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# Large-scale Atomic Effective Pseudopotential Method for the Electronic Structure of Semiconductor Nanostructures

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Large-scale Atomic Effective Pseudopotential  
Method for the Electronic Structure  
of Semiconductor Nanostructures

Applications

- Optoelectronic devices
- Quantum information

Nanostructures

- Quantum wells, wires & dots
- Nano, but ...  
 $10^3 - 10^5$  atoms

Capabilities of current  
atomistic methods (DFT):  
**1000 – 5000** atoms at most

# Large-scale **Atomic Effective Pseudopotential** Method for the Electronic Structure of Semiconductor Nanostructures

## Screened effective pseudopotentials

- No self-consistency cycle as in DFT
- **No total energy**

Empirical pseudopotential method

$$V(\mathbf{r}) = \sum_{\mathbf{G}} v(\mathbf{G})S(\mathbf{G})e^{i\mathbf{G}\mathbf{r}}$$

Semi-empirical pseudopotentials

DFT, different structures & lattice constants



Atomic  
Effective  
Pseudopotentials

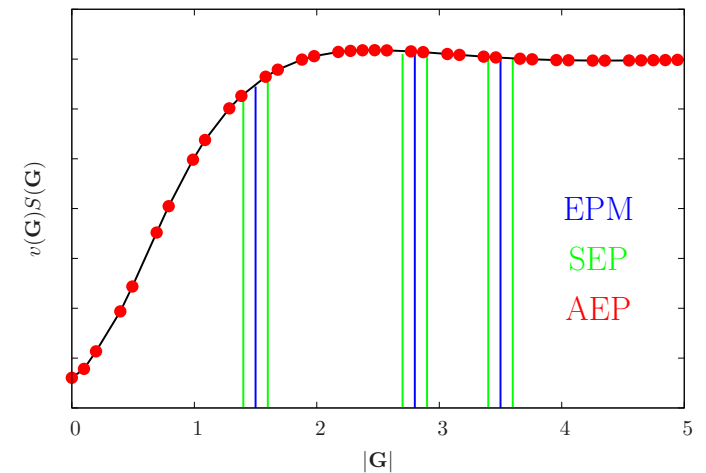
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## Large-scale calculations

- **Few selected eigenstates** of eigenvalue spectrum  
Electronic & optical properties  $\leftrightarrow$  States close to VB & CB
- Real space implementation  $\rightarrow \mathcal{O}(N)$

▶ Large-scale Atomic Effective Pseudopotential Program  
(LATEPP)

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▶ Atomic Effective Pseudopotentials

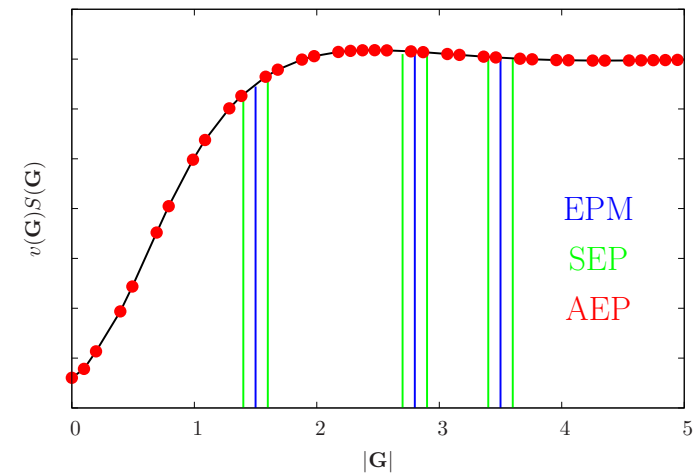
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# Atomic Effective Pseudopotential — AEP

## Basic idea

Screened local effective crystal potential from self-consistent DFT

$$V_{\text{eff}}^{\text{sc}} = V_{\text{local}}^{\text{Pseudo}} + V_{\text{Hartree}} + V_{\text{xc}}$$



Atomic Effective Pseudopotentials

$$V_{\text{eff}}^{\text{sc}}(\mathbf{r}) = \sum_{\alpha, n} v_{\alpha}(\mathbf{r} - \boldsymbol{\tau}_{\alpha, n}) \approx \sum_{\alpha, n} v_{\alpha}(|\mathbf{r} - \boldsymbol{\tau}_{\alpha, n}|)$$

