

TOPOLOGICAL LOCALISATION OF DEFECTS AT ATOMIC SCALE

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ABSTRACT

The problem addressed in this paper is the detection of defects on atomic structures. The procedure proposed is in two steps. At first a tessellation is built starting from the atoms. It consists of a partition of the space into cells, and is used to define the neighbourhood relationships between the atoms. Then, the local contribution to a topological parameter, namely the Euler-Poincaré characteristic, is defined and measured for each cell. Within a regular tessellation, made of identical cells, this local contribution is equal to zero. Any local deviation from regularity corresponds to a tessellation containing cells with non-zero contributions. This allows us to locate the defects from a topological criterion and opens the way to a fully automatic detection of interfaces at atomic scale. The procedure is applied in 2D space for the detection of edge dislocations, grain boundaries and twins from HREM models and images. A 3D example is also given to illustrate its generality.

Keywords: defects, Euler-Poincaré characteristic, HREM images, interfaces, tessellations, topology.

INTRODUCTION

In materials science, it has long been recognised that topology offers a basic quantitative description of microstructures. Several examples of such a topological analysis can be found in the literature for sintering processes (Rhines, 1967; De Hoff *et al.*, 1972), grain growth (Smith, 1964; Rivier, 1986), multiphase grain structures (Cahn, 1966) or in relation with flow through porous media (Macdonald *et al.*, 1986; Jouannot *et al.*, 1995).

Topological descriptions have also been proposed for cellular structures: polycrystalline materials, biological cells, foams, etc (Smith, 1952; Aboav, 1980; Rivier, 1985). The relations are mainly expressed as a function of the mean number of sides per cell and based on Euler's equation and the local equilibrium of cells under surface tension. They are verified only when the rules governing this local equilibrium are fulfilled i.e. not on random tessellations of the space or on periodic structures (except for regular stackings of hexagons or tetrakaidecahedra).

We propose here a description of the spatial cell's organisation valid for any kind of cellular structure. It is based on the measurement of a new parameter: the contribution of each cell to the Euler-Poincaré characteristic of the space. This approach is general because no prerequisite (local equilibrium of cells,

geometrical unit, etc.) is needed. As far as grain boundaries between metal crystals are concerned, they have already been described as packings of polyhedral units (Ashby *et al.*, 1978). The present analysis complements, from a topological point of view, this geometrical description.

In this article, the contributions of the cells to the Euler-Poincaré characteristic of the space are used to locate the defects occurring on a regular tessellation. Then, on an atomic structure, the atoms belonging to a defect can be automatically selected using this topological criterion. To illustrate the method, edge dislocations or grain boundaries have been extracted from HREM images.

DEFINITIONS

Euler-Poincaré characteristic

In order to simplify the presentation, we limit ourselves to the 2D case. As will be seen in the last section, the extension to 3D space presents no major difficulties.

Consider a polygon; it is well known that it has the same number of vertices, V , and edges, E :

$$V = E. \quad (1)$$

This equation is nothing but Euler's equation:

$$V - E + P = 1, \quad (2)$$

where P , the number of faces, is equal to 1.

Consider now an aggregate of polygons. Their union satisfies the more general equation:

$$V - E + P = N, \quad (3)$$

where V , E and P stand respectively for the number of vertices, edges and faces of the aggregate (see Fig. 1). N is called the Euler-Poincaré characteristic (EPC) or connectivity number (Serra, 1982). It can be topologically interpreted as the number of connected components minus the number of their holes.

