# The Equilibrium Shapes of Crystals and of Cavities in Crystals

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Surface free energies are assumed to be the sum of the excess free energies of bonding of molecules in or near the surface, and the stable form of a crystal or cavity is assumed to be the form that makes the sum of these excess free energies a minimum. When only plane surfaces are allowed, this model predicts the same shapes for crystals as an equation of Wulff (2. Kristallogr. 34, 449 (1901)), which is based on the macroscopic thermodynamic relation of Gibbs ("The Collected Works, Vol. 1.: Thermodynamics," Longmans, Green, New York (1931)). The model predicts rounding of edges and corners of kinds which are not allowed by the Wulff relation and predicts that spherical forms of particles and cavities can be stable despite anisotropic surface free energies. The model provides a useful framework for analysis of whether unstable crystal or cavity shapes will evolve into stable or metastable forms. Some crystals and cavities that have been assumed to have equilibrium shapes instead have metastable shapes.

## I. Introduction

Gibbs (1), and, independently, Curie (2), derived as the condition which determines equilibrium shapes of crystals

$$\sum_{i} \sigma_{i} A_{i}$$
 is a minimum (1)

where  $\sigma_i$  is the specific surface free energy per unit area of crystal face i and  $A_i$  is its area. Wulff (3) first recognized that Eq. (1) implies that the facets of crystals or of cavities in crystals should obey the relation

$$\sigma_i/h_i = \sigma_i h_i = \dots$$
 (2)

where  $h_i$  is the distance from a common center drawn normal to crystal or cavity facet i.

Equation (1) describes the minimum work of surface formation derived from macroscopic thermodynamics and, there-

fore, (1) and (2) both appear almost selfevidently correct. But under conditions that favor equilibrium, cavities in metals are reported to be bounded by faceted, lowindex surfaces (4, 5). In contrast, exterior surfaces of high-purity metal crystals often show more rounded surfaces (6-8).

The difference is usually regarded as reflecting experimental error in one or the other kind of experiment. It has been suggested, for example, that the faceting of cavities may not be an equilibrium property (8). But annealing causes cavities of apparently constant volume formed by inert gas ion bombardment of metal foils to evolve from spherical to faceted shapes (4, 5). This evidence that these faceted cavities are more stable than rounded ones cannot be dismissed.

The fault for the apparent discrepancy between shapes of crystals and cavities

may lie partly in limitations in the analysis. Gibbs' proof was developed from macroscopic thermodynamics specifically for faceted crystals. It is shown in the next section of this paper that interpretations which have been given of rounded crystal surfaces in terms of Eq. (2) may not adequately explain particle properties. Then it is shown that if the free energy of a crystal surface is viewed as the sum of the excess free energies of those molecules of the crystal which are in or near the surface, the condition found for equilibrium is equivalent to Eq. (1) as long as rounding of surfaces is arbitrarily excluded. But the microthermodynamic model shows that rounded surfaces need not obey Eq. (2) and that spherical crystals or cavities may have wide differences in local specific surface free energies.

In a discussion section, the microthermodynamic model is used to suggest that metastable shapes are often likely to be produced and that some presumed equilibrium shapes of crystals and cavities probably are metastable forms. An accepted determination of the relative temperature dependences of surface tensions of different facets of cavity surfaces is shown to be questionable.

## II. Theory

A defect in the macroscopic model is that it provides no means of describing separately the thermodynamic behavior of molecules which are at corners or edges of surface planes. Edge and corner atoms are less tightly bonded than molecules in other parts of a surface. In consequence, a rounding of edges and corners that is not predicted by Eq. (2) might be expected (9). Herring (10) addressed this problem by assuming that, in the vicinity of an edge, the surface tension can be expressed as a series formed of constants multiplied by the radius of curvature to the  $0, -1, -2, \ldots$ 

powers. He then sought values for the constants and radius of curvature that are plausible and that would be consistent with Gibbs' fundamental relation, Eq. (1).