## Formalism for extended supercells

Anion & cation potentials:  $v_{\pm} = v_a \pm v_c$

(1)

$$V_{\text{eff}}^{(1)}(\mathbf{r}) = \sum_{i=1}^{N/2} v_a(\mathbf{r}_i) + \sum_{j=1}^{N/2} v_c(\mathbf{r}_j)$$

(2)

$$V_{\text{eff}}^{(2)}(\mathbf{r}) = \sum_{j=1}^{N/2} v_a(\mathbf{r}_j) + \sum_{i=1}^{N/2} v_c(\mathbf{r}_i)$$

Analytic connection by Fourier transform

$$F[V_{\text{eff}}^{(\pm)}(\mathbf{r})] = \frac{1}{\Omega} \int_{\Omega} [V_{\text{eff}}^{(1)}(\mathbf{r}) \pm V_{\text{eff}}^{(2)}(\mathbf{r})] e^{-i\mathbf{G} \cdot \mathbf{r}} d\mathbf{r} = \beta_{\pm} v_{\pm}(\mathbf{G})$$

$$V_{\text{eff}}^{(1)}(\mathbf{r}) \pm V_{\text{eff}}^{(2)}(\mathbf{r}) = \sum_{n=1}^N (\pm 1)^{n+1} v_{\pm}(\mathbf{r}_n)$$

