Hard-sphere displacive model of extension twinning in magnesium.

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Abstract
A crystallographic displacive model is proposed for the extension twins in magnesium. The atomic displacements are established, and the homogeneous lattice distortion is analytically expressed as a continuous angular-distortive matrix that becomes a shear when the distortion is complete. The calculations prove that a volume change of 3% occurs for the intermediate states. The twinning plane, even if untitled and restored when the distortion is complete, is not fully invariant during the transient states. The crystallographic calculations also show that the (90°, a) twins observed in nano-pillars and the (86°, a) twins observed in bulk samples differ only by a slight obliquity angle (± 3.4°). Continuous features in the pole figures between the low-misoriented (86°, a) twin variants are expected; they are confirmed by EBSD maps acquired on a single crystal of magnesium. As the continuous mechanism of extension twinning is not a simple shear, a “virtual work” criterion using the value of the intermediate distortion matrix at the maximum volume change is proposed in place of the usual Schmid’s law. It allows predicting the formation of extension twins for crystal orientations associated with negative Schmid factors.

Keywords: Extension twins; hexagonal close-packed; magnesium; Schmid factor; angular-distortive matrix; hard sphere.

1. Introduction

1.1. The choice of paradigm
It seems important to make clear the paradigm used in this work. Indeed, the manuscript in its first formulation, i.e. without the sections 1.1, 5, 6.2 and 6.3 [1], was rejected twice from other journals mainly because it does not imply “disconnections” and because it is an “oversimplification”. We hope that this preamble will help the reader to understand why we think that the mechanism of deformation twinning is not compatible with the “disconnection” theory, and why the “displacive” paradigm has been chosen. We admit that the electronic structure of the crystal is ignored in first approximation, but a simple model is not a false model; it is just a first step toward a more accurate model.

Deformation twins appear in the face centered cubic (fcc) and body centered cubic (bcc) metals deformed at high speeds or low temperatures. They are more commonly formed in hexagonal close-packed (hcp) metals because of the lowest number of slip systems. Deformation twinning share many characteristics with martensitic transformations; they are formed at very high speed (close to the speed of sound) and they take the shape of lenticular plates that become highly intricate at high deformation rates: both deformation twinning and martensitic transformations belong to the wide class of displacive transformations [2]. Deformation twinning has been mathematically treated as a homogeneous shear for more than one century [1]-[9]. Bevis and Crocker [7] adopted the definition “twinning shear is any shear which restores a lattice in a new orientation” and used it to build a generalized theory that predicts the possible twinning matrices. The more realistic ones were chosen among those with the minimum shearing magnitude. The use of shear matrices relies on the observations of planar interfaces between the parent and its twins. As explained by Christian and Mahajian [2]: “Since a parent crystal and its twin remain in contact at the interface plane during the formation of the twin, the relation between the structures must be such that this plane is invariant in any
deformation carrying one lattice into the other”. Consequently, the classical displacive “shear” theory of twinning makes a large use of shear planes and shear directions, or their conjugates. Concerning more particularly the hcp metals, fifteen deformation twinning modes could be established in titanium by Crocker and Bevis [8], and a more recent theoretical study also based on lattice correspondence resulting from shear deformation has been proposed by Niewczas to analyze the correspondence of the dominant slip modes in the parent and its twins [9]. Another point should be mentioned. The Bravais cell of a hcp crystal is composed of two atoms, roughly speaking the 8 atoms at the “corners” of the hexagonal lattice (counting for 1/8) and one atom in the position (2/3, 1/3, 1/2) “inside” the hexagonal cell. The atoms at the corners have a trajectory directly given by the homogeneous linear shear transformation, while the trajectory of the atom inside the cell is given by an affine transformation, i.e. the same linear shear distortion but associated with a translation component called “shuffle”. The homogeneous lattice distortion is in full agreement with the fact that the atoms can move collectively in a coordinated way at very high velocities.

Although the “displacive shear” theory is established for more than a century, another approach was proposed twenty years ago by Pond, Hirth, Serra, and Bacon [10][11][12]. They introduced the concept of “disconnection”, derived from the “twinning dislocations” introduced by Orowan in 1954 [13], the “emissary dislocations” imagined by Sleenwyk in 1963 [14], and the “zonal dislocations” introduced by Mendelson in 1969 [15]. Their theory, also named “topological model”, supposes that the lattice is inhomogeneously transformed by the movement of disconnections. A disconnection is a “customized” defect; it is a mixture of a classical dislocation with its usual Burgers vectors and a “step height” introduced in order to accommodate the geometrical misfit at the interface. The disconnections are determined with the help of the dichromatic pattern i) by overlapping the parent and twinned lattices according to their orientation relationship (OR), and ii) by finding the smallest displacements vectors between the nodes of the two lattices. Disconnections are a substitute to the displacements resulting from the homogenous lattice distortion in the classical displacive theory. The disconnection theory seems to rally an active community despite of its major weaknesses:

a) The concept of “shuffle” has strongly digressed from the one used in the classical shear theory. In the disconnection theory, the shuffles are associated with the motion of the step part of the disconnection, without clear justification of this choice and no apparent link with the initial meaning of “shuffle” because the notion of homogeneous lattice distortion was completely suppressed.

