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The models that have been discussed assume fluid molecules without any geometric structure. However, in some silicon oils the molecules have a clear chain-like structure which implies that excluded volume and chain entanglement need to be taken into account. These aspects, which are treated fully in [12], essentially involve details beyond the scope of this brief review of the status of microscopic treatments of droplet spreading.

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**Lattice-gas model**

The molecular dynamics analysis remains computationally very heavy despite ignoring the substrate structure. Moreover, the recent ideas about the spreading mechanisms put forward by Forcada and Mate need further investigation and clarification. For these reasons we returned to the Monte Carlo method used in an Ising or lattice-gas model that is richer from a statistical mechanical point of view [11]. The fluid particles were assumed to interact with their nearest neighbours in a cubic lattice and with the flat, unstructured substrate through the long-range potential \( V(z) = -A/z^3 \). The dynamics were assumed to obey Kawasaki spin-exchange dynamics, which conserves the amount of fluid matter in the simulation system. The model therefore allows for volatility, the amount of which can be changed by adjusting the Hamaker constant \( A \) and the temperature in the simulations.

As in the molecular dynamics simulations, we found dynamic layering and linear precursor growth at early times crossing over to diffusive spreading at late times. These features are shown in Fig. 3 giving snapshots of the development in time of the profile of an initially ridge-shaped droplet profile as well as the average densities of the droplet for early and late times.

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**Tracer Studies**

We have studied the fluxes at different times of tracer particles within the fluid phase and through the vapour phase via the evaporation-condensation mechanism. At early times, when growth is linear or nearly so, the flow of matter is concentrated in the fluid interface region of the droplet cap and the flux of particles is predominantly vertically downwards towards the substrate. This results in fast precursor growth and a purely horizontal flux in the precursor. For late times, when growth is diffusive, the fluxes are purely horizontal in the precursor and predominantly horizontal in the second and third layer. Increased vertical fluxes were only observed in the neighbourhood of the layer edges. Under these conditions, growth of the precursor is dominated by particles migrating off from the second layer edge and performing random walks on the precursor layer until they find holes to fill or rough precursor edges to stick with. It is believed that these mechanisms explain the experimental results of Forcada and Mate more satisfactorily than the evaporation-condensation process these authors invoked.

The models that have been discussed assume fluid molecules without any geometric structure. However, in some silicon oils the molecules have a clear chain-like structure which implies that excluded volume and chain entanglement need to be taken into account. These aspects, which are treated fully in [12], essentially involve details beyond the scope of this brief review of the status of microscopic treatments of droplet spreading.

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**A TRANSFERABLE TIGHT-BINDING POTENTIAL FOR TRANSITION METAL SILICIDES**

**The Intriguing Case of FeSi2**

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Semi-empirical methods for electronic structure calculations which combine flexibility and limited computational requirements allow for a reliable description of the link between structural features and the physical properties of technologically relevant complex materials.

A renaissance of the well-established tight-binding method for the semi-empirical calculation of electronic states in solids has taken place in the last few years owing to its ability to provide reliable interatomic potentials for total energy and molecular dynamics calculations in semiconductors and carbon structures [1]. The method has now been extended to cover transition-metal silicides, an important class of materials, both scientifically and technologically.

Metallic silicides play an increasingly important role in the fabrication of microelectronic devices owing to their excellent compatibility with technologies for manufacturing integrated circuits based on silicon. Refractory disilicides such as WSi2 and Ti2Si2 are used at present, but their poor crystallographic matching with silicon generates defect-rich interfaces. On the other hand, near-noble disilicides such as NiSi2 and CoSi2 with the simple fluorite (CaF2) structure display very sharp interfaces with silicon, to the extent that epitaxial CoSi2 has been experimented with for transistors of the permeable-basis type [2]. Attention of a different sort has recently been devoted to another member of the same family of compounds — namely FeSi2 — because it exists as a semiconductor (\( \beta \)-phase) at room temperature. This phase has a distorted fluorite structure with a large number of atoms in the (orthorhombic) unit cell. Being very complex, it can only be dealt with using semi-empirical methods and thus represents a meaningful example of the predictive power of tight-binding potentials.

The possibility of using \( \beta \)-FeSi2 for optoelectronic devices has been explored. However, this application is partially frustrated by light absorption measurements and first-principles calculations [3] showing that an indirect band gap as large as 0.78 eV is present, a value which is slightly smaller than the direct gap at the Brillouin zone centre (0.85 eV). Secondly, owing to the difference...
the crystal symmetry between the orthorhombic cell and the (111) surface of silicon, it is also difficult to obtain macroscopic epitaxial samples of FeSi2 with a good crystal quality. Nonetheless, interest in fundamental aspects of the semiconducting properties of β-FeSi2 remains so some effort has been devoted to investigating related phases which could improve understanding. In particular, the fluorite form of the γ-FeSi2 phase and the defective CsCl structure of Fe0.5Si (where the iron sites are randomly occupied at the 50% level) have been deposited successfully by Hans van Kaenel and co-workers [4] at the ETH in Zurich using molecular-beam epitaxy. These metallic materials do not exist in the bulk, but are stabilized by a very good matching with silicon substrate, so that epitaxial films can be grown up to tens of Ångströms in thickness. At larger thicknesses, depending on temperature, a transition to the stable β-phase occurs, together with the unexpected appearance of the tetragonal α-phase (see Fig. 1), the high-temperature bulk phase which has recently been deposited by various epitaxial techniques at temperatures well below its stability range in the bulk.