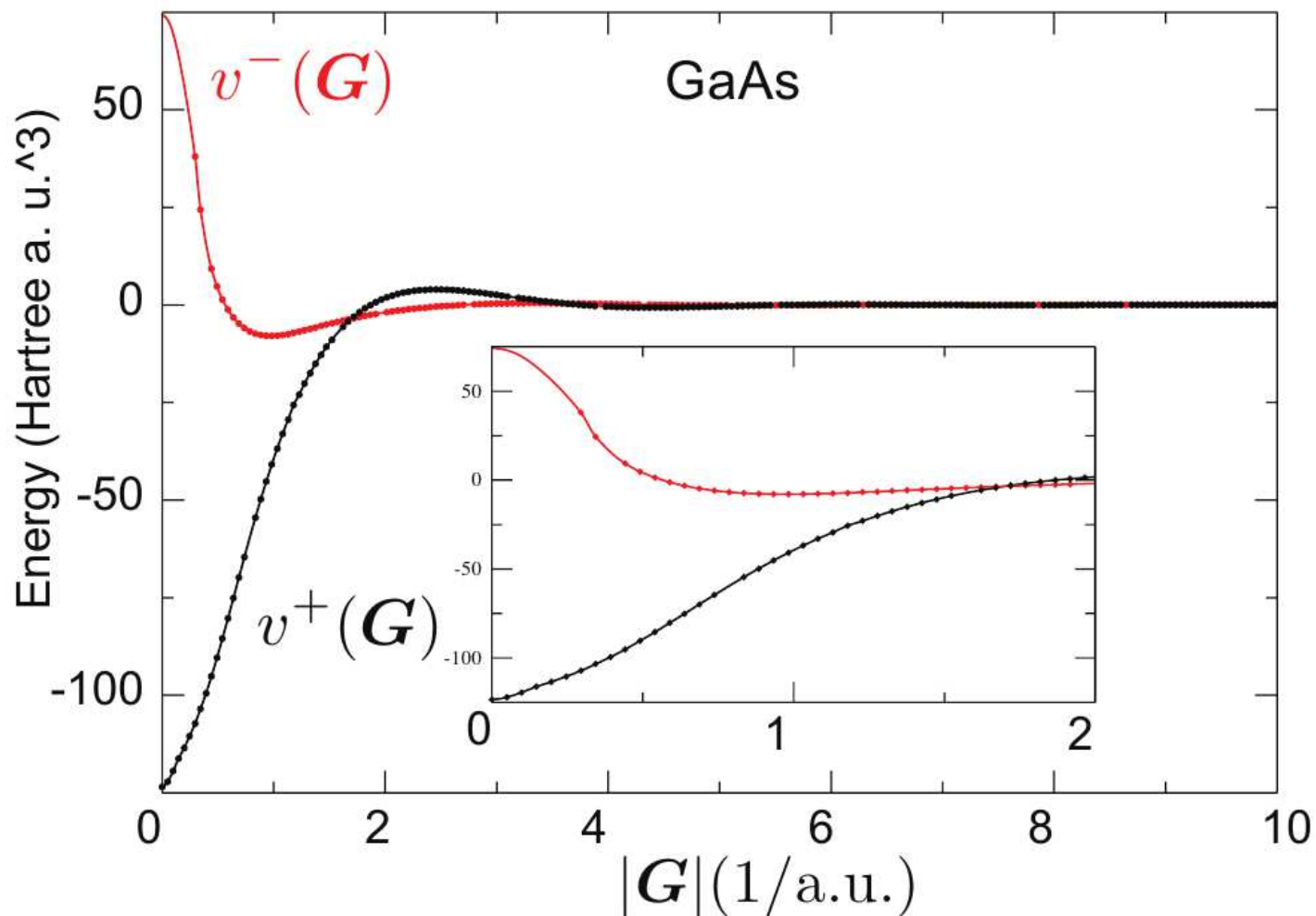
Spherical approximation

$$v_{\pm}(|\mathbf{G}|) = \Re[v_{\pm}(\mathbf{G})] = f(\beta_{\pm}, F[V_{\text{eff}}^{(\pm)}(\mathbf{r})])$$

J. R. Cárdenas and G. Bester  
Phys. Rev. B **86** 115332 (2012)

# Atomic Effective Pseudopotential — AEP

From elongated & slightly deformed supercells



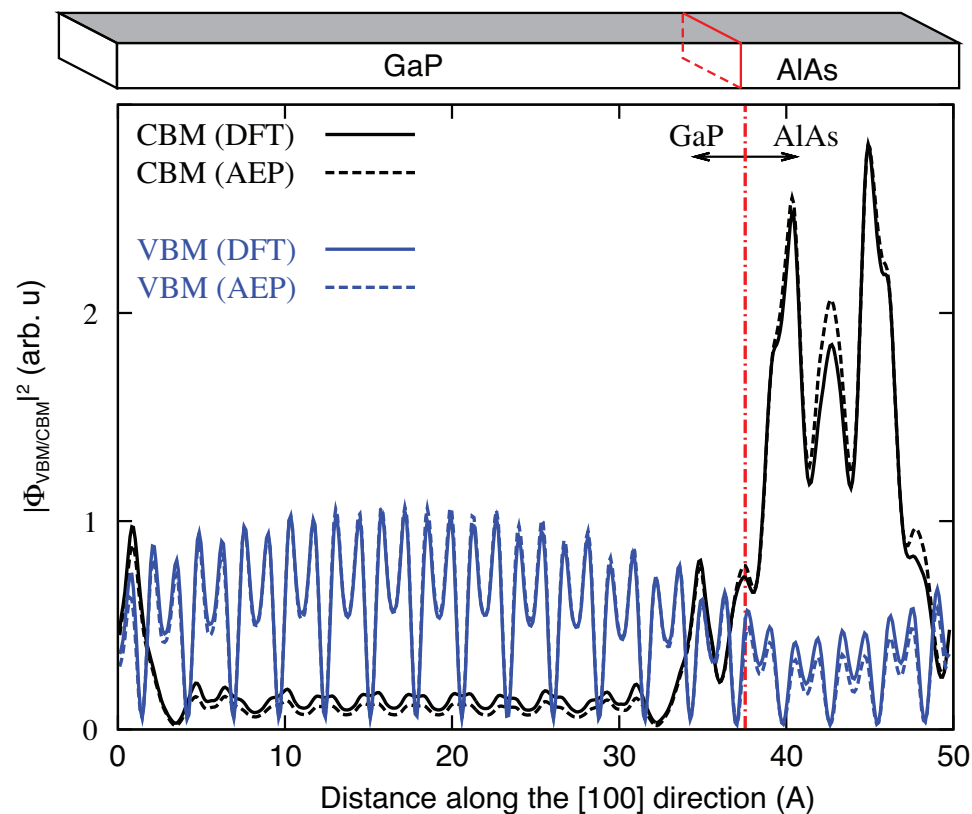
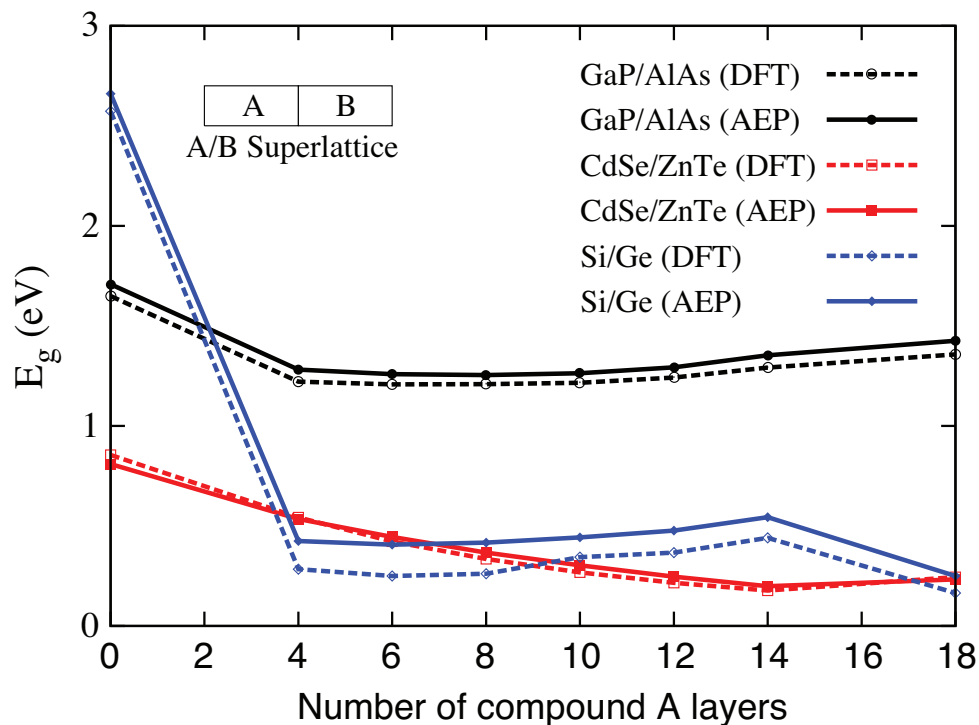
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$$v_{\pm}(|\mathbf{G}|) = \Re[v_{\pm}(\mathbf{G})] = f(\beta_{\pm}, F[V_{\text{eff}}^{(\pm)}(\mathbf{r})])$$

