from 1. Faraday's law:

$$m = \frac{M \cdot I \cdot t}{z \cdot F} \implies t_f = \frac{M \cdot I \cdot t}{A \cdot \rho \cdot z \cdot F} \implies r = \frac{M \cdot J}{\rho \cdot z \cdot F}$$

m	Mass	M	Molar mass	1	Current
t	Deposition time	Z	Valency	F	Faraday constant
$t_f$	Film thickness	Α	Active area	$\rho$	Density
j	Current density	r	Deposition rate		



## **Transport mechanisms**

- Convection
  - Forced convection induced by agitating the bath (stirring, air bubbling)
  - Natural convection due to density gradients caused by depletion at cathode surface and due to gravitation (minor impact)
- Diffusion
  - By concentration gradient in diffusion layer caused by depletion of copper ions at the cathode surface because of the reduction reaction
  - Flux through diffusion follows Fick's law
- Migration
  - Movement of copper ions within the electrical field between the electrodes
  - Can be neglected, since conc. of copper ions < conc. sulfur ions
- Overall flux is sum of flux of each mechanism:

$$j = \underbrace{(c \cdot v)}_{convection} + \underbrace{\left(-\frac{D}{\delta_{N}} \cdot \nabla c\right)}_{diffusion} + \underbrace{\left(-z \cdot \mu \cdot F \cdot c \cdot \nabla \varphi\right)}_{migration}$$

С	Ion concentration	V	Flow velocity	D	Diff. Coefficient
$\delta_{\!N}$	Length of Difflayer	Z	Valency	μ	Ion mobility
F	Faraday constant	$\varphi$	Potential		



# **2.9 Advanced Wet Chemical Deposition Processes**

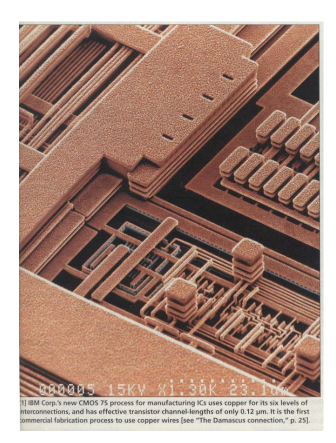
2.9.1	Repetition of Fundamentals on ECD
2.9.2	Application of Copper ECD in Damascene Interconnect Systems
2.9.3	Equipment for ECD in Microelectronics Front End Processing
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#### **Application of Copper ECD in Damascene Interconnects**

#### **Background**

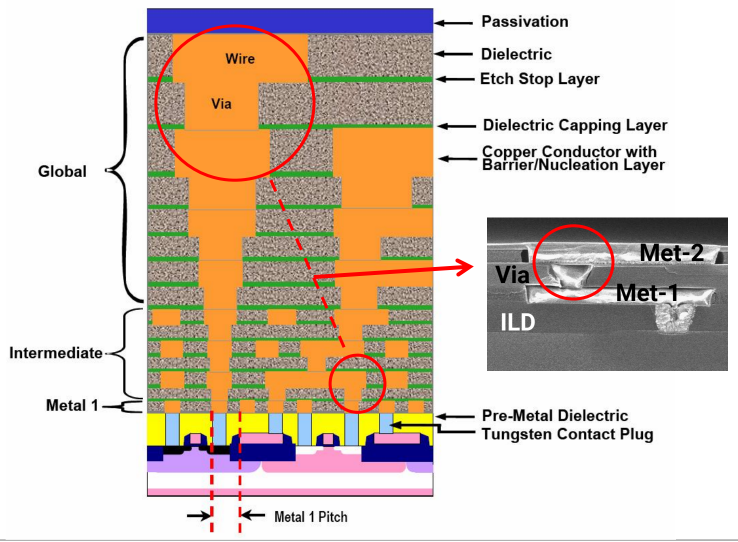
- Copper has replaced Aluminum in recent years because of higher conductivity and better electromigration resistance
- Copper ECD has better ability to fill high aspect ratio trenches compared to PVD and CVD and moreover its less expansive
- Filling of multilayer structures in one step can be done (Dual Damascene)