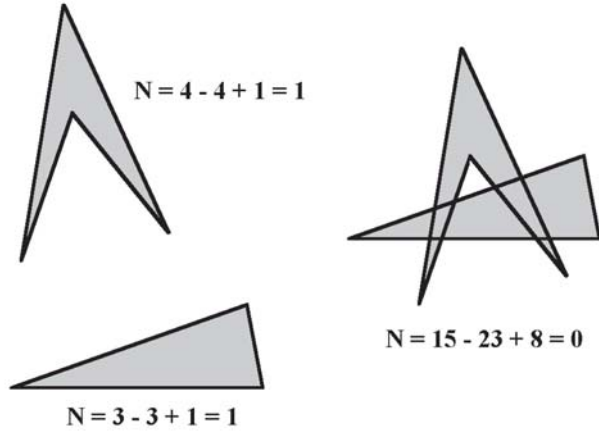


Fig. 1. Euler-Poincaré characteristic of isolated or aggregated polygons.

Planar tessellations

A planar tessellation is defined as a family of compact and convex subsets (Z_i) indexed by $i \in I$ and satisfying the following conditions:

- i) the union of all the Z_i is equal to \mathbb{R}^2 .
- ii) each interior of Z_i is non-empty.
- iii) the interiors of Z_i are pairwise disjoint.
- iv) the number of Z_i intersecting a bounded domain of \mathbb{R}^2 is finite.

The elements Z_i are the cells of the tessellation. Due to these conditions, any planar tessellation is made up of a countably infinite number of polygonal cells.

A facet F of the tessellation is defined as an intersection between cells. Its dimension, $\dim F$, is equal to 0 for a point, 1 for a segment and 2 for a polygon; its order, $\text{ord} F$, is the total number of cells to which it belongs.

Local contribution of a cell to the EPC

Let X be a compact subset of \mathbb{R}^2 which has been intersected by a tessellation ($Z_i, i \in I$). Under some

mild assumptions on the shape of X , it has been proved (Jernot *et al.*, 2001a) that the EPC of X can be written as:

$$N(X) = \sum_{F \in \mathcal{F}} (-1)^{2-\dim F} N(X \cap F), \quad (4)$$

where \mathcal{F} is the set of facets of the tessellation. The number 2 in the exponent is not fortuitous: it is simply the workspace dimension.

The local contribution of one cell to $N(X)$ is defined by sharing the term $N(X \cap F)$ uniformly between all the cells containing the facet F . Therefore the local contribution to the EPC of X with respect to the cell Z_i (Jernot *et al.*, 2001b) can be defined as:

$$C_i(X) = \sum_{F \in \mathcal{F}(Z_i)} \frac{(-1)^{2-\dim F}}{\text{ord} F} N(X \cap F), \quad (5)$$

where $\mathcal{F}(Z_i)$ is the family of all facets contained in Z_i .

The term ‘local contribution’ is justified by the fact that the EPC of X is the sum of all the contributions $C_i(X)$ associated with each cell Z_i :

$$N(X) = \sum_{i \in I} C_i(X). \quad (6)$$

In the case where $X = Z_i$, the Eq. 5 simplifies and gives:

$$C_i(Z_i) = \sum_{F \in \mathcal{F}(Z_i)} \frac{(-1)^{\dim F}}{\text{ord} F}, \quad (7)$$

which can be expanded as:

$$C_i(Z_i) = \sum_{F \in \mathcal{F}_0(Z_i)} \frac{1}{\text{ord} F} - \sum_{F \in \mathcal{F}_1(Z_i)} \frac{1}{\text{ord} F} + \sum_{F \in \mathcal{F}_2(Z_i)} \frac{1}{\text{ord} F}, \quad (8)$$

where $\mathcal{F}_d(Z_i)$ denotes the set of facets of dimension d contained in Z_i .

$\mathcal{F}_2(Z_i)$ has a unique element: the cell Z_i itself. Facets of dimension 1 and 0 of Z_i may not coincide with the edges and the vertices of the cell Z_i as can be seen in Fig. 2.