From his analysis Herring concluded that when the Wulff construction predicts faceted crystals, rounding of "at most a few tens of atom spacings" can occur, and that when the Wulff construction predicts that all or part of the equilibrium shape will be smoothly rounded, "for a specimen of observable size the amount of rounding will correspond fairly closely to that demanded by the Wulff construction without any further refinements."

Herring applied these deductions to interpret the observation that metal tips which have been used in field emission studies approach smoothly rounded shapes with perhaps a few flat regions in the crystallographically simplest directions. He concluded that either anisotropy of surface tensions must be low enough to give a Wulff construction of the smoothly rounded type, or that the rounded shape of the field emission tips is not an equilibrium one.

Careful efforts made subsequent to Herring's analysis to establish equilibrium conditions in tips of field emitters continue to yield essentially hemispherical forms (11, 12). Drechsler and Nicholas (13) have calculated theoretical equilibrium shapes using Morse or Mie potentials that are in excellent agreement with experimental observation when constants are used that yield surface energies of only slight directional anisotropy. Consequently, most investigators would probably now accept Herring's first suggested interpretation—that the tip shapes reflect an equilibrium consistent with Wulff's analysis.

Direct observations of diffusion in the adsorption layers of field emitter tips, however, show substantial dependence on surface orientation (14) and suggest, therefore, that adsorption energies vary substantially

with surface orientation; such variations imply that the bonding energies of atoms in the surface layer also vary substantially with orientation.

To develop a thermodynamic model that may be in better accord with molecular behavior in and on crystal surfaces, a single component or pseudo-single component crystal is here assumed to have a free energy of formation from its constituent molecules that is an additive function of the free energies of bonding of each molecule to other molecules, whether next or more distant neighbors. The crystal is in its most stable form when its constituent molecules are arranged in whatever way makes its total surface free energy a minimum. A crystal or cavity that is bounded only by plane surfaces  $i, j, \ldots$  is considered first.

The excess free energy of molecules in the outermost layer of the i surface over molecules in the bulk crystal is  $G_{ila}$ ,  $G_{ilb}$ , . . . and in the next outermost layer is  $G_{i2c}$ ,  $G_{i2d}$ , . . . , where a and b, for example, identify molecules of the layer with different excess free energies. The i surface is thus considered to include those subsurface layers in which, because of the presence of the surface, average molecular free energies are raised above the free energies of molecules in the bulk by some arbitrarily small fraction of kT, where k is the Boltzmann constant. Molecules at edges between crystal surfaces and molecules near edges have free energies  $G_{ij1e}$ ,  $G_{ij1f}$ ,  $G_{ij2g}$ , . . . , where the first two subscripts identify the surface planes to which the edge molecules belong, the number identifies the layer, and the last subscript identifies possible different excess free energies for molecules of the same edge and layer.

An approximation in Eq. (1) is that the free energy of formation of edges or corners is simply the sum of the free energies of formation of an equivalent area of surfaces of the planes bounded by the edges or corners. When this approximation is adopted,

 $G_{ij1e}$ , for example, is equal to  $G_{i1a} + G_{j1a}$ , that is, the free energy of edge and corner molecules can be assigned to the surfaces which the edge and corner molecules terminate. Then when  $n_i$  is defined as the sum of all molecules with excess free energies because of the presence of the surface i,

$$\sum_{i} n_i G_i = \sum_{i} \sigma_i A_i \tag{3}$$

and for crystals which are restricted to having only plane surfaces, the microscopic model is equivalent to the macroscopic model.

The restriction that only planar crystal faces are allowed can be dropped, and the most stable form of a crystal or cavity in a crystal is then that for which

$$\sum_{i} n_i G_i$$
 is a minimum (4)

where the summation is now understood to be over all molecules in which the free energy is raised over that of molecules in the bulk by the presence of exterior or cavity surfaces, whether planar or curved.