CMOS interconnects [IBM]

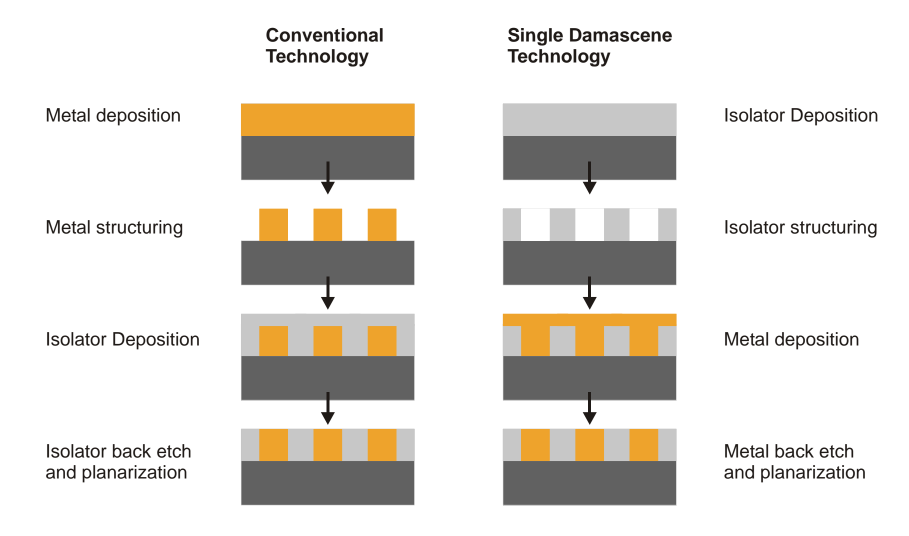


#### **Application of Copper ECD in Damascene Interconnects**





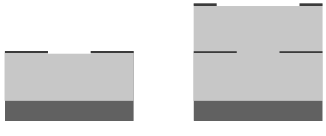
## **Basic Principle**



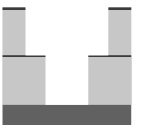


## **Dual Damascene Technology**

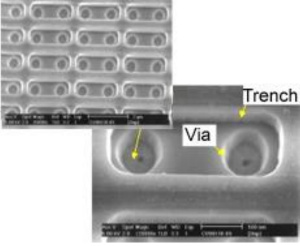
- Vias and trenches filled in single step ECD
- Barrier (Ta/TaN, Ti/TiN) and Cu-seed deposition e.g. by ionized PVD
- CMP for planarization



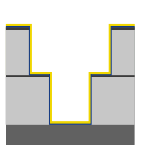
Isolator deposition and Hardmask deposition / Structuring (Layer 1+2)



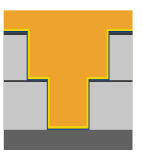
Isolator structuring



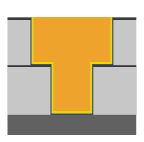
Trench and Vias [IMEC]



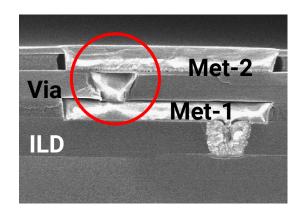
Barrier and Seed layer deposition



Cu deposition



CMP





## **Basic requirements for Copper electroplating**

- Plating Cell
  - Simplest form: Beaker (research, initial tests)
  - Advanced semiconductor plating tool (Fountain Plater)
  - Rack Plater for PCB-industry
- Electrolyte
  - copper sulfate, conducting salt
  - Additives
- Convection: Stirrer or circulation Pump
  - Electrolyte movement
  - Wafer rotation or lateral movement
- Anode
  - soluble copper anode
  - inert anode (Ti+Pt)
- Cathode (piece to be deposited => i.e. Wafer)
  - Seed Layer (e.g. Cu) + barrier (e.g. TiN) + adhesion layer (e.g. Ti)
  - Wafer holder
- Power Supply (DC or Pulsed)