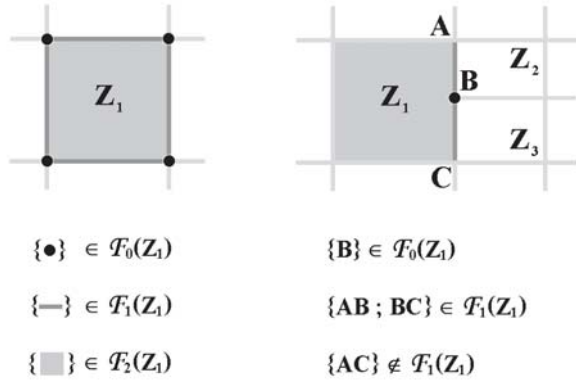


Fig. 2. The elements of $\mathcal{F}_2(Z_i)$ result from intersections between Z_i and its neighbouring cells. (AC) is an edge of the cell Z_i but is not a facet of the tessellation. B is not a vertex of the cell Z_i but is a facet of the tessellation.

In all examples given in this paper, the facets of dimension 1 and 0 coincide respectively with the edges and the vertices of the cells. Moreover, we always have $\text{ord}F = 1$ for facets of dimension 2 and $\text{ord}F = 2$ for facets of dimension 1. Then, the previous equation can be simplified as:

$$C_i(Z_i) = \sum_{V \in Z_i} \frac{1}{\text{ord}V} - \sum_{E \in Z_i} \frac{1}{2} + 1 \quad (9)$$

This equation is illustrated in Fig. 3. One can see at once that the local contributions obtained depend on the cells surrounding Z_i in the tessellation and bring a topological information that cannot be obtained from the mere number of edges of a cell. This point appears clearly in Fig. 8.

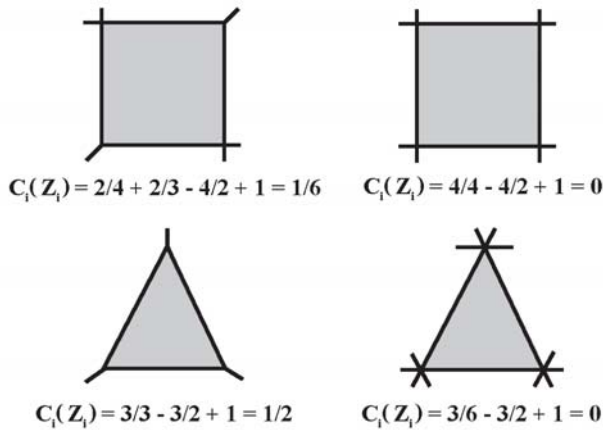


Fig. 3. Local contributions of one cell, Z_i , to the EPC of several tessellations of the 2D space.

For a regular tessellation, made of identical cells, the contribution of each cell to the EPC of the space

is equal to 0 (if not, the EPC of R^2 would be infinite instead of 1). This is not true if the tessellation is complex (e.g. Voronoï partition or Penrose tilings). In that case, only the mean value of the cell contributions is equal to 0. For cellular structures in equilibrium under surface tension (Smith, 1952; Rivier, 1985), assuming $\text{ord}V = 3$ in Eq. 9 leads to $C_i(Z_i) = 0$ if the number of vertices (or edges) of the cell is equal to 6.

If a deviation from regularity is encountered in some cells of a regular tessellation, then their contributions are not zero. This difference is now used for detecting defects on crystallographic structures.

DETECTION OF DEFECTS ON CRYSTALLOGRAPHIC STRUCTURES

The structures of interest are directly related to HREM images. First, the procedure is explained on theoretical structures. Then, model structures derived from HREM images are studied. Finally, an example is given for an HREM image. In all cases, the projected atomic columns (white or black spots according to the experimental conditions) are referred to as 'atoms'.

Illustration of the method on theoretical structures

Consider the images of Fig. 4a which represent partially incoherent and coherent interfaces according to Smith (1952). Two zones where atoms are regularly arranged are observed on each image. The purpose of this exercise is to detect the transition between both zones.

The first step is to build a tessellation starting from the set of atoms. It must be pointed out that several tessellations can be designed from the same set. In a standard one, we can consider the Voronoï cell associated with each atom. Others are based on graphs: the cells are limited by edges between adjacent atoms, an adjacency relationship defining the graph. If the atoms are represented by spots, the cells can be defined as the zone of influence of each spot (Lantuéjoul, 1978).