# Atomic Effective Pseudopotential — AEP

## Band gaps and wavefunctions of quantum well heterostructures



- Nice agreement with self-consistent DFT results
- Excellent transferability
- Errors
  - Spherical averaging  
→ Few meV differences
  - Charge transfer at the interface  
→ Small deviations in state densities



	DFT [eV]	AEP [eV]	$\Delta$ [meV]
CdSe	0.244	0.233	-11
CdSe (wz)	0.297	0.289	-8
CdSe (rs)	0.653	0.662	10

**Band gaps of different crystal structures**

# Large-Scale Atomic Effective Pseudopotential Program

$$\left( -\frac{\nabla^2}{2} + V_L + \widehat{V}_{NL} + \widehat{V}_{SO} \right) \Psi_{i,\mathbf{k}} = \epsilon_{i,\mathbf{k}} \Psi_{i,\mathbf{k}}$$

## Kinetic energy

- $T_{\mathbf{G},\mathbf{G}'}(\mathbf{k}) = \langle \mathbf{k} + \mathbf{G} | T | \mathbf{k} + \mathbf{G}' \rangle = \delta_{\mathbf{G},\mathbf{G}'} \frac{1}{2} (\mathbf{k} + \mathbf{G})^2$
- Real space: Finite difference scheme  $\rightarrow \mathcal{O}(N)$