The intriguing phenomenon involving the epitaxial growth of a crystalline phase which does not appear in the bulk phase diagram is called pseudomorphism. It highlights the possibility of using epitaxy to explore structural phase transitions induced by changing either the pressure or the temperature [4]. Transition metal silicides are particularly interesting from this point of view, since their main metallic character allows for an easy transformation from one phase to another owing to the absence of high-energy barriers to atomic motion. A suitable annealing of the film, together with the substrate, even allows a transition from one stoichiometry to another (see Fig. 1). It must be appreciated, however, that the distinctive p-d bonding of silicides still preserves a fair amount of directional, covalent character. This probably explains why semiconducting gaps appear in the band structure depending on the atomic coordination, as is the case for β-FeSi2 and why a localized-orbitals scheme such as tight binding should offer an appropriate description.

**Semi-Empirical Schemes**

By considering the relationship between phase stability and film thickness, as described in Fig. 1, it is clear that the interface with the substrate plays a very important role in the epitaxial phase diagram of iron silicides. It is therefore very useful to have an accurate method for calculating the internal energy of structurally different and complex phases which takes account of the biaxial strain imposed by the silicon substrate in thin-film conditions. Unfortunately, since integration over the entire Brillouin zone of metallic systems demands considerable computing power, calculating the total energy is presently beyond the capabilities of first-principles techniques. These techniques are limited to estimating the electronic density of states (DOS) and the total energy of the most simple phases in the bulk. On the other hand, fully empirical interatomic potentials do not provide electronic information, and fitting several parameters defining the total energy is difficult because very few experimental data are available, even for the bulk properties of silicides.

What is needed is a semi-empirical scheme which can be fitted to the first-principles results in simple test cases and then used to extrapolate them to more realistic situations to provide electronic features, cohesion energies and elastic or vibrational information. This can be accomplished in the framework of the well-established tight-binding scheme for the calculation of the electronic states in the solids — an approach that has been applied successfully to FeSi2.

Since the late-1970s, tight-binding electronic bands have been satisfactorily included in total-energy calculations of the surface relaxation of polar semiconductors (s-p bonding) and transition metals (d-d bonding) [5]. This approach can be extended to bulk transition metal silicides (p-d bonding), where one also requires that the parameters be transferable from one phase to another in order to obtain a coherent representation of phase diagrams. The tight-binding approach is based on the partition of the total energy \( E \) of the solid into a band structure term \( E_{\text{bs}} \) given by the summation in reciprocal space of occupied tight-binding states \( E_{\text{bs}} \) and a repulsive contribution \( E_{\text{rep}} \) generated by a summation over the relevant neighbours \( j \) of a short-range, two-body potential \( f(r_{ij}) \). Thus we have

\[
E = E_{\text{bs}} + E_{\text{rep}} = \sum_{\kappa} E_{\text{bs,} \kappa} + \sum_{i<j} f(r_{ij})
\]

\( E_{\text{bs}} \) is negative and represents the covalent attraction provided by filling the valence electronic states which are generated by diagonalizing the Hamiltonian matrix in the localized orbitals representation. The latter includes s and p orbitals for silicon and s, p and d orbitals for the transition metal, with non-zero interactions up to the third-nearest neighbours (Si-Fe, Si-Si and Fe-Fe) for a precise description of the DOS near the Fermi level. One then obtains the dependence of the tight-binding Hamiltonian (and of \( E_{\text{bs}} \) in turn) on the atomic positions by scaling its parametrized elements with distance and angle following the universal rules formulated by J.C. Slater and W.A. Harrison.
$E_{\text{rep}}$, on the other hand, is positive and phenomenologically represents the quantum mechanical repulsion between occupied orbitals. It is obtained by fitting $f(r)$ to the equilibrium lattice parameters and bulk modulus of a few simple phases, using equilibrium and stability conditions. The parameters entering this two-body potential (and the parametrized Hamiltonian) remain limited in number, and in the case of iron sili-

des one obtains very satisfactory compari-

Some Interesting Results

Fig. 2 gives the calculated cohesion energy curves as a function of lattice parameter for the four FeSi$_2$ phases. The $\beta$-phase is correctly predicted to be at the lowest energy for a zero-temperature situation. The hierarchy in stability between the high-temp-

ature dependent forces are derived from the

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and then evaluating the time-frequency

spectrum would be very interesting.

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