b) The disconnection theory can’t explain where the dislocations/disconnections come from and how they can be produced in a very precise sequential and coordinate order. Serra and Bacon [11] considered (as we do) that the “pole mechanism” is not realistic and proposed another model where “a matrix dislocation becomes a new source of twinning dislocations”, but after twenty years the existence of these sources remain to be proved experimentally. In addition, nothing is said about their driving forces.

c) The disconnection theory is unable to account for the elastic phase transformations and for the second order phase transitions because in these cases dislocations are not required to accommodate the parent/product interfaces.

d) It is mathematically possible to associate an inhomogeneous distribution of disconnections to any homogeneous displacement field; however, the notions of lattice distortion and correspondence that specifies how the crystallographic directions are transformed are then lost. By assuming that the shear in inhomogeneous, it is nearly impossible to figure out how the atoms move. For example, the Bain correspondence of fcc-bcc martensitic transformation can’t exist within the disconnection theory.

e) Any theory should be judged by its predictions. The results of the disconnection theory are often compared with High Resolution Transmission Electron Microscopy (HRTEM) images or molecular dynamic simulations, but the macroscopic features of the transformations (habit planes, variant pairing, accumulated strains etc.) are clearly beyond its scope.

f) Because dislocations and plasticity are the results of irreversible mechanisms, the “disconnection” theory is unable to explain the detwinning mechanisms observed in fcc [16] and hcp metals [17][18] or the reversibility of the transformations in shape memory alloys.

g) The disconnection theory is unable to explain how deformation twins can grow at velocities close to the speed of sound. How is it possible that coordinate arrays of dislocations can move so fast? In the same order of ideas, why decreasing the temperature favors deformation twins if twinning is generated by disconnections?
By ignoring the basic principles of the classical displacive theory, the disconnection theory becomes unable to respond to these fundamental questions. To our opinion, dislocations are not the cause of the twinning distortion but its consequence; they are the defects let at the interface, and also far away from the interface in the surrounding matrix; they are created to locally accommodate the deformation caused by the lattice distortion. Moreover, we think that the concept of “moving interface” gives a false image of the mechanism. Martensite and deformation twinning is suddenly formed in a cooperative (military) way, and all the atoms move similarly relatively to their neighbors: the interface does not move, it is the product phase that is transformed by bursts inside the surrounding matrix. The classical displacive theory uses homogeneous distortion (linear algebra) that is well adapted to describe the formation of twins or martensite that grows at high velocities as a “wave” [19][20]. For all these reasons, we have chosen to treat deformation twinning within the displacive paradigm, i.e. by using distortion matrices, correspondence matrices etc. This is the approach we have already followed in our recent developments of the crystallography of martensitic transformation in steels [21], and more generally between the fcc, bcc and hcp phases [22]. The case of mechanical twinning in fcc crystals was also treated in Ref.[22]. The main difference between our model and the classical “shear” theory is that we believe that the shear matrices are inappropriate to catch the details of the distortion mechanism because the atoms would interpenetrate too much during a continuous simple shear. Therefore, we have replaced the shear matrices by the more general “angular distorting” matrices [22] that can be explicitly calculated by assuming that the atoms are hard-spheres of constant size. The displacive paradigm does not mean that the concept of disconnection is useless but, to our opinion, disconnections and disclinations should be introduced in a second stage, as consequences of the lattice distortion. The disconnections are the dislocation arrays that accommodate the translation parent/twin misfits at the interface, and the disclinations are the dislocation arrays that accommodate the rotational misfits in the parent matrix around the twin.