In the present case, we have used the Gabriel graph (Gabriel and Sokal, 1969): two atoms a_1 and a_2 are said to be adjacent if the disk with diameter (a_1a_2) does not contain any other atom. This tessellation has turned out to be more stable than the Voronoï partition.

The cells of Fig. 4b have been obtained using the Gabriel graph. Observe that only the cells located within the transition zone may have non-zero contributions. Then, the interface is localised from the set of atoms contained in cells with non-zero

contributions (Fig. 4c) but these cells depend on the tessellation design.

Edge dislocations

The analysis of the core structure of edge dislocations is usually carried out in three steps: i) observation of the sample along an appropriate zone axis ii) generation of atomic models using anisotropic elasticity, and iii) image simulation, followed by a

comparison between simulated and experimental images. An atomic model corresponding to the projection of an edge dislocation in GaN layers grown on (0001) sapphire and observed along the [0001] axis is shown in Fig. 5.

The local contribution of the cells is zero ($6/3 - 6/2 + 1$) except for the 5 and 7 sided ones whose values are respectively $1/6$ and $-1/6$. These cells correspond to the core of the dislocation.

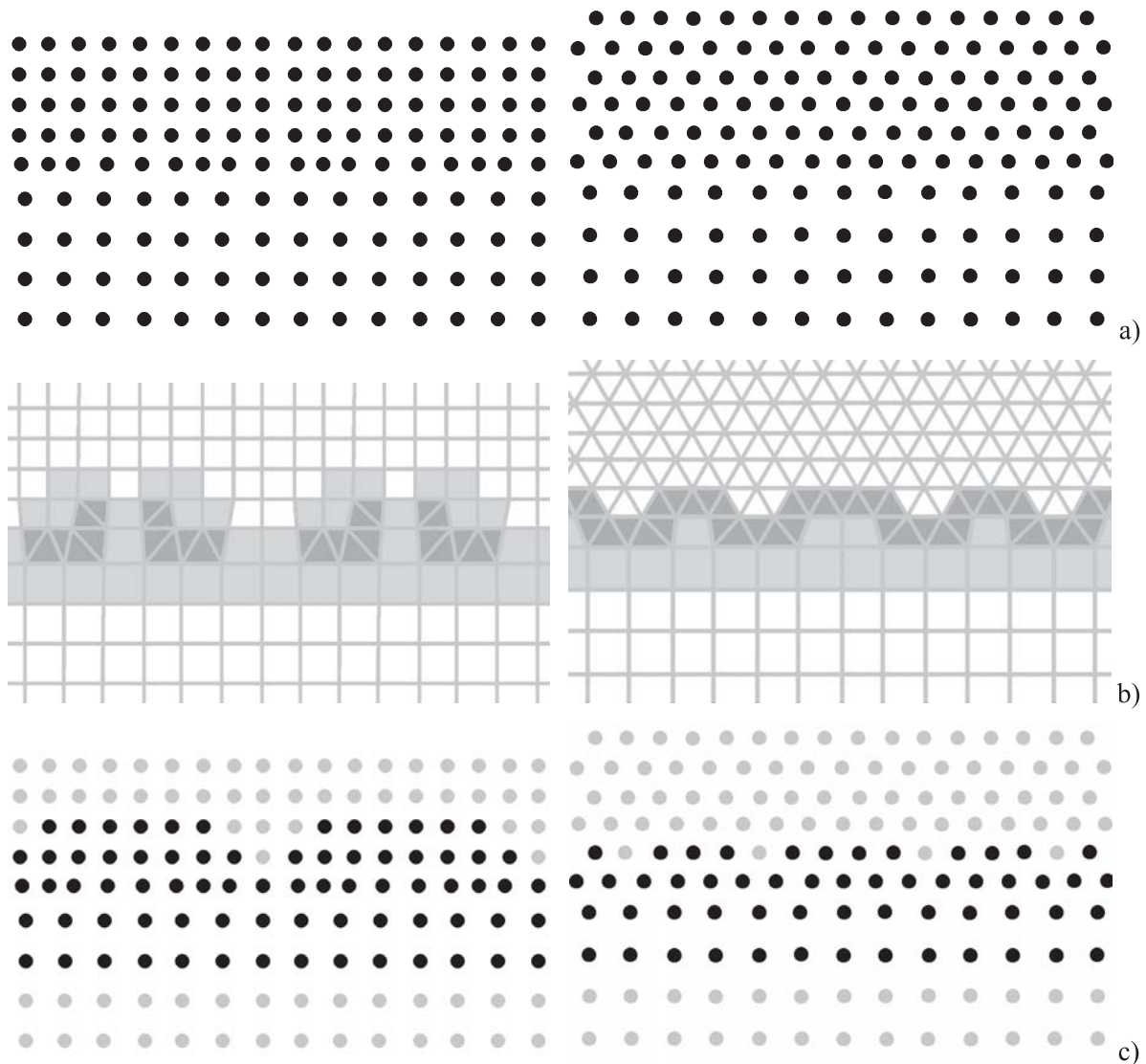


Fig. 4. Two schematic examples of interfaces between crystals. (a) atom locations, (b) tessellation associated with atom locations (Gabriel graph): the contributions of the white, light grey and dark grey cells are respectively zero, negative and positive, (c) the atoms that belong to cells with nonzero contributions are represented in black.

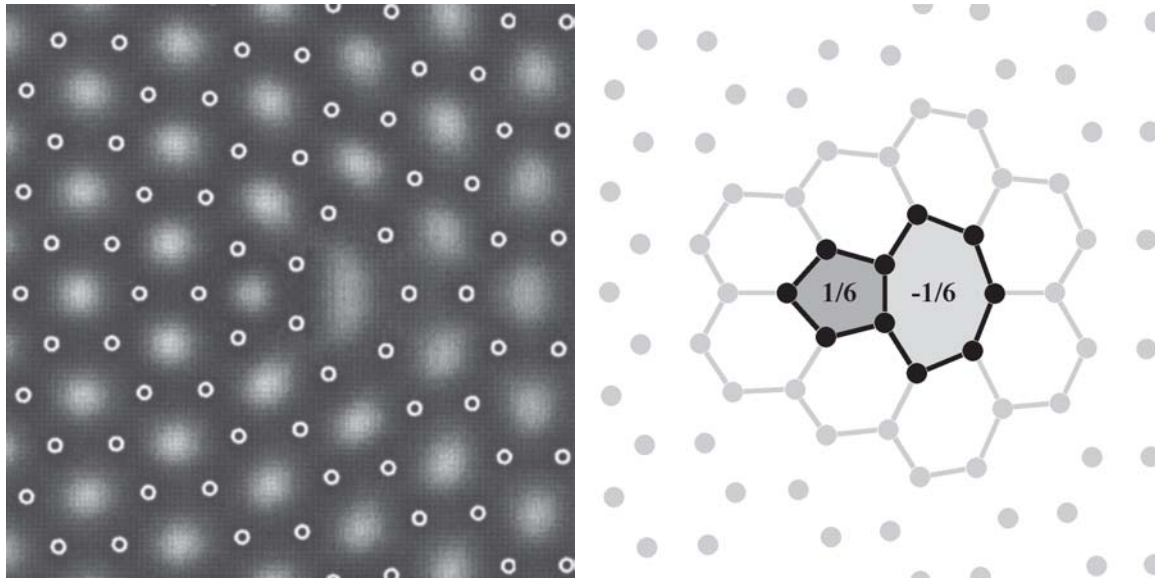


Fig. 5. Example of edge threading dislocation in GaN (adapted from Potin et al., 2000). For the dislocation core, the sum of the contributions of the cells to the EPC of the space is zero.

