## Electronic Structure Pseudopotential Calculations of Large (~1000 Atoms) Si Quantum Dots

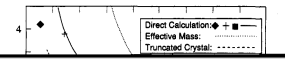
## Lin-Wang Wang and Alex Zunger\*

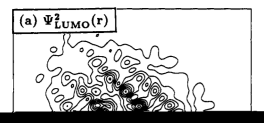
National Renewable Energy Laboratory, Golden, Colorado 80401 Received: September 30, 1993; In Final Form: December 15, 1993\*

	The electronic structure of quantum dots containing $N \ge 1000$ atoms is difficult to calculate by conventional molecular methods since the effort scales as $N^3$ . Our newly developed method allows calculation of eigenstates							
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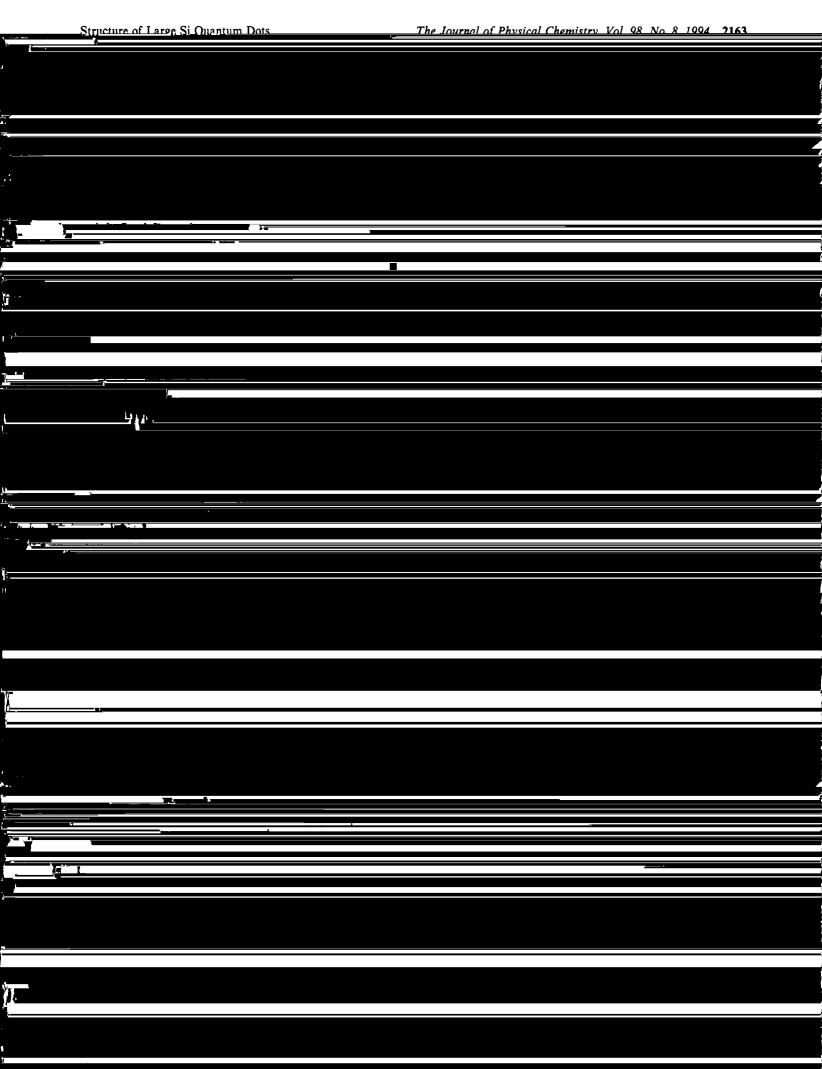
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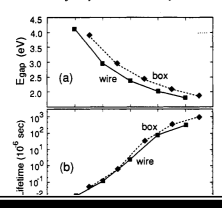
shape of the quantum dot does not change. The precise relaxation of the quantum dot surface atoms is taken from data on these three primary surfaces of H-covered Si films. The reconstructed





different treatments of the surface H potentials: In ref 38, the energy levels of the SiH4 molecule are fitted to give the matrix elements of the TB Hamiltonian. We have tested this procedure using EPM and find a similar surfacelike LUMO state. However, we feel that a SiH4 molecule is not an adequate model for H-covered Si surfaces. On the (001) film surface, there are two H atoms from neighboring H:Si:H groups which can be quite





lifetimes for these two systems are very close, within a factor of 2. The fact that the gap of a dots is larger than that of the wire is consistent with the larger quantum confinement for finite  $L_z$  (i.e., box). However, the nearly identical radiative lifetimes of these two systems is surprising given that in the wire  $k_z = 0$  for the HOMO and LUMO while in the box  $k_z \neq 0$  for both of them. This implies that, in this case, the radiative lifetime is mostly determined by the x, y directions, while the z direction has little effect.

## IV. Conclusions

We have used the empirical pseudopotential method to calculate