1.2. Current issues about extension twinning in magnesium

Magnesium is a very good example of hcp metal in which deformation twinning play a major role on the mechanical properties. Magnesium, thanks to its lightness, is used in some automotive parts and is considered as a good candidate for many other applications; however, it suffers from a poor ductility because its number of slip systems is low. Twinning comes as an additional deformation mechanism that improves the elongation, but in a very anisotropic way. A better understanding of deformation twinning could thus help to improve the mechanical properties and enlarge the use of magnesium alloys in industry. The main twinning mode observed in magnesium is “extension” twinning on the {1012} planes. These twins are formed when the c-axis is close to the tensile axis. Other twinning modes can also be observed, such as the “compression” twins on the {1011} planes that are created when the c-axis crystal is close to the compression axis, and the so-called {1011}-{1012} “double-twins that are supposed to result from a double-step compression-extension mechanism. The twinning modes were historically determined from the trace of the habit planes in deformed single crystals [23], and they are now currently and automatically identified by Electron Back Scatter Diffraction (EBSD) by measuring the specific misorientations between a parent grain and the twin (see ref. [24] for example). The twin boundaries of these three twinning modes are all characterized by a rotation around the a-axis and differ only by their rotation angle: 86° for the {1012} extension twins, 56° for the {1011} contraction twins, and 38° for the {1011}-{1012} “double-twins”. Many questions concerning the twinning mechanisms remain open. The {1012} extension twins and the {1011} compression twins were on the list of the fifteen twin modes predicted by Crocker and Bevis [8], but not the “double-twins”, and many of the predicted modes could not be observed. More surprisingly, many experimental studies report that twins can form despite low or even negative Schmid factors; this effect was called “anomalous twinning” or “non-Schmid behavior”. Anomalous twinning was reported mainly for the double-twins and compression twins [23][25][26][27], and more recently for the extension twins [28]-[31]. Another unsolved question is the atomic displacements during twinning. As mentioned in the previous section, according to the classical “shear” theory, the atoms at the lattice nodes follow the homogeneous lattice shear displacements, and the other atoms shuffle. Even if it is agreed that both shear and shuffles are concomitant, the “shear” theory first establishes the lattice shear, and only then estimates the shuffles without rigorous justification. Very recently, Yu et al. [32] developed a two-step rotation-shear model of the atomic trajectories during extension twinning in which the shear and shuffle displacements occur simultaneously. The model we will propose can be understood as a one-step model, where the combination of rotation and shear appears naturally as a consequence of the hard-sphere assumption.
Another point is puzzling. Recently, Liu et al. [33][34] reported in-situ transmission electron microscopy (TEM) observations of extension twins in a submicron-sized single crystal magnesium pillar induced by compression along the [110] axis. Surprisingly, the orientation relationship between the parent crystal and its twin is a rotation of 90° around the a-axis, in place of the expected 86° rotation. In addition, the parent/twin interface is made of basal/prismatic terrace-like interfaces instead of a straight boundary along a (1012) plane. They also observed that the interface can “propagate” reversibly without obvious shear. These results made the authors assume that the extension twins they observed were not produced by a simple shear but by a “newly discovered deformation mode”. This mode implies a direct conversion between the basal and prismatic planes of the parent crystal and its twin resulting in a “tetragonal compression” of the lattice, which was called “unit cell reconstruction”. This “reconstructive” model contradicts the disconnection theory, but is also unable to account for the displace nature of mechanical twinning; it was received with reservations [10][35]. For example, Ostapovets et al. [35] gave another explanation to the experimental results. They showed that the deformation matrix is the stretch component that appears in the polar decomposition of the usual simple shear matrix, and they proposed that the 90° twins are “produced by an average of two conjugate simple shears”. However, the traces of the two compensating shearing planes could not be observed by Liu et al. in the submicron-sized pillars. The controversy that then emerged between the partisans of the disconnection theory and those of the reconstructive model is explained in details in Ref.[36]. Most of the recent debates are linked to the formation of prismatic/basal interfaces. However, to the point view explained in the preamble, the interface features are a consequence of the mechanism and not the cause; and we estimate that it is only when the mechanism is understood that the structure of the dislocations at the interface and in the surrounding matrix can be deduced. Neither the disconnection theory nor the reconstructive model can account for the displace nature for deformation twinning because the concept of homogenous lattice distortion is voluntarily ignored. To our opinion, it is great time to rehabilitate this concept that was at the core of crystallographic metallurgy for more than a century and that seems to have fallen out of favor for these last decades in some scientific communities.

1.3. Objectives of the study

The aim of the present paper is to propose an alternative theory to the “shear”, “disconnection” and “reconstructive” theories in the case of the extension twins in magnesium and other hcp metals with c/a ratio close to the ideal ratio. This approach shares with the shear theory the concept of homogenous distortion in order to preserve the displace nature of the transformation. It follows our recent developments in the crystallography of martensitic transformation between the fcc, bcc and hcp phases [21] [22]. The model considers that the atoms are hard spheres of constant size and that the lattice distortion must respect the size of the atoms. The calculations will prove that the habit plane of the {1012} extension twins is not fully invariant during the distortion, and that the unit volume and distances in the twinning plane are not constant in the transient states. Such a volume change can indeed be expected because it is known from Kepler (17th-century) that the densest packings of hard spheres are only the hcp and fcc ones, which means that the intermediate states between the initial parent hcp and its final hcp should be less dense [37]. The paper responds to the questions on how the volume change occurs and what is its amplitude. It will be shown that the change of reticular distances are small (a few percent) but larger than the elastic strain of magnesium (~0.3 %). This implies that extension twinning in magnesium can’t be elastically accommodated and that there is no way for the atoms to follow a simple shear path during the twinning distortion, even by taking into account an elastic deviation from the hard sphere assumption. The model will be used to rigorously determine the orientational and distortional twinning variants. In analogy with martensitic transformations and ferroelectric domains, orientation continuities will be expected between the low-misorientated twin variants. They will be experimentally confirmed by Electron Back Scatter Diffraction (EBSD) maps acquired on a single crystal of magnesium. As extension twinning differs from simple shearing, an energy criterion generalizing the Schmid factor will be proposed. It will be shown to predict twinning for “anomalous” conditions, in agreement with some experimental results reported in literature.

2. Notations, calculation rules, and experimental details

The three-index notation in the hexagonal system is preferentially chosen for the calculations. The planes will be sometimes written in four-index notation, but mainly to refer to literature. A reader not familiar with the conversion between the three-index and four-index notations can refer to classical textbooks [38]; special
attention should be given to the conversion of directions which is more complicated than for planes. The vectors are noted by bold lowercase letters and the matrices by bold capital letters. In order to calculate the continuous paths of the atoms, we made the approximation that the atoms are hard spheres of constant size. The ratio of lattice parameters is then the ideal hcp ratio:

$$\gamma = c/a = \sqrt{8/3}$$

This approximation is a good start for a model of magnesium because this metal is hcp with a ratio $\gamma = 1.623$ very close to the ideal ratio of hard-sphere packing.