## Local effective potential $V_L$

- $V^{\text{loc,eff}}(\mathbf{G}) = \frac{1}{\Omega_c} \sum_{\alpha}^{N_{\text{species}}} v_{\alpha}(\mathbf{G}) \sum_n^{N_{\alpha}} \exp(-i\mathbf{G}\boldsymbol{\tau}_{\alpha,n}) w_{\alpha,n}$
- Iterative solvers  $\xrightarrow{\text{FFT}} V^{\text{loc,eff}}(\mathbf{r}) \Psi_{i,\mathbf{k}}(\mathbf{r})$

## Nonlocal potential $\widehat{V}_{NL}$

- Fully separable formulation of Kleinman & Bylander

$$\sum_{l,m} |l,m\rangle \delta V_l(\mathbf{r}) \langle l,m| \xrightarrow{\text{KB}} \sum_{l,m} |\chi_{lm}^{\text{KB}}\rangle E_l^{\text{KB}} \langle \chi_{lm}^{\text{KB}}|$$

- Evaluation in real or reciprocal space

## Spin-Orbit potential $\widehat{V}_{SO}$

- Spin-Orbit coupling in norm-conserving pseudopotential

$$\widehat{V}_{SO} = \sum_{l,m} |l,m\rangle \left[ V_l^{\text{SO}}(\mathbf{r}) \mathbf{L} \cdot \mathbf{S} \right] \langle l,m|$$

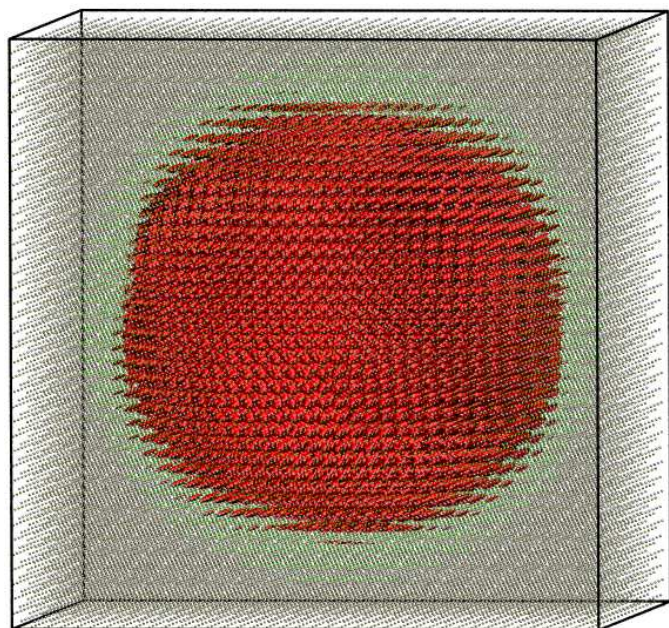
- Scalar relativistic part included in pseudopotential

## LATEPP

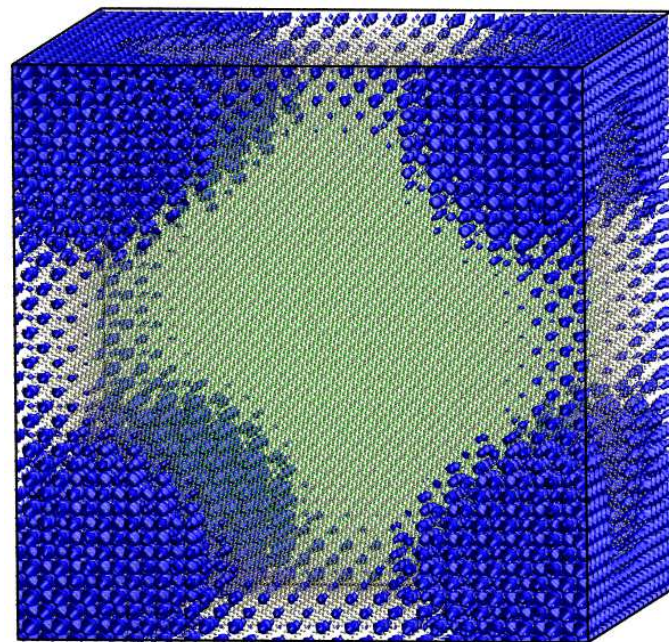
- $\Psi_{i,\mathbf{k}}$  represented & evaluated
  - in plane wave basis
  - on real space grid
- Reciprocal space  $\leftrightarrow$  Real space  
FFTW & FFTE
- Weights  $w_{\alpha,n}$   
for alloys & interfaces
- Atomic Effective Potentials  
No self-consistency required
- Compute few eigenstates
  - Iterative solver
  - Arnoldi restart method (ARPACK)
  - Eigenstates around  $E_{\text{ref}}$   
 $H\Psi \rightarrow H\Psi - E_{\text{ref}}\Psi$   
 $\Rightarrow$  Large-scale eigenproblems
- OpenMP parallelization



