Is there an elastic anomaly for a (OO1) monolayer of InAs embedded in GaAs? >Ua Yg'9"'6YfbUfX'UbX'5"YI 'Ni b[Yf'

CdhWu`WkUfUvWyfjnUhjcbcZ=b5gacbc`Unyfghfi Whifyg[fckbcbff/% Ł5 UbXf\$\$\$\frac{1}{2}; U5ggi VqhfUhyg

5dd`"D\mg"@Yhh"62ž%\$\$\$ f% - ' \!/%\$"%\$*' #%"%\$,) *%

Is there an elastic anomaly for a (001) monolayer of InAs embedded in GaAs?

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/December 100 Dec. 120 1000 110 12 12 10 04 17 1004

When a coherently grown (001)-oriented layer of InAs is embedded in a GaAs host, the coherency strain induces a perpendicular distortion of the embedded layer, predicted by continuum elasticity theory to be $\epsilon_{\perp} = 7.3\%$. Brandt, Ploog, Bierwolf, and Hohenstein, [Phys. Rev. Lett. 68, 1339 (1992)]

report on an investigation into whether a first-principles local-density total energy minimization shows such an elastic anomaly in the monolayer limit. We find that it does not.

When a layer of material with cubic lattice parameter a_0 is grown coherently in a host with cubic lattice parameter $a = \pm a_0$, the hyrigh layer undergoes a distortion of its lattice

from harmonicity. Regarding (2), it was previously shown⁴ that for certain interface orientations nonuniform atomic re-

[UU1], and the nonuniformity of these interplanar relaxations

near the interfaces is quite small. We will see below that

sponse to the distortion of its lattice parameter parametro the plane. Since surface effects are absent, the amount of the distortion can be accurately calculated by minimizing the bulk energy as a function of all the structural degrees of freedom. The simplest way to constitute this is to use harmonic continuum elasticity theory (see, e.g., Ref. 1). For example, for cubic materials with a layer orientation parallel to (001), the perpendicular strain $\epsilon_{\perp} = (c - a_0)/a_0$ is related to the parallel strain $\epsilon_{\parallel} = (a_0 - a_0)/a_0$ by

these do not substantially alter the applicability of the simple picture provided by continuum elasticity theory. Regarding (a), the question of interacts dependence has, that recently, been untested in semiconductor layers. One might imagine,

 $\epsilon_{\perp} = -2 \frac{C_{12}}{C_{11}} \epsilon_{\parallel}, \tag{1}$

small, interaction between them might substantially alter the

here the C greenesting anstante of the ambadded materi

theory.

For example, with the measured $C_{11}=0.8329$ and $C_{12}=0.4526$ (Tdyn/cm²) for InAs, Eq. (1) predicts $\epsilon_{\perp}=7.27\%$ for a layer of InAs buried in GaAs.

In view of the unprecedented result of Brandt et al.,³ we tested the validity of Eq. (1) via an atomistic, nonharmonic, first-principles theory of the relaxations of such a system to determine whether it would reveal the preakdown ostensibly demonstrated in the experiment. Our first-principles results in no way depend upon Eq. (1) or the approximations on which it is based. We found good agreement with Eq. (1) and no theoretical evidence for such a breakdown, in spite of using state of the cert techniques for relaxing the test leaves.

Recently, Brandt et al.³ examined the strain of buried (001) layers of InAs in GaAs experimentally, via a high-

Our tests improve on harmonic continuum elasticity theory is two store First me suffer the proach by a discrete atomistic approach, still limited to purely elastic energies. This is done with the Vesting valence-roice-near (VFF) model, which uses a description in terms of microscopic quantities the two-body bond

sponding to $\epsilon_L = 12.46\%$, much greater than the prediction of plied to a three monolayer film of InAs revealed a lattice distortion corresponding to $\epsilon = 6.06\%$ in good sometime edge. (1). Thus, they concluded that the widely used harmonic elasticity theory breaks down in the extreme limit of a

strain energy is minimized as a function of all the structural

Equation (1) involves several approximations: (1) the

ness of the embedded layer. Regarding (1), we can expect the

mation (LDA). With the total energy represented as a func-

tions to the total energy, not just the elastic contribution

strains; nowever, the lattice mismatch between GaAs and InAs (~7%) is not sufficient to cause a substantial departure

to it are treated on the same quantum-mechanical footing. This avoids all the potential pitfalls discussed above, and can be regarded as giving the best available theoretical estimate