We call $\mathbf{B}_{\text{hex}} = (a, b, c)$ the usual hexagonal basis, and $\mathbf{B}_{\text{ortho}} = (x, y, z)$ the orthonormal basis represented in Fig. 1 and linked to $\mathbf{B}_{\text{hex}}$ by the coordinate transformation matrix $\mathbf{H}_{\text{hex}}$:

$$\mathbf{H}_{\text{hex}} = \left[ \mathbf{B}_{\text{ortho}} \rightarrow \mathbf{B}_{\text{hex}} \right] = \begin{pmatrix} 1 & -1/2 & 0 \\ 0 & \sqrt{3}/2 & 0 \\ 0 & 0 & \gamma \end{pmatrix}$$

In order to follow the displacements of the atoms during extension twinning, some labels are given to the atomic positions, as illustrated in Fig. 1. We note O, the “zero” position that will be let invariant by the distortion. We call X, Y and Z the lattice parameters, i.e. $\mathbf{OY} = a + 2b = [120]_{\text{hex}}$ and $\mathbf{OZ} = c = [001]_{\text{hex}}$. It can be checked with the matrix $\mathbf{H}_{\text{hex}}$ that $\mathbf{OX} = [100]_{\text{ortho}}$ and $\mathbf{OU} = O + [100]_{\text{ortho}}$ and $\mathbf{OZ} = [0 0 \gamma]_{\text{ortho}}$. The nodes O, X, Y, Z define a non-primitive cell that will be noted XYZ. Other vectors are noted $\mathbf{OP} = [5/3, 4/3, 1/2]_{\text{hex}}$. The atom at the center of the face (O, X, Y, S) is noted M, and the atom close to the face (O, X, Z, T) is noted N; their position vectors are $\mathbf{OM} = [110]_{\text{hex}}$ and $\mathbf{ON} = [2/3, 1/3, 1/2]_{\text{hex}}$. At the same z-level as N, there are the atoms P and Q given by $\mathbf{QP} = [5/3, 4/3, 1/2]_{\text{hex}}$ and $\mathbf{OQ} = [2/3, 4/3, 1/2]_{\text{hex}}$. The bases in which the vectors and matrices are expressed are specified in the text or as indices in the equations.

To describe the crystallography of extension twinning $p \rightarrow t$ from a parent crystal $p$ to its twin $t$, three important matrices will be used: the distortion matrix, the coordinate transformation matrix, and the correspondence matrix. Let us briefly explain them.

The distortion matrix $\mathbf{D}$ gives the image $x'$ of a vector $x$ by a linear distortion: $x' = \mathbf{D}x$. The displacement field is given by $x'-x = (\mathbf{D}-\mathbf{I})x$ where $\mathbf{I}$ is the 3x3 identity matrix. The deformation matrix is given by its gradient; it is simply the matrix $\mathbf{D}\mathbf{I}$. The letter $\mathbf{F}$ is often used in place of $\mathbf{D}$ in the textbooks on finite strain theory. The letter $\mathbf{U}$ is sometimes preferred to specify that the matrix is symmetric. The distortion matrix can be calculated as follows. The vectors of the initial parent basis are transformed by the distortion into new vectors: $\mathbf{a}_p \rightarrow \mathbf{a}'_p$, $\mathbf{b}_p \rightarrow \mathbf{b}'_p$ and $\mathbf{c}_p \rightarrow \mathbf{c}'_p$. The distortion matrix $\mathbf{D}_{\text{hex}}^{p\rightarrow t}$ is the matrix formed by the images $\mathbf{a}'_p$, $\mathbf{b}'_p$, and $\mathbf{c}'_p$ expressed in the initial hexagonal basis, i.e. $\mathbf{D}_{\text{hex}}^{p\rightarrow t} = \left[ \mathbf{B}_{\text{hex}}^{p} \rightarrow \mathbf{B}_{\text{hex}}^{t} \right] = \mathbf{B}_{\text{hex}}^{t}$ with $\mathbf{B}_{\text{hex}}^{t} = (\mathbf{a}_t, \mathbf{b}_t, \mathbf{c}_t)$ and $\mathbf{B}_{\text{hex}}^{p} = (\mathbf{a}_p, \mathbf{b}_p, \mathbf{c}_p)$. The distortion matrix is simply expressed by writing in column the coordinates of $\mathbf{a}'_p$, $\mathbf{b}'_p$ and $\mathbf{c}'_p$ in the basis $\mathbf{B}_{\text{hex}}^{t}$. The crystallographic studies on displacive phase transformations and mechanical twinning often consist in finding the distortion matrices close to the identity matrix in order to minimize the atomic displacements.