## **Principle of acid copper plating**

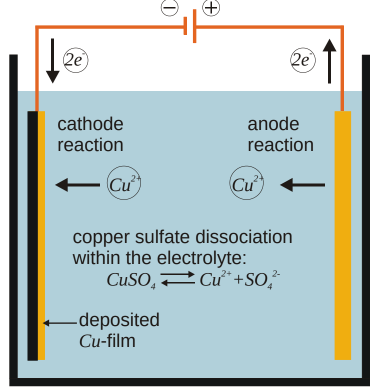
• Electrolyte is a dissociated copper sulfate salt:

$$CuSO_4 \xrightarrow{\longleftarrow} Cu^{2+} + SO_4^{2-}$$

• Redox reaction for acid copper plating is:

$$Cu \xrightarrow{Ox.} Cu^{2+} + 2 \cdot e^{-}$$

- Anode types:
  - Soluble copper anode (provides cupric ions)
  - non soluble inert anode (ions are added from external source)
- Seed layer: mostly Cu
- Additives to control film properties (grain size, ductility, stress, brightness)



Copper deposition with soluble anode



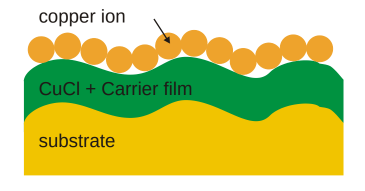
# Composition of a acid copper plating bath

	Component	Chemical basis	Meaning
<u> </u>	Copper sulfate	Metal salt	- Provision of cupric ions
Basic	Sulfuric acid	Acid	- Increase in conductivity
	Chlorine	 Halogen	- Weak inhibition of deposition
			- Necessary for operation of other additives
	Carrier	Polyether	- Inhibition of deposition
	(suppressor)	(PEG, PAG)	- Uniformity of deposit
Additives	Brightener	Organic Sulfur	- Acceleration of deposition
	(accelerator)		- Bright deposits
490		(SPS, MPS)	- Incorporation in deposit
			- Weak grain refiner
	Leveler	Organic Nitrogen	- Strong inhibition of deposition
	C	compound (JGB)	- Smooth surfaces (leveling)
			- Strong grain refiner



## **Effect of plating additives**

- Chlorine (CL)
  - · CL is a surface active complexing agent
  - CuCl-film is formed at the surface
  - · Film slightly inhibits reaction
  - · Required for action of other additives





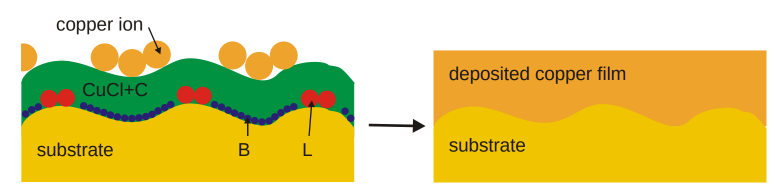
- Suppressor or Carrier (C)
  - · Adsorption of C promoted by CL
  - Inhibition of Cu deposition (suppresses current density)
  - Deposited film has even thickness (uniformity)

deposited copper film substrate

Deposition with CL and Carrier



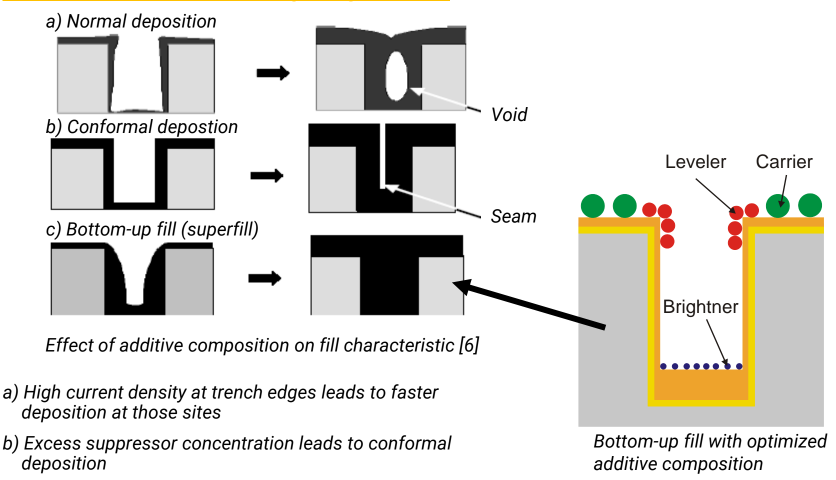
- Accelerator or Brightener (B)
  - Copper and B form Cu-thiolate film
  - Film interacts with CL to accelerate copper ion reduction ( $C^{2+} \rightarrow Cu$ )
  - Grain refinement → bright deposit (mirror like films)
- Leveler (L)
  - Inhibition of copper deposition
  - Adsorbs preferentially ad peaks and valleys thus levels the deposit
  - · Refines grain structure