# Large-Scale Atomic Effective Pseudopotential Program



Highest occupied states

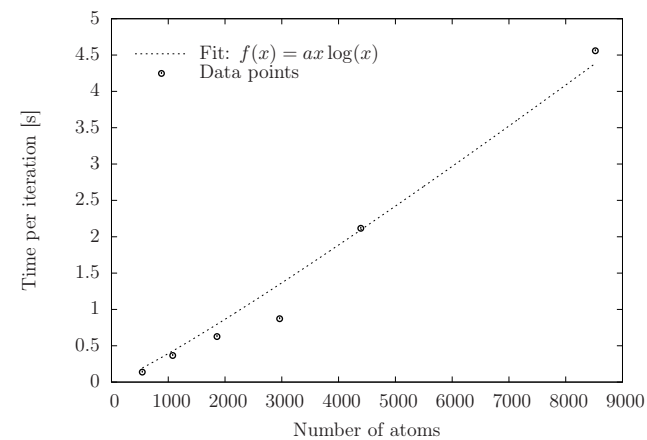
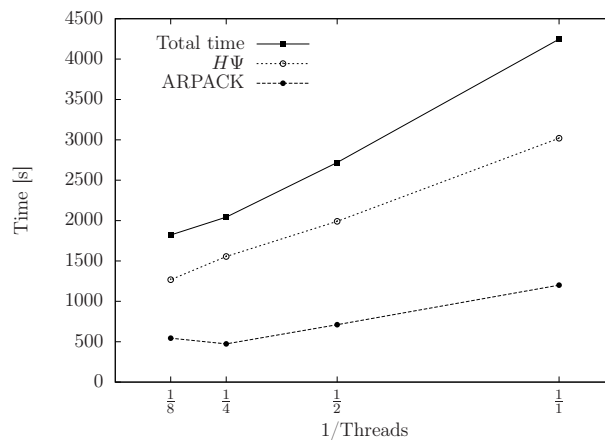


Lowest unoccupied states

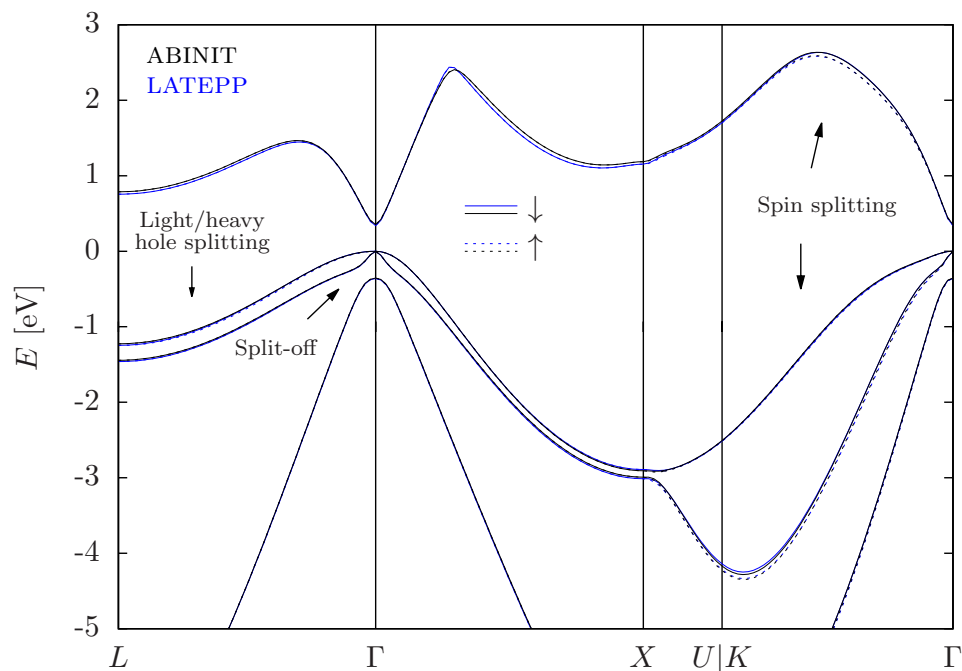
## GaAs quantum dot in AlAs

- GaAs QD radius: 5 nm
- AlAs host: 20 lattice constants
- Total: **64,000** atoms
- 7 million plane waves
- 20 states close to CB & VB
- Single node