In the first step, we applied the VFF model, with the bond lengths, to the calculation of the relaxed geometry of a structure containing an embedded (001)-oriented layer of InAs in a GaAs host, all confined to the GaAs lattice constant in the (001) plane. We found values ranging from ϵ_{\perp} =7.09% with 1 monolayer (ML) of InAs, to ϵ_{\perp} =7.17% with 10 ML.⁹ This demonstrates good agreement with the

large distortion in the ultrathin limit.

We note that in a recent paper Massies and Grandjean¹⁰ used a one-dimensional variant of the VFF model to investigate the behavior of monolayer-height InAs surface islands on GaAs, finding a relaxation away from the surface larger

ever, the clear agreement of our bulk three-dimensional VFF calculations with continuum elasticity theory demonstrates that this comparison of a one-dimensional model of surface relaxation with bulk strain is inappropriate.

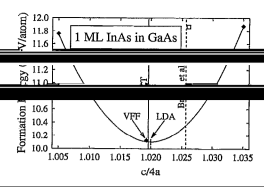
The inability of the VFF model to reproduce the microuse electronic elects as wen. Our total-energy calculations were performed using the LDA in the form of the

equivalent of 10 special points¹⁴ in the irreducible zinca 5%, far less man me more man 70% difference between me

stants of zinc-blende GaAs and InAs were determined by fitting to a Birch equation of state¹⁵ the total energies from seven calculations with differing lattice constants. The fit gives the values (Å) 5.60±0.005 (5.65) for GaAs and 6.02

constants induces only about a 4% (0.3 percentage points) overestimation in our calculated ϵ_1 .

For the first-principles calculations of the properties of



containing 1 monolayer of InAs vs its c/4a ratio. Filled diamonds denote direct first-principles LDA results; the curve is a cubic fit to those points. The minimum of the fit curve is marked by a vertical tick, labeled "LDA." The diamond symbol labeled "VFF" lies at the c/4a value predicted by the VFF model. The solid vertical line, labeled "CET," marks the prediction of

the InAs layer has $\epsilon_{\rm L}\!=\!12.5\%$, illustrating the large energy penalty incurred if the strain is concentrated in the InAs layer.

was used to find the equilibrium c/a_h ratio. While the symmetry of the structure permits additional relaxations beyond the relaxation of morely the LaAs hand leastly the effect of these was rough, where the structure small.

Our results are illustrated for the single-monolayer case

also shown. The aquilibrium a/a ratios for the three moth

lattice constants). We estimate the error in the LDA value ratio to be about 170. Similarly good agreement among the

recent pseudopotential calculation by Shiraishi and Yamaguchi. However, correspondence with one of the authors has clarified that their results do not, in fact, show evidence of an elastic anomaly in the monolayer limit.

so because the determination of $\epsilon_{\rm L}$ is based on the overall measured shift in the lattice. Because the VFF model provides a sufficiently accurate reproduction of the structural parameters obtained from the LDA calculations, we used it

to ohter the etructural effect of a thick GaAs substrate Tests

more than adequate to converge the structural reatures of the interface and the GaAs layer to match those with much thicker layers. We minimized the SL total energy with respect to all structural degrees of freedom, subject to the co-

to relay the atomic positions and a fit of total energy us clas-

respectively can raise a to 13.05% comparable to the

permitting more precise determination of the individual spacings, suggests that they should be able to detect excess In, at least in the two larger amounts we have suggested.²⁰

first subsequently grown GaAs laver would be to change the

²¹C. Giannini, L. Tapfer, S. Lagomarsino, J. C. Boulliard, A. Taccoen, B.

Capalle, M. R. O. Broadt and K. W. Bloom Phys. Phys. B 48, 11496 (1995).