If the distortion is known in the basis $\mathbf{B}_{\text{ortho}}$, and noted $\mathbf{D}_{\text{ortho}}^{p\rightarrow t}$, a formula of coordinate transformation can be used to express it in the basis $\mathbf{B}_{\text{hex}}$; it is:

$$\mathbf{D}_{\text{hex}}^{p\rightarrow t} = \mathbf{H}_{\text{hex}}^{-1} \mathbf{D}_{\text{ortho}}^{p\rightarrow t} \mathbf{H}_{\text{hex}}$$

with $\mathbf{H}_{\text{hex}}$ given by equation (2). Inversely, if the distortion matrix is found in $\mathbf{B}_{\text{hex}}$ and it can be written in $\mathbf{B}_{\text{ortho}}$ by the inverse formula:

$$\mathbf{D}_{\text{ortho}}^{p\rightarrow t} = \mathbf{H}_{\text{hex}} \mathbf{D}_{\text{hex}}^{p\rightarrow t} \mathbf{H}_{\text{hex}}^{-1}$$

The coordinate transformation matrix $\mathbf{T}^{p\rightarrow t}$ allows the change of the coordinates of a fixed vector between the parent and twin bases. It is given by the vectors forming the basis of the twin $\mathbf{B}_{\text{hex}}^{t} = (\mathbf{a}_t, \mathbf{b}_t, \mathbf{c}_t)$ expressed in the parent hexagonal basis, i.e. $\mathbf{T}^{p\rightarrow t} = \left[ \mathbf{B}_{\text{hex}}^{p} \rightarrow \mathbf{B}_{\text{hex}}^{t} \right]$ and is calculated from the orientation relationship
between the parent and its twin experimentally obtained from TEM or EBSD. The coordinate transformation
matrix for the reverse twinning operation is simply \( T^{-1} \). The correspondence matrix \( C^{t-p} \) gives the images of the parent basis vectors by the distortion, i.e. \( \mathbf{a}_p', \mathbf{b}_p' \) and \( \mathbf{c}_p' \), expressed in the twin basis. These images are obtained from the coordinate transformation matrix and the distortion matrix: 

\[
C^{t-p} = \begin{pmatrix} \mathbf{a}_p' & \mathbf{b}_p' & \mathbf{c}_p' \end{pmatrix} / \begin{pmatrix} \mathbf{a}_p & \mathbf{b}_p & \mathbf{c}_p \end{pmatrix} = T^{t-p} \frac{\begin{pmatrix} \mathbf{a}_p' & \mathbf{b}_p' & \mathbf{c}_p' \end{pmatrix}}{\begin{pmatrix} \mathbf{a}_p & \mathbf{b}_p & \mathbf{c}_p \end{pmatrix}} = T^{t-p} B^p_{\text{hex}} = T^{t-p} D^{p-t}. 
\]

The correspondence matrix is thus:

\[
C^{t-p} = T^{t-p} D^{p-t} \tag{5}
\]

The correspondence matrix is used to calculate in the twin basis the coordinates of the image by the distortion of a vector written in the parent basis, i.e.

\[
x_p' / b_{\text{hex}} = D^{p-t} x_p / b_{\text{hex}} \rightarrow x_p' / b_{\text{hex}} = C^{t-p} x_p / b_{\text{hex}} \tag{6}
\]

Most of the symbolic and numerical calculations have been performed with Mathematica. The .nb programs (one for the lattice distortion and another one for the energy criterion) are available in Supplementary Material 1a and 1b, respectively.

Although the paper is mainly theoretical, we used an EBSD map acquired for another study related to compression twinning because this map confirms the intrinsic link between the (86°, a) twins and the (90°, a) twin modes. A single crystal of magnesium, with a cylinder shape of diameter 10 mm and length 6 mm, was compressed by 5% at 100°C in direction along the c-axis with a strain rate of 0.05 s⁻¹; the experiment was performed on a Gleeble 3800 system. The deformed crystal was mechanically polished with abrasive papers and clothes with diamond particles down to 1 µm, and then electropolished at 12V with an electrolyte made of 85% ethanol with 5% HNO₃ and 10% HCl just taken out of the fridge (10°C). The EBSD map was acquired on a field emission gun (FEG) XLF30 scanning electron microscope (FEI) equipped with the system Aztec (Oxford Instruments). The sample will be shown to exhibit large extension twins. This is surprising because the single crystal was deformed in compression, but this phenomenon was already observed on Mg polycrystalline alloys [39] and Mg single crystals [40] and attributed to the unloading stage. It is also possible that the twins appeared when the samples were cut with an abrasive disk saw.

3. Distortion matrices of (90°, a) twinning

3.1. Matrix of complete distortion

The case treated in this section and in the next one aims at calculating the stretch component of the lattice distortion. The tilt needed to let \{10\overline{1}2\} plane untilted will be introduced only in a second stage in section 4. This approach is similar to that used for fcc-bcc martensite transformations in steels, where the Bain (stretch) matrix is first calculated, and then an additional rotation is added to compensate the tilt and let a line invariant [21][41]. This section directly calculates the matrix of lattice distortion in the case of a complete transformation, without considering the continuous path that leads to it. The calculation of the continuous expression of the stretch matrix will be the subject of the section 3.2.