Equation (4) allows rounded corners and edges to be treated in the same framework as curved surfaces; within the overall limit set by Eq. (4) there is no restriction set on their radii of curvature, and a surface with essentially equal radii of curvature over two solid angles from a common center can have different surface free energies over those two solid angles if the resultant crystal form minimizes the free energy of the crystal as a whole.

Application of Eq. (4) can be illustrated by comparing theoretical relative stabilities at  $0^{\circ}$ K of a faceted face centered cubic (fcc) crystal to a spherical fcc crystal of the same volume. The expected facets are on (100) and (111) planes. For present purposes, the excess atomic enthalpy  $H_i$  for each atom can be assumed to be proportional to the difference between the number of neighbor atoms in the bulk crystal and the number of

neighbors  $N_i$  for atom i, that is,  $H_i = k(12 - N_i)$ , where k is the enthalpy per bond. This assumption makes the ratio of specific surface enthalpies in (100) surfaces to specific enthalpies in (111) surfaces,  $H_{100}n_{100}/H_{111}n_{111} = (4 \sin 60^\circ)/3 = 1.15$ , where  $n_{100} = n_{111} \sin 60^\circ$  because of the lower packing density in (100) planes.

With these relative surface enthalpies, the Wulff relation makes the ratio of the lengths of the normals from a common center to the 100 and 111 faces of the cubo-octahedron 1.15. Equation (4) asserts that a sphere of the same volume will have an equal total surface free energy when

$$\frac{\sigma_{\text{ave}}}{r} = \frac{\sigma_{111}}{h_{111}} = \frac{\sigma_{100}}{h_{100}} \tag{5}$$

where  $\sigma_{\text{ave}}$  is the average specific surface free energy per unit area over the sphere and r is its radius. For equal volumes,  $r = 0.985 \ h_{100}$ , and therefore a spherical crystal can be more stable than the Wulff cubo-octahedron as long as  $\sigma_{\text{ave}} < 0.985 \cdot 1.15 = 1.13$ . Local values of  $\sigma$  for the sphere can range from  $\sigma_{111} = 1.00$  to values larger than  $\sigma_{\text{ave}}$ . The Wulff relation permits spherical crystals only if  $\sigma$  values are the same over the entire surface.

Data of Van Hardeveld and Hartog (15) can be used to show that essentially spherical crystals with surfaces of varying local surface free energies can have low total free energies. As part of a statistical study of adsorption sites on metal crystals, they calculated the number of nearest neighbors for all the surface atoms of a number of polyhedral crystals and of a nearly spherical crystal, all of which could be formed from 683 atoms packed in an fcc structure. The nearly spherical crystal was formed in two steps. Atoms were assigned to all lattice sites lying within a sphere with a radius equal to 4.9 times the atomic diameter and with the center at a lattice site. A so-called rearranged sphere of a larger number of nearest neighbors per atom was then formed by movement of 8 atoms of low atomic coordination to vacant lattice sites with 5 neighboring occupied sites. The polyhedral shapes considered by Van Hardeveld and Hartog included several truncated octahedra; these had (111) and (100) surfaces, but the faces were not formed at the distances required by the Wulff construction. Evidently a symmetrical cubooctahedron cannot be formed of 683 atoms.

Crystals formed of 683 atoms are so small—if formed of gold atoms, only  $\sim 3$ nm in cross section—that a significant fraction of the atoms of the polyhedral crystals are in edge or corner sites; consequently it is not surprising to calculate that the unsymmetrical pair-bonded polyhedra have higher surface enthalpies than does the rearranged sphere. But the fact that the average number of neighbors per surface atom in the rearranged sphere is 8.1, making  $\sigma_{ave}$ = 1.11  $\sigma_{111}$  in this pair-bonding model for the small sphere, is pertinent. The optimum crystal packing must be size dependent, but this result suggests that spherical particles may be stable, despite markedly anisotropic bonding, for larger crystals as well.