## Performance of LATEPP

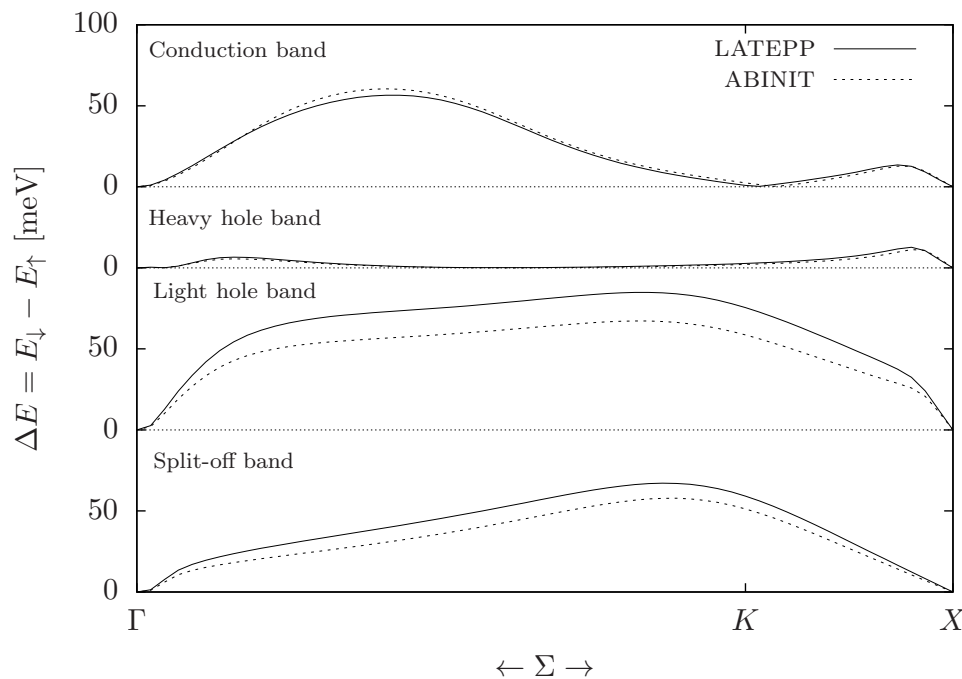


# Large-Scale Atomic Effective Pseudopotential Program



## GaAs band structure

- Perfect agreement to *ab initio* results
- Spin-orbit effects
  - Kleinman-Bylander / real space
  - Light/heavy hole splitting very accurately reproduced