In order to obtain the stretch component of extension twinning, we consider the case in which the axis \( a \) remains invariant, the axis \( a + 2b \) of the parent crystal is transformed into the axis \( c \) of the twin, and the axis \( c \) of the parent is transformed into the axis \( a + 2b \) of the twin:

\[
a_p' = a_t, \quad a_p' + 2b_p' = c_t, \quad c_p' = a_t + 2b_t \tag{7}
\]

which by linear combination gives

\[
a_p' = a_t, \quad b_p' = \frac{1}{2}(c_t - a_t), \quad c_p' = a_t + 2b_t \tag{8}
\]

These vectors written in columns form the correspondence matrix \( C^{t-p} \) of extension twinning expressed in the basis \( B_{\text{hex}} \).
$$C_{\text{hex}}^{t-p} = \begin{pmatrix} 1 & -1/2 & 1 \\ 0 & 0 & 2 \\ 0 & 1/2 & 0 \end{pmatrix}$$  \hspace{1cm} (9)$$

Its use is given by equation (6). It can be checked for example the vector of the parent crystal [1,2,0]_\text{hex} is transformed by distortion into the vector [0,0,1]_\text{hex} of the twin.

The coordinate transformation matrix $T_{\text{hex}}^{p-t}$ is given by the vectors of the twin basis expressed in the parent basis. In the present case, this matrix is similar to the correspondence matrix, excepted that now the metrics appears:

$$T_{\text{hex}}^{p-t} = \begin{pmatrix} 1 & -1/2 & \gamma \sqrt{3} \\ 0 & 2\gamma \sqrt{3} & 0 \\ 0 & \gamma & 0 \end{pmatrix} = \begin{pmatrix} 1 & -1/2 & 2\sqrt{2}/3 \\ 0 & 4\sqrt{2}/3 & 0 \\ 0 & 3/4\sqrt{2} & 0 \end{pmatrix} \text{ for } \gamma = \frac{8}{\sqrt{3}}$$  \hspace{1cm} (10)$$

The coordinate transformation matrix transforms the hexagonal parent basis into the hexagonal twin basis. The equivalent matrix that transforms the orthonormal basis of the parent crystal into the orthonormal basis of the twin crystal is given by

$$\bar{R}_{\text{ortho}}^{p-t} = \mathbf{H}_{\text{hex}} T_{\text{hex}}^{p-t} \mathbf{H}_{\text{hex}}^{-1} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & 0 \end{pmatrix}$$  \hspace{1cm} (11)$$

which, as expected, is a mirror symmetry across the plane Y+Z=0 or equivalently an improper rotation of axis $OX \rightarrow a_p = a_t$ and angle 90°.

The distortion matrix $U_{\text{hex}}^{p-t}$ is obtained from the correspondence and coordinate transformation matrices by using the relation (5):

$$U_{\text{hex}}^{p-t} = T_{\text{hex}}^{p-t} C_{\text{hex}}^{t-p} = \begin{pmatrix} 1 & -1/2 + \frac{\gamma}{2\sqrt{3}} & 0 \\ 0 & \frac{\gamma}{\sqrt{3}} & 0 \\ 0 & 0 & \frac{\gamma}{\sqrt{3}} \end{pmatrix} = \begin{pmatrix} 1 & -1/2 + \sqrt{2}/3 & 0 \\ 0 & 2\sqrt{2}/3 & 0 \\ 0 & 3/2\sqrt{2} & 0 \end{pmatrix} \text{ for } \gamma = \frac{8}{\sqrt{3}}$$  \hspace{1cm} (12)$$

This matrix is given in the basis $\mathbf{B}_{\text{hex}}^{p}$. It can be written in the basis $\mathbf{B}_{\text{ortho}}^{p}$ by using equation (4):

$$U_{\text{ortho}}^{p-t} = \mathbf{H}_{\text{hex}} U_{\text{hex}}^{p-t} \mathbf{H}_{\text{hex}}^{-1} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \frac{\gamma}{\sqrt{3}} & 0 \\ 0 & 0 & \frac{\gamma}{\sqrt{3}} \end{pmatrix}$$

$$\Rightarrow U_{\text{ortho}}^{p-t} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 2\sqrt{2}/3 & 0 \\ 0 & 3/2\sqrt{2} & 0 \end{pmatrix} \approx \begin{pmatrix} 1 & 0 & 0 \\ 0 & 0.94 & 0 \\ 0 & 0 & 1.06 \end{pmatrix} \text{ for } \gamma = \frac{8}{\sqrt{3}}$$  \hspace{1cm} (13)$$

The general formula (13) proves that the (90°, a) twin is an extension twin along the z-axis for $\gamma \leq \sqrt{3}$, and a contraction twin for $\gamma \geq \sqrt{3}$, as already noticed by Yoo [42].

This matrix is diagonal; it is a stretch equivalent to the Bain distortion known in martensitic transformations. Its values are $x'/x$, $y'/y$, $z'/z$; they mean that the distortion lets the x-axis invariant, shorten the y-axis by $\Delta y/y = -$
6% and extends the z-axis by $\Delta z/z = +6\%$ in the case of magnesium. The ratio of contraction $y'/y$ is exactly the inverse of the ratio of extension $z'/z$. One can now raise the question: how the two ratios evolve during the lattice distortion? Do they follow this inverse relation continuously, which would mean that the volume is unchanged during the lattice distortion? To answer this question one needs to go deeper into the details of the distortion.