Deposition with optimized additive composition



#### Fill characteristics - effect of plating additives



fill characteristic

b) Optimized composition of additives leads to perfect trench-



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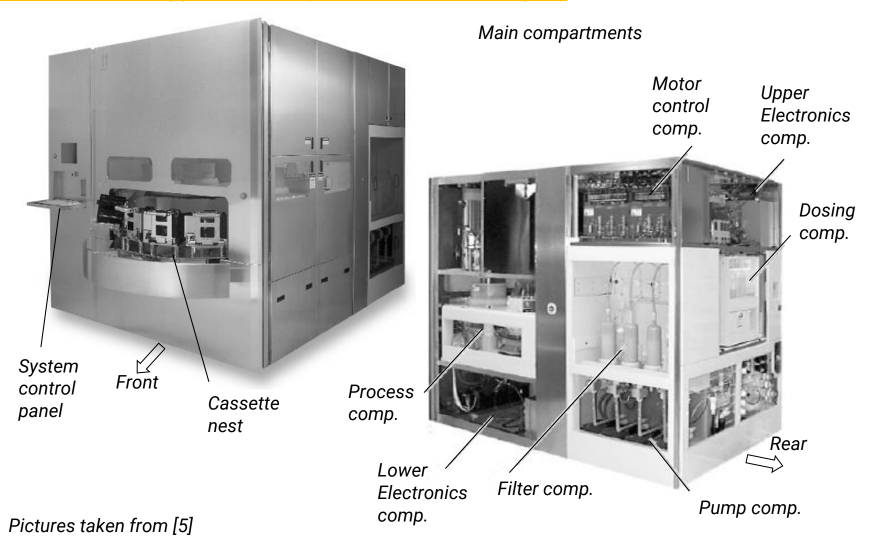
#### **Equipment for Copper ECD**

#### **Basic facts**

- Pre/post treatment cells for cleaning, rinsing and drying
- Plating cells:
  - Contain plating solution (cupric ion solution)
  - (Soluble) Anode
  - Cathode i.e. workpiece (Wafer) in a holder
- Two basic concepts of plating cells:
  - Vertical cathode-anode-arrangement used in older PCB-plating tools for high troughput
  - Horizontal cathode-anode-arrangement in new PCB tools and in Semiconductor fabrication for more precise deposition processes
- Main manufacturers of electroplating tools:
  - Lam Research (former Novellus), AMAT (former Semitool)
  - Semsysco, Classone (spin off from Semitool); Tel Nexx (f. Nexx Systems)
- Bath chemistry suppliers:
  - Enthone (Cubath Viaform), Atotech, BASF

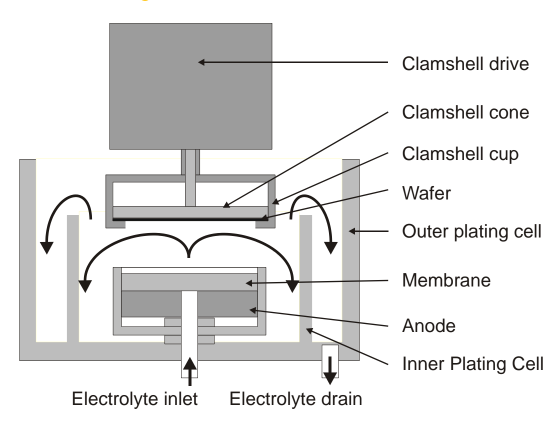


#### **Semiconductor Copper ECD tool (Novellus Sabre Concept II)**





#### **Fountain Plating Cell**



Principle of fountain plating cell



Plating cells and post plating cells in the process compartment [5]