## Spin splitting in GaAs

- Semi-local / real space
- Good qualitative agreement

# Summary — Conclusion — Outlook — Acknowledgements

## LATEPP

Atomic  
Effective  
Pseudopotentials

+

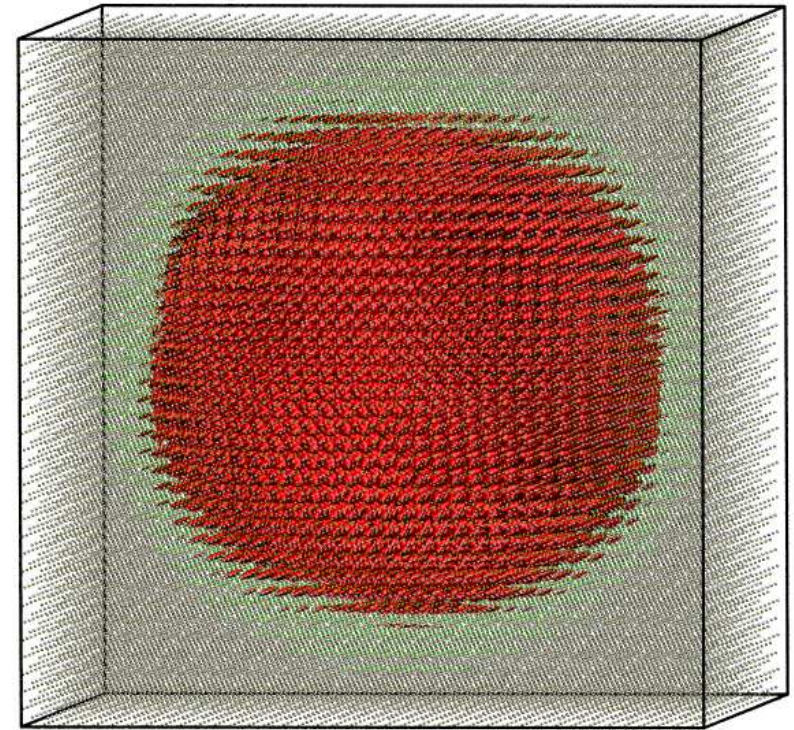
Efficient iterative solver  
Selected states of eigenspectrum

⇒ Large structures on *ab initio* level

## Outlook

- *ab initio* quality wavefunctions  
→ Configuration Interaction
- Beyond DFT: Correction of quasiparticle ...
  - effective masses: Nonlocal self-energy potential (empirical parametrization)
- band gaps: Energy dependence of  $\Sigma$  — shifting eigenvalues by scissors operator

$$E_g^{\text{LDA}} \longrightarrow E_g^{\text{LDA}} + \Delta$$



## Acknowledgements



Thank you for your attention!

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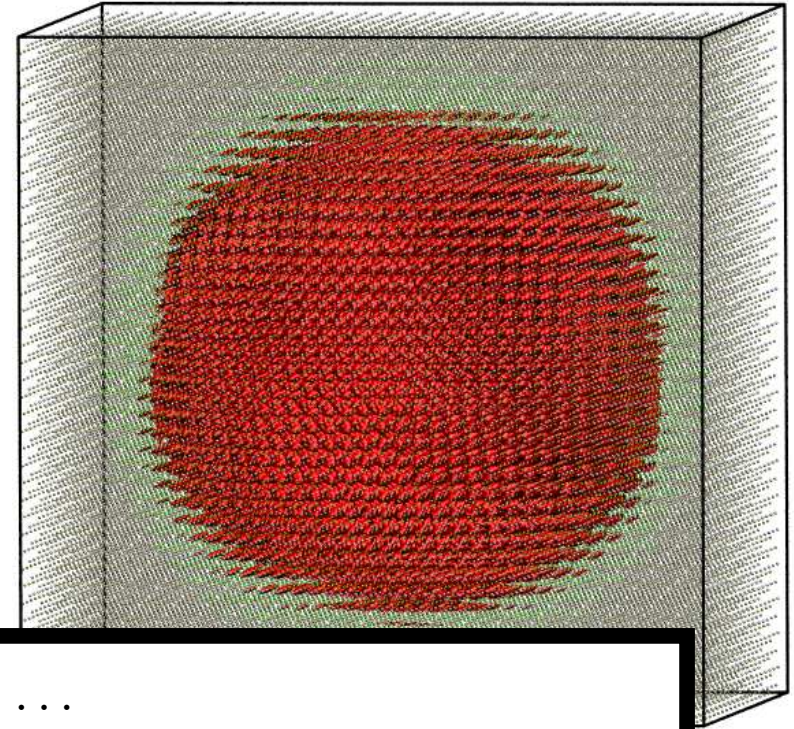
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## Outlook

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LATEPP & AEPs in use ...

- Mo, 1:39pm — Room: 325  
**Optical properties of  $\text{Ga}_{1-x}\text{Mn}_x$  from large scale *ab initio* calculations**  
JEROME JACKSON, RICARDO CÁRDENAS AND GABRIEL BESTER
- Th, 10:48am — Room: 307  
**Calculation of the optical properties of the nitrogen-vacancy center in diamond**  
DENIS ANTONOV, JÖRG WRACHTRUP AND GABRIEL BESTER

## Acknowledgements



Thank you for your attention!