3.2. Matrix of continuous distortion

In the rest of the paper, it will be assumed that the atoms are hard spheres and that these atoms are perfectly packed, i.e. $y = \sqrt{\frac{B}{3}}$. The mechanic reason for the simultaneous $y'/y$ contraction and $z'/z$ extension is the coordinated displacement of the M and N atoms. Indeed, these atoms, taken as hard spheres, are in “contact” and keep contact during their movements. The atom M initially in the basal plane $(O,X,Y,S) = (001)_b$ goes out of the plane such that after twinning this plane is transformed into the prismatic plane $(O,X,Y,S) = (010)_b$, and the atom N goes toward the prismatic plane $(010)_b$ such that after twinning this plane is transformed into the basal plane $(001)_b$, as illustrated in Fig. 1 and in Fig. 2a and b. During the displacements of the atoms M and N, the distance $OZ$ increases and the distance $OY$ decreases. It is assumed that the atoms O and X do not move, and that the atoms M and N keep contact with the atoms O and X, and between each other during their displacements. The trajectories of the atoms M and N can then be described by a unique parameter, which is the angle $\eta$ made by the vector $\textbf{IM}$ with the basal plane $(O,X,S,Y)$, i.e. $\Phi = \text{ArcCos} \left( \frac{1}{\sqrt{3}} \right) \approx 70.5^{\circ}$. During extension twinning, the angle $\eta$ increases from the start value $\eta_s = 0$ to the finish value $\eta_f = \frac{\pi}{2} - \Phi \approx 19.47^{\circ}$. The coordinates of the atoms M and N in the orthonormal basis $\textbf{B}_{ortho}$ are

$$\textbf{OM} = \begin{pmatrix} \frac{1}{2} \cos(\eta) \\ \frac{\sqrt{3}}{2} \cos(\eta) \\ \frac{\sqrt{3}}{2} \sin(\eta) \end{pmatrix} \quad \text{and} \quad \textbf{ON} = \begin{pmatrix} \frac{1}{2} \cos(\eta + \Phi) \\ \frac{\sqrt{3}}{2} \cos(\eta + \Phi) \\ \frac{\sqrt{3}}{2} \sin(\eta + \Phi) \end{pmatrix}$$

(15)

In the present case, it is supposed that $OY'$ remains parallel to $OY$ during the transformation. The atom M moves such that it keeps contact with the atoms O, X and Y. Thus, as shown in Fig. 2d, the point J in the middle of OY keeps the same y-coordinate as M, i.e. $OY = 2 OJ = \sqrt{3} \cos(\eta)$. It is also assumed that OZ' remains parallel to OZ. The atom N moves such that it keeps contact with the atoms O, X and Z. Thus, as shown in Fig. 2d, the point K in the middle of OZ keeps the same z-coordinate as N, i.e. $OZ = 2 OK = \sqrt{3} \sin(\eta + \Phi)$. Let us write the vectors $\textbf{OX}$, $\textbf{OY}$ and $\textbf{OZ}$ forming the basis $\textbf{B}_{XYZ}$ of the XYZ cell in the orthonormal basis $\textbf{B}_{ortho}$. The coordinate transformation matrix between these bases is

$$[\textbf{B}_{ortho} \rightarrow \textbf{B}_{XYZ}(\eta)] = \textbf{B}_{XYZ}(\eta) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \sqrt{3} \cos(\eta) & 0 \\ 0 & 0 & \sqrt{3} \sin(\eta + \Phi) \end{pmatrix}$$

(16)

The continuous distortion matrix at each step $\eta$ of the distortion is given in the basis $\textbf{B}_{ortho}$ by the matrix $\textbf{U}^{p-t}_{ortho}(\eta) = \textbf{B}_{XYZ}(\eta) \cdot \textbf{B}^{-1}_{ortho}(\eta_0)$ (see equation 1 of Ref.[1]). The calculation leads to

$$\textbf{U}^{p-t}_{ortho}(\eta) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \cos(\eta) & 0 \\ 0 & 0 & \frac{3}{2\sqrt{2}} \sin(\eta + \Phi) \end{pmatrix}$$

(17)

In order to get simpler expressions, we introduce the variable $\kappa = \sin(\eta)$. The start value becomes $\kappa_s = 0$ and the finish value $\kappa_f = 1/3$. The distortion matrix is now a function of $\kappa$:

$$\textbf{U}^{p-t}_{ortho}(\kappa) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \sqrt{1 - \kappa^2} & 0 \\ 0 & \kappa \frac{3}{2\sqrt{2}} + \sqrt{1 - \kappa^2} \end{pmatrix}$$

(18)
The matrix of complete transformation is given for $\kappa_f = 1/3$; it can be checked that it is the matrix already given in equation (14). This proves that the ratio of contraction $y'/y$ is the inverse of the ratio of extension $z'/z$ only in the final state, but not during the intermediate states of the distortion.