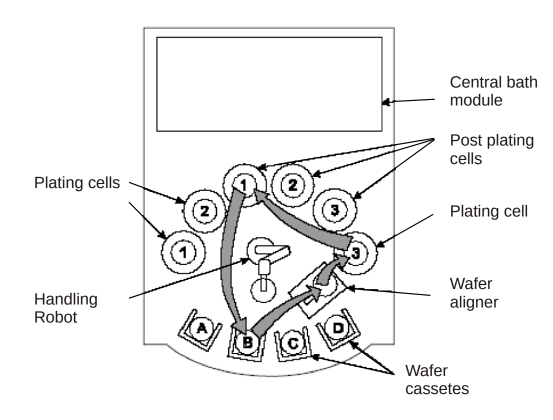
Clamshell [5]





#### **Process flow**

- Wafer cassettes are placed in the cassette Load port
- Wafer picked by handling robot out of cassette and placed into wafer aligner
- Centering and alignment of wafer for processing
- Handling robot places wafer into clamshell of plating cell
- Wafer is submerged into the plating solution and DC current is applied
- After processing the wafer is transported to a post plating cell for cleaning, rinsing and drying
- · Return of wafer to cassette



Schematic of process flow [5]

Feed lines for DI

Delivery system for Additives

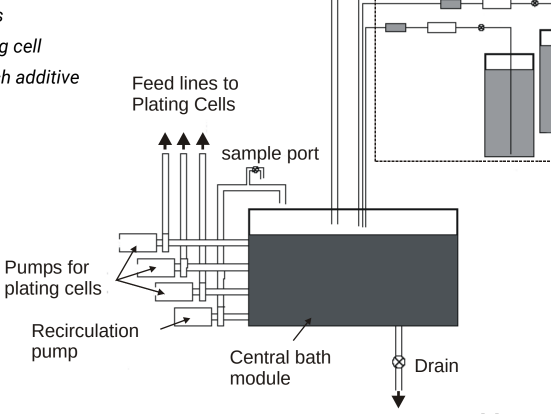
Water and Plating

Additives



## **Dosing System**

- Central bath module: 150l tank contains plating solution
- Recirculation pump for proper mixing of the electrolyte
- Sample port for analysis
- 3 pumps for each plating cell
- Separate delivery of each additive





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## **Electroless Deposition**

#### **Fundamentals**

- · Deposition process without any external power supply
- Two types of electroless deposition processes:
- 1. Reduction process
  - Reducing agent contained in the electrolyte
  - Electrons provided by oxidation of the reducing agent

$$Me^{z^+} + Re d \rightarrow Me + Ox$$
Metalion Reducing Metal Oxidized form of Reducing agent

- 2. Ion exchange process
  - Substrate surface has more negative standard potential
  - Electrons from exchange reaction with substrate

Oxidation:

Reduction:

$$Me_1 \to Me_1^{z+} + z \cdot e^- \qquad Me_2^{z+} + z \cdot e^- \to Me_2$$

- Mostly for protection of bare metals: e.g. Zn, Sn on steel; Au on brass
- · Only thin films because of self-inhibiting effect



## Typical chemistry for the reduction process type

Chemistry	Meaning
Metal salt	- Provision of metal ions (e.g. sulfate based)
Complexing agent	- Prevents precipitation of metal salt
Reducing agent	<ul> <li>Supplies electrons for metal ion reduction (e.g. formaldehyde HCHO or hypophosphite H<sub>2</sub>PO<sub>2</sub>-)</li> </ul>
Activator	- Supports/activates oxidation of red. Agent (e.g. Pd)
Buffer	- pH maintenance (e.g. NaOH)
Stabilizers	- Retards homogeneous reactions (e.g. thiourea)
Accelerators	- Increase in deposition rate
Surfactants	- Wetting agent
	- Removal of evolved gases

