

### **Growth and Pattern Formation for Epitaxial Surfaces**

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# Outline

- Strain in epitaxial systems
  - Leads to structure
  - Quantum dots and their arrays
- Atomistic strain model
  - Lattice statics model
  - Lattice mismatch
- Numerical methods
  - Algebraic multigrid (AMG)
  - Artificial boundary conditions (ABC)
- Application to nanowires and nanocrystals
  - Step bunching instability
- Summary

# Strain in Epitaxial Systems

- Lattice mismatch leads to strain
  - Heteroepitaxy
  - E.g., Ge/Si has 4% lattice mismatch
- Relief of strain energy can lead to geometric structures
  - Quantum dots and q dot arrays

# Quantum dots and Q Dot Arrays



Ge/Si, Mo et al. PRL 1990



Si<sub>.25</sub>Ge<sub>.75</sub>/Si, (5 μm)<sup>2</sup> MRSEC, U Wisconsin

# Directed Self-Assembly of Quantum Dots

- •Vertical allignment of q dots in epitaxial overgrowth (left)
- Control of q dot growth over mesh of buried dislocation lines (right)



B. Lita et al. (Goldman group), APL 74, (1999)





H. J. Kim, Z. M. Zhao, Y. H. Xie, PRB 68, (2003).

In both systems strain leads to ordering! CSCAMM, 4/24/2007

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### **Atomistic Modeling of Strain in Thin Films**

- Lattice statics for discrete atomistic system,
  - minimize discrete strain energy (Born & Huang, 1954)
  - Application to epitaxial films, e.g.
    - E.g., Stewart, Pohland & Gibson (1994), Orr, Kessler, Snyder & Sander (1992),
- Idealizations
  - Harmonic potentials, Simple cubic lattice
  - General, qualitative properties
    - Independent of system parameters
  - Computational speed enable additional physics & geometry
    - 3D, alloying, surface stress
- Atomistic vs. continuum
  - atomistic scale required for thin layer morphology
    - strain at steps
  - continuum scale required for efficiency
    - KMC requires small time steps, frequent updates of strain field

## Microscopic Model of Elasticity with Harmonic Potentials Continuum Energy density - isotropic $E = \lambda (S_{xx} + S_{yy})^2 + \mu (S_{xx}^2 + S_{yy}^2 + 2S_{xy}^2)$ - cubic symmetry $E = \alpha (S_{xx}^2 + S_{yy}^2) + \beta S_{xy}^2 + \gamma S_{xx} S_{yy}$ • Atomistic Energy density - Nearest neighbor springs

$$E = k(S_{xx}^{2} + S_{yy}^{2})$$

- Diagonal springs  $E = \ell (S_{xx} + 2S_{xy} + S_{yy})^2 + \ell (S_{xx} - 2S_{xy} + S_{yy})^2$
- Bond bending terms

$$E = mS_{xy}^{2}$$

• Elastic equations  $\partial_u \mathbf{E} [\mathbf{u}] = \mathbf{0}$ 

## **Strain in an Epitaxial Film Due to Lattice Mismatch**



- lattice mismatch  $\bullet$ 
  - lattice constant in film a
  - lattice constant in substrate h
- CSCAMM, 4/24/2007 relative lattice mismatch  $\epsilon$ =(a-h)/h

# Deformation of Surface due to Intrinsic Surface Stress

Surface stress included by variation of lattice constant for surface atoms

No misfit in film

film misfit

## Strain Tensor Step with No Surface Stress



# 

#### Strain Tensor Step with Surface Stress No lattice mismatch







# **Interaction of Surface Steps**

- Steps of like "sign"
  - Lattice mismatch  $\rightarrow$  step attraction
  - Surface stress  $\rightarrow$  step repulsion





Step attraction due to lattice mismatch

Repulsion of nearby steps due to surface stress

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## Numerical method for Discrete Strain Equations