Grain boundaries

This procedure can also be applied to grain boundaries which are based on periodic arrays of dislocations. A first example is given in Fig. 6: for $\Sigma 5$, rotation of 36.87° around a $\langle 001 \rangle$ axis, the grain boundary built of one type of dislocation is perfectly planar. The set of cells with non-zero contributions is easily detected and defines the grain boundary plane.

In a second example (Fig. 7), the boundary plane

is slightly rough. It corresponds to $\Sigma 11$ which is described by a rotation of 50.48° around a $\langle 11 \rangle$ axis. Two variants A and B have been identified in silicon and germanium under different experimental conditions (Bourret and Bacmann, 1987; Putaux and Thibault-Desseaux, 1990). Again, a modification of the local contribution to the EPC allows us to accurately locate the grain boundary plane: the two configurations $\Sigma 11_A$ and $\Sigma 11_B$ are clearly differentiated and, as in the preceding example, the contribution of each period is zero.

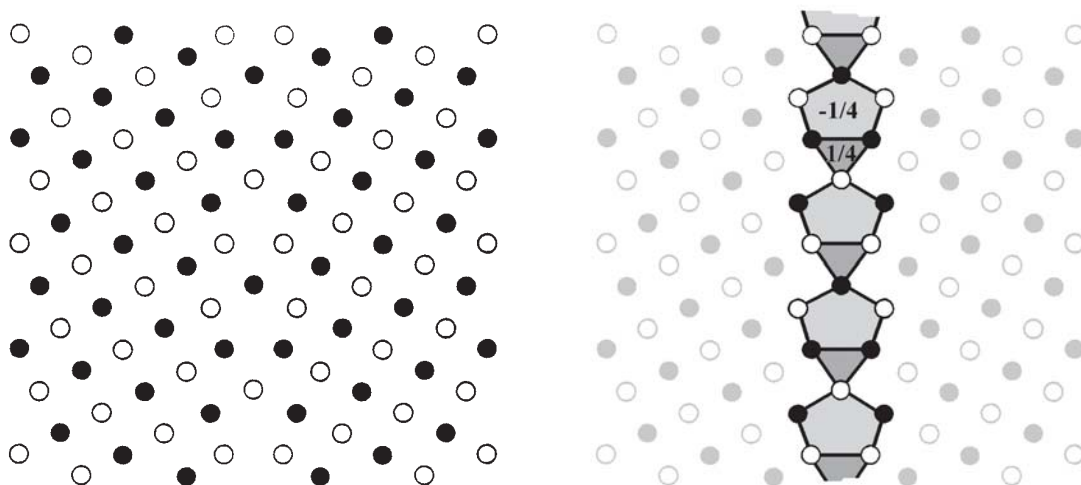


Fig. 6. $\Sigma 5$ tilt grain boundary in Cu (adapted from Grigoriadis et al., 1999). The disks and the circles correspond respectively to atoms at heights $z = 0$ and $z = 1/2$.