#### III. Discussion

The microthermodynamic model that leads to Eq. (4) and the macrothermodynamic model that leads to Eq. (2) are both straightforward derivations based on the same initial assumption. The two approaches might have been expected to lead to identical results, but the microthermodynamic analysis suggests that particle equilibrium may be achieved by molecular packing arrangements that violate Eq. (2). In particular, Eq. (2) allows spherical crystals to be formed only if the specific surface free energies in each solid angle of the crystal are identical, while the microthermodynamic model allows greater surface free energy variations and allows corners and edges to take forms not allowed by Eq. (2). That crystal forms which do not obey the Wulff relation can be stable is demonstrable by model calculations of the kind illustrated by use of the Van Hardeveld and Hartog data. Experimental demonstration that such forms are stable is more difficult. The experimental evidence for substantial anisotropy in surface free energies which was presented above is qualitative. It would be desirable to study the surface anisotropy of field emitter tips by excitation methods that provide measurements of individual bond energies.

Equation (2) cannot be used to describe metastable crystal or cavity shapes. The microthermodynamic model provides a useful framework for understanding their formation and persistence. The persistence of metastable forms was recently subjected to a useful analysis in terms of nucleation theory (16). The present model has a different emphasis. Nucleation is usually considered to occur by statistical fluctuations in a local portion of a system. Shape changes for surfaces may be slow because the change from one shape to a more stable one requires the surface as a whole to assume intermediate shapes of higher total surface free energy than that of the initial shape.

The difference in stability between two different forms of a crystal—form b and form a—is given by  $G_{tb}-G_{ta}$ , where  $G_{tb}$  and  $G_{ta}$  are total surface free energies for the crystal or cavity. But provided the free energy change is negative, its magnitude plays no direct role in determining whether the transformation from a to more stable form b will take place. The driving forces for shape changes result from differences in free energies in subareas of the existing surface at any given time, say subareas  $\alpha$  and  $\beta$ .

Because shape changes are never directly driven by the free energy difference between initial and final forms, structure-sensitive kinetic factors can often play major roles in shape evolution. If, for exam-

ple, an initially spherical particle or cavity is unstable relative to a symmetrical polyhedral form, the sphere is almost certainly also unstable relative to a variety of other forms with the same facets, but with relative areas that violate both Eqs. (2) and (4). Statistical fluctuations in dislocation densities, which have been shown to be important in causing marked variations in rates of cavity migration (17), might cause different particles or cavities of a particular solid to evolve into forms which have the same surfaces, but with different relative areas. When a particle or cavity has reached a shape that is bounded by surfaces that approximate those of the most stable form. further evolution may become too slow to observe in experimentally practicable times.

Experimental observations should be evaluated with this expectation in mind. Perhaps, for example, although transfer of a monolayer from one surface to another would reduce the total free energy, transfer of one half of a monolayer would produce an intermediate form of higher total free energy. This kind of possibility could be tested by calculations of the kind made by Van Hardeveld and Hartog.

This analysis suggests that greater caution should be exercised in accepting persistent crystal forms as stable, or as consequences of impurities. For example, the observation (18) that initially spherical particles of gold, silver, and copper are transformed on annealing to partially rounded polyhedral shapes is definitive evidence that those latter shapes are more stable than the initial spheres. But, for gold particles annealed at 1000°C in dry H<sub>2</sub>, contamination by the furnace atmosphere is unlikely, and observations of imperfections in "nearly all" profiles may not indicate nonuniform contamination, as suggested by Sundquist (6), but rather the development of metastable shapes that cannot further evolve. The larger the crystal, the more unlikely it is that the form of maximum stability will be achieved—a conclusion also reached from nucleation theory (16).

The assumption (6, 18) that the relative specific surface free energies of (100) and (111) surfaces can be deduced from relative areas of facets in partially faceted crystals may not be warranted, even if one accepts Eq. (2), because the relative areas may simply be those that evolve by the kinetically most favorable process and then persist as metastable forms.