- Algebraic multigrid with PCG
- Artificial boundary conditions at top of substrate
  - Exact for discrete equations
- 2D and 3D, MG and ABC combined
- Russo & Smereka (JCP 2006), Lee, REC & Lee (SIAP 2006), REC, et al. (JCP 2006)



### Multigrid





CPU speed (sec) vs. lattice size for strain computation in a 2D quantum dot system

Strain energy density for 160 atom wide pyramid in 2D with trenches, for various trench depths



#### **Artificial Boundary Conditions**



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### Nanowires

- Growth catalyzed by metal cluster (Au, Ti, ...)
- Epitaxial
- Application to nano-electronics
- Stability difficulties

# 

### **Ti-Nucleated Si Nanowires**

Kamins, Li & Williams, APL 2003



FIG. 1. (a) Plan-view and (b) cross-sectional scanning-electron micrographs of Ti-nucleated Si nanowires (60 min growth) after annealing in  $\rm H_2$  at 850 °C for 1 h,



#### Nanowire



InP wire
20nm Au cluster
at tip
Scale bar =5 nm
Oxide coating,
Not present
during growth
TEM

Gudiksen, Wang & Lieber. JPhysChem B 2001

## Instability in Metal Catalyzed Growth of Nanowires

- Epitaxial structure
  - Tapered shape due to side attachment
- Instability at high temperature
  - Tapered shape → terraced shape
  - Step bunching



1μm



Kamins, Li & Williams, APL 2003

### **D UCIA 2D** Simulation of Nanowires

- 2 steps looking for step bunching by energy minimization
- Model
  - Harmonic potential
  - Surface stress
- No step bunching in 2D



#### Strain Energy vs. distance between steps

### **3D Simulation of Interaction between Steps on Nanowires**

- Homogeneous, epitaxial nanowire with surface stress
- Interactions of two steps
  - $r = R_1$  for  $z < z_1$
  - $r = R_2$  for  $z_1 < z < z_2$
  - $r = R_3 \text{ for } z_2 < z$
  - $L=z_2-z_1 = inter-step distance$
  - z = axial distance, r = wire radius
- Energy minimum occurs for small L
  - Step bunching
- Results are insensitive to parameters
  - Step size  $(R_1 R_2 \text{ or } R_2 R_3)$
  - Surface stress
  - Wire radius, shape
- Lowest value of energy E occurs for small value of separation L
  - System prefers bunched steps



 $(R_1, R_2, R_3) = (3,4,5)$ 





## Nanocrystals

- Sphere or rod (diameter =10 50 nm)
- Coated by shell (thickness = 1 10 nm)
- Epitaxial structure
- Wide range of new properties and applications
- Difficulty with instability of shell due to strain

#### Epitaxial Nanocrytals core/shell=CdSe/CdS R<sub>core</sub>=34Å, R<sub>shell</sub>=9Å



#### HRTEM

<sup>100 Å</sup> Peng, Schlamp, Kadavanich, Alivisatos. JACS 1997



#### Epitaxial Nanorods core/shell=CdS/ZnS



Left to right
Increasing shell thickness
Epitaxial structure breaks
down at larger shell size
HRTEM

#### 10 nm Manna,Sher, Li, Alivisatos. JACS 2002

## Simulation of Strain Field in Nanocrystals

- Core  $0 < r < R_{core}$
- Shell  $R_{core} < r < R_{shell}$
- Strain model
  - Harmonic potential
  - Equal elastic parameters
  - lattice mismatch
  - No surface stress
- Max energy density occurs at critical shell thickness
- Critical shell thickness is at peak in photoluminescence
- Robust results: same in 2D, variation of parameters





### Comparison to Critical Size for Photoluminescence





Peng, Schlamp, Kadavanich, Alivisatos. JACS 1997

Peak in photoluminescence is at same shell thickness as peak in elastic energy density CSCAMM, 4/24/2007

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- Strain model
  - Harmonic potential
  - Minimal stencil
  - Surface stress represented by variation in lattice constant
- Numerical methods
  - AMG
  - ABC
- Nanowires
  - Surface stress
  - No step bunching in 2D
  - Step bunching in 3D