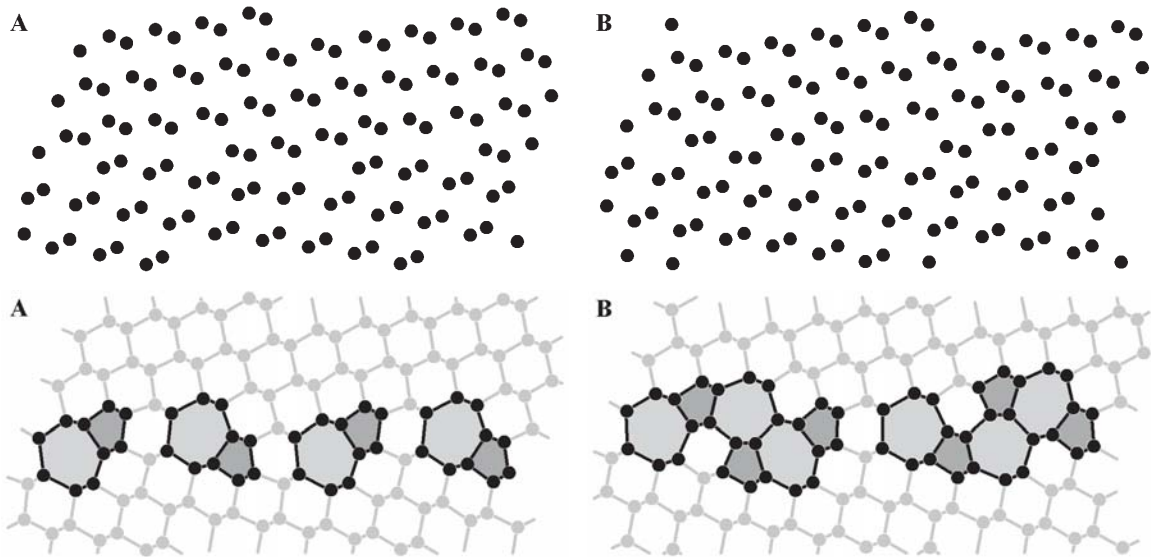


Fig. 7. Two examples of grain boundaries in silicon and germanium (adapted from Chen et al., 1999). $\{5 \text{ atom cell}\}$ and $\{7 \text{ atom cell}\}$ whose contributions are respectively $+1/6$ and $-1/6$ are grouped to form structural units with zero contribution.

Twins

The measurement can also be carried out in twins if a local modification of the cell contribution is observed. For instance, in the $\{111\}$ twin of the face-centred cubic system there is only a modification of

the orientation of the cells without any change in their topology, so this approach is not appropriate. For the $\{10\bar{1}2\}$ twin in the hexagonal system the situation is more favourable since the local topology is changed at the level of the twin plane (Fig. 8).