# Potential Applications of ELD in Damascene Technology

- Alternative method for Cu seed layer deposition (conformal deposition)
- Deposition of cap layers on Cu lines
- Also alternative method for Cu trench superfill



# **Examples for the reduction process type**

- Nickel deposition
  - Deposition in alkaline or acidic solution with nickel provided from NiSO<sub>4</sub>-salt
  - Complexing agents: organic acid or amine (ethylene diamine, citrate acid)
  - Reducing agent: hypophosphite (e.g. Sodium-hypophosphite: NaH<sub>2</sub>PO<sub>2</sub>)
  - Buffer for pH-maintenance is e.g. NaOH
  - Basic reaction steps

Oxidation of red. agent: 
$$H_2PO_2^- + H_2O \rightarrow H_2PO_3^- + 2 \cdot H^+ + 2 \cdot e^-$$

Reduction of Ni ions: 
$$Ni^{2+} + 2 \cdot e^{-} \rightarrow Ni$$

- Copper deposition
  - Mostly alkaline solutions with CuSO<sub>4</sub> as metal salt
  - Complexing agents
  - Reducing agents mostly formaldehyde (HCHO)
  - Reaction steps:

Oxidation of red. agent: 
$$HCHO + 3OH^- \rightarrow HCOO^- + 2 \cdot H_2O + 2 \cdot e^-$$

Reduction of Cu ions: 
$$Cu^{2+} + 2 \cdot e^{-} \rightarrow Cu$$



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2.9.1	Repetition of Fundamentals on ECD



#### **Electroless deposition of copper diffusion barriers**

#### **Background**

- Conventional barrier deposition shows several drawbacks for depositions in narrow trenches (non defect free, pinches off at trench top)
- Several process steps (blanket deposition, patterning)

#### Advantages of electroless deposition

- Conformal deposition even within complex structures (narrow trenches)
- Deposition on non-conductive materials
- Possibility of area selective plating as cap layers on Cu interconnects reduced process steps

#### Possible materials

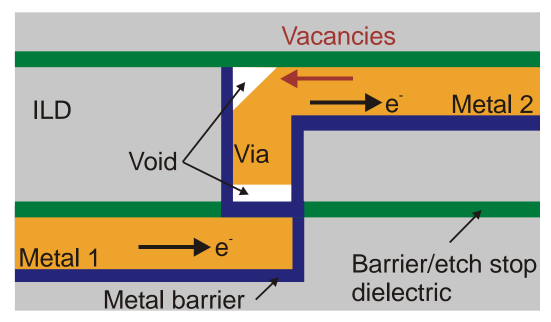
- Ni alloys (NiMoP)
- Co alloys (CoWP, CoWB)





#### **Electroless CoWP**

- Used as barrier metal and capping barrier layer
- Co forms metal-metal bonds with Cu while W and P stuff grain boundaries to inhibit Cu-diffusion
- CoWP has strong adhesion to copper and prevents Cu-diffusion/vacancy-movement and thus prevents void formation
- Copper reliability improvements through CoWP capping layers
- Device performance improvement due to lower k<sub>eff</sub> compared to previous capping layers (SiN, SiC, SiCN)

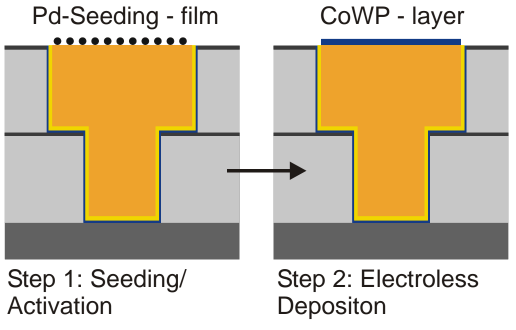


Void formation through weak Cu-barrier/etch stop layer adhesion [10]



#### **Electroless CoWP - Process**

- Usually 90% Co, 2% W and 8% P
- Selective deposition onto Cu-lines after Pd-Activation (seeding) Capping layer deposition with fewer process steps



Electroless CoWP as capping layer - Process steps

 One to two orders of magnitude increased electromigration lifetime of capped Cu interconnects compared to non-capped ones



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