Development of rounded crystals from the partially rounded polygonal crystals when silver is heated through 775°C and copper is heated somewhat above 1000°C must mean that the entropy of transition to the essentially spherical crystals is positive, as noted by Sundquist (7). The positive entropy may not be a consequence of surface roughening as Sundquist suggested, however. It seems likely that the particles observed by Sundquist have atomically smooth surfaces like those of field emitter tips. If so, the positive entropy of rounding is not due to surface roughening of the kind considered in the model of Burton et al. (19).

Because polygonized cavities in magnesium, cadmium, and zinc evolve from initially spherical cavities (4, 5), the polygonal forms must be more stable than spherical ones. Observations that the ratio of cavity dimensions along the c axis to dimensions along the a axis of the hexagonal cavities show wide statistical fluctuations for zinc by a factor of two-constitute clear evidence that the cavities commonly approach metastable shapes. The assumption (20) that relative surface entropies can be calculated from the temperature dependence of the average axial ratios must be questioned; the average depends on unevaluated kinetic variations.

To many scientists the attraction of chemical thermodynamics lies in using our understanding of structure and bonding to predict thermodynamic behavior under conditions that have not been directly studied. The most important conclusion from this study is that an analysis of the structure and bonding in surfaces can provide new insights in surface thermodynamics. Papers are in preparation on the utilization of this approach to analysis of particle-vapor equilibria and of multilayer adsorption equilibria.

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#### References

- J. W. Gibbs, "The Collected Works, Vol. 1. Thermodynamics" p. 320, Longmans, Green, New York (1931).
- 2. P. Curie, Bull. Soc. Mineral. Fr. 8, 145 (1885).
- 3. G. WULFF, Z. Kristallogr. 34, 449 (1901).
- W. A. MILLER, G. J. C. CARPENTER, AND G. A. CHADWICK, *Philos. Mag.* 19, 305 (1969).
- H. O. K. KIRCHNER AND G. A. CHADWICK, Philos. Mag. 20, 405 (1969).
- 6. J. C. HEYRAUD, J. Cryst. Growth 50, 571 (1980).
- 7. B. E. SUNDQUIST, Acta Metall. 12, 585 (1964).
- 8. J. Friedel, Ann. Phys. 1, 257 (1976).
- C. HERRING AND M. H. NICHOLS, Rev. Mod. Phys. 21, 185 (1949).
- 10. C. HERRING, Phys. Rev. 82, 87 (1951).
- A. Muller and M. Drechsler, Surf. Sci. 13, 471 (1969).
- E. W. MULLER, in "Chemistry and Physics of Solid Surfaces, R. Vansalow and S. Y. Tong (Eds.), Vol. 1, p. 2, CRC Press, Cleveland, Ohio (1977).
- M. DRECHSLER AND J. F. NICHOLAS, J. Phys. Chem. Solids 28, 2609 (1967).
- 14. H. P. Bonzel, in "Chemistry and Physics of Solid Surfaces, R. Vanslow and S. Y. Tong, (Eds.), Vol. 1, p. 87, CRC Press, Cleveland, Ohio (1977).
- R. VAN HARDEVELD AND F. HARTOG, Surf. Sci. 15, 189 (1969).

- P. J. GOODHEW AND D. A. SMITH, Scr. Metall. 16, 69 (1982).
- P. J. Lemaire and H. K. Bowen, J. Amer. Ceram. Soc. 65, 49 (1982).
- 18. B. E. SUNDQUIST, Acta Metall. 12, 667 (1964).
- W. K. Burton, N. Cabrera, and F. C. Frank, Philos. Trans. R. Soc. (London) A243, 299 (1951).
- H. O. K. KIRCHNER AND G. A. CHADWICK, Philos. Mag. 22, 449 (1970).