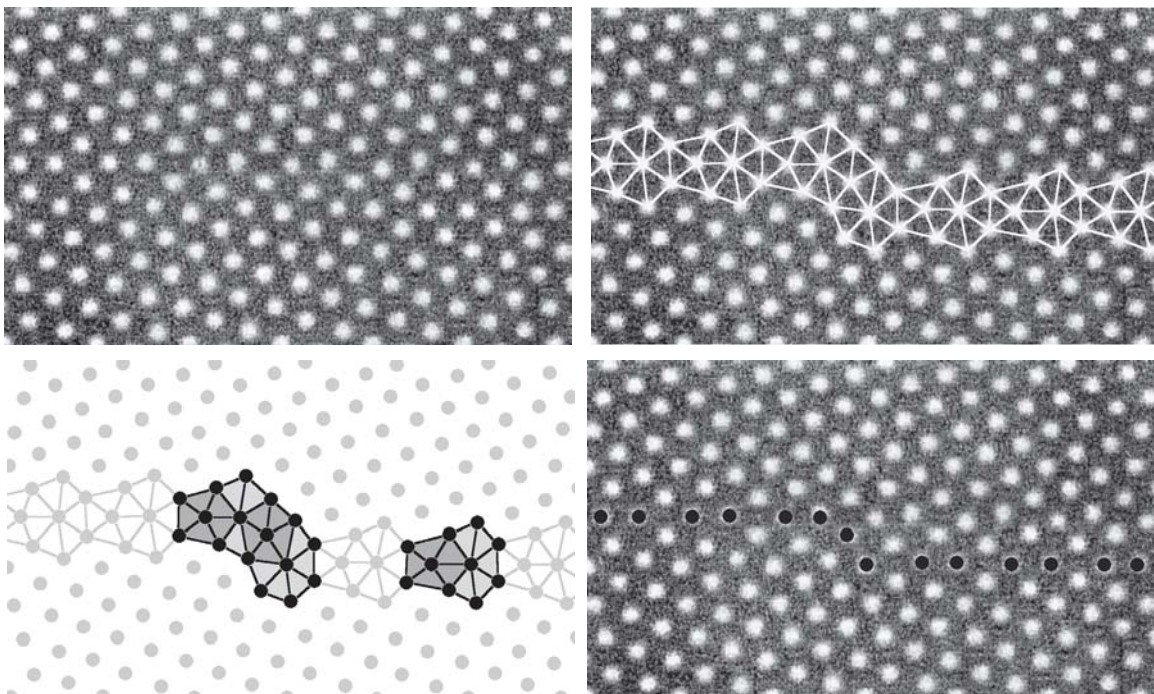


Fig. 8. Core defect inside a twin boundary from deformed α -Ti (adapted from Braisaz et al., 1996). On the bottom left figure, two different elementary features are clearly delineated: one for the grain boundary and one for the step.

Contrary to the preceding examples, the number of the edges of the cells is constant even in the defect zone. Nevertheless, the twin plane is still localised from the contributions of the elementary triangular cells of the tessellation. Their values are equal to zero everywhere $(3/6) - (3/2) + 1$ except in the twin plane for which a structural unit is detected. This structural unit is made up of 10 cells whose contributions are $-5/210$ (5 cells), $+2/210$ (2 cells) and $+7/210$ (3 cells). The total contribution of the structural unit is then zero. Inside the twin plane, a step corresponding to the total contribution $8(-5/210) + 6(2/210) + 4(7/210) = 0$ is observed. Then, even small atomic steps such as the step $b_{2/2}$ associated with the twinning dislocation (Braisaz *et al.*, 1996) are localised.

EXTENSION TO 3D SPACE

The procedure developed may be applied in 3D space provided that 3D images are available. In 3D space, the local contribution to the EPC of X with respect to the cell Z_i is defined as:

$$C_i(X) = \sum_{F \in \mathcal{F}(Z_i)} \frac{(-1)^{3-\dim F}}{\text{ord}F} N(X \cap F) \quad (5')$$

In the case where $X = Z_i$, this equation simplifies and gives:

$$C_i(Z_i) = - \sum_{V \in Z_i} \frac{1}{\text{ord}V} + \sum_{E \in Z_i} \frac{1}{\text{ord}E} - \sum_{P \in Z_i} \frac{1}{2} + 1 \quad (9')$$

since $\text{ord}F = 1$ for facets of dimension 3 and $\text{ord}F = 2$ for facets of dimension 2.

As in 2D space, a regular 3D stacking of identical cells leads to a contribution equal to zero for each cell. For instance, the contribution of each cell is $-(8/8) + (12/4) - (6/2) + 1$ in the case of a simple cubic stacking and $-(24/4) + (36/3) - (14/2) + 1$ for a compact stacking of tetrakaidecahedra. Any local topological modification of the tessellation will induce non-zero contributions in the vicinity of the defect.

A simple 3D interface can be given as an illustration: two crystals with simple cubic structures are translated $1/2, 1/2, 1/2$ one from the other, so as to create tetrahedra and square-based pyramids in the interface zone. Then, the non-zero local contributions correspond to:

$-(4/13) + (4/8) + (8/4 + 4/5) - (6/2) + 1 = -1/130$
for the cubes sharing a face with the pyramids,
 $-(5/13) + (4/4 + 4/5) - (5/2) + 1 = -11/130$ and

$-(4/13) + (4/4 + 2/5) - (4/2) + 1 = +12/130$ respectively for the pyramids and for the tetrahedra. The association of these three cells generates the elementary topological unit of the interface.

CONCLUSIONS

In this article, a method has been proposed for detecting the interfaces of an atomic structure provided that the associated tessellation departs from a regular one. It is based on a new topological criterion and has been illustrated on several examples with a particular reference to HREM images.

Associated with an automatic extraction of atomic positions (Chaix, 1998), this opens the way to a fully automatic detection of interfaces at atomic scale but requires images with a good contrast and a sufficient magnification.

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