The distortion matrix can be written in the hexagonal basis $\mathbf{B}_{\text{hex}}$ by using equation (3):

$$
\mathbf{U}_{\text{hex}}^{p-t}(\kappa) = \begin{pmatrix}
1 & -1 + \sqrt{1 - \kappa^2} & 0 \\
0 & \sqrt{1 - \kappa^2} & 0 \\
0 & 0 & \frac{\kappa}{2\sqrt{2}} + \sqrt{1 - \kappa^2}
\end{pmatrix}
$$

(19)

The ratio of volume change $V'/V$, where $V$ is the initial volume of the XYZ cell and $V'$ is the volume of the distorted cell, is directly given by the determinant of the distortion matrix:

$$
\frac{V'}{V}(\kappa) = \det(\mathbf{U}_{\text{hex}}^{p-t}) = \det(\mathbf{U}_{\text{ortho}}^{p-t}) = 1 - \kappa^2 + \frac{\sqrt{2}}{4}\kappa \sqrt{1 - \kappa^2}
$$

(20)

The curve $V'/V$ is presented in Fig. 3. The maximum of volume change, close to 1.0303 is obtained for the intermediate value $\kappa_i \approx 0.1691 (\eta_i \approx 9.73^\circ)$. It proves that, although it returns to its initial value when the distortion is complete $V/V(\kappa_f) = 1$, the volume is not constant during the twinning process. A similar conclusion was drawn on fcc-fcc mechanical twinning in a previous study (sections 3 and 7.5 of Ref.[1]).

The matrix $\mathbf{U}_{\text{hex}}^{p-t}(\kappa)$ gives the stretch (Bain) matrix during the process of extension twinning. The atoms at the first corners of the XYZ cell (i.e. O, X, Y, Z) follow a trajectory whose equation in the basis $\mathbf{B}_{\text{ortho}}$ is directly given by this matrix, with $\kappa$ continuously varying from $\kappa_s = 0$ to $\kappa_f = 1/3$. The atoms M, N, Q inside the XYZ cell do not follow the same trajectory; they “shuffle”. The trajectory equation of M and N is deduced from equation (15). Actually, the initial position vectors in the basis $\mathbf{B}_{\text{ortho}}$, which are $\mathbf{O} = [1, \frac{\sqrt{3}}{2}, \frac{\sqrt{2}}{2}, 0]$, $\mathbf{O}N = [0, \frac{2\sqrt{3}}{3}, \frac{\sqrt{2}}{3}, 0]$, are all rotated by the same “shuffle rotation” that is simply

$$
\mathbf{R}_{\text{shuffle}}(\eta) = \begin{pmatrix}
1 & 0 & 0 \\
0 & \cos(\eta) & -\sin(\eta) \\
0 & \sin(\eta) & \cos(\eta)
\end{pmatrix}
$$

(21)

Consequently, in the XYZ cell, the corner atoms O, X, Y, Z, S, T, U, V (each of them count for 1/8 in the cell) follow the distortion (18), and the atom M (counts for 1/4), N (counts for 1), Q (counts for 1/2) follow the shuffling rotation (21). This means that twinning is obtained with 1/3 of distortion and 2/3 of shuffle.

Some movies displaying the atomic displacements in a magnesium crystal during the twinning transformation were computed with VPython. The first movie (Supplementary Material 2) shows the transformation of a Bravais unit cell and the second movie (Supplementary Material 3) shows the transformation of a 4x4x4 XYZ supercell. Three snapshots taken from the initial, intermediate and final states are extracted from the two movies; they are given in Fig. 4 and Fig. 5, respectively.

4. Distortion matrices of (86°, a) twinning

4.1. Compensating angle and matrix of continuous distortion

The distortion matrix (18) is diagonal and only one value equals 1, which means that there is no plane that is let invariant. The orientation relationship between the parent and its twin is a (90°, a) (improper) rotation, as it was observed in submicron-sized Mg single-crystal [33]. However, in bulk magnesium, it is known that extension twins let “invariant” the \{1012\} planes, which implies that the orientation relationship is (86°, a) and not (90°, a). In order to make this plane untilted during the lattice distortion, a rotation should be added to the matrix (18), in the same way as a rotation is added to the Bain tensor in fcc-bcc transformations in steels (see for example ref. [41]). Since the direction $\mathbf{OX} = a$ is already invariant, the (01̅12) plane can be
continuously maintained until by compensating the rotation of the direction \( \mathbf{OV} = [121]_{\text{hex}} \parallel [0\ 3\ 2\sqrt{2}]_{\text{ortho}} \). Let us call \( \xi \) the angle \( (\mathbf{OV}, \mathbf{OV}') \) illustrated in Fig. 6a. The cosine of this angle \( C_\xi \) is calculated by the scalar product \( \mathbf{OV}_{\text{ortho}} \cdot \mathbf{U}_{\text{ortho}}^{\text{p-t}} \mathbf{OV}_{\text{ortho}} \) using the matrix (18). It is

\[
C_\xi(k) = \frac{2\sqrt{2}k + 17\sqrt{1 - k^2}}{\sqrt{17(1 - 16k^2 + 4k\sqrt{2} - 2k^2)}}
\]

The angle \( \xi \) varies from 0 to \( \xi_f = 3.4^\circ \) during the diagonal distortion. The rotation matrix that compensates the angle \( \xi \) is thus given in the basis \( \mathbf{B}_{\text{ortho}} \) by

\[
\mathbf{R}(k) = \begin{pmatrix}
1 & 0 & 0 \\
0 & C_\xi(k) & 0 \\
0 & -\sqrt{1 - C_\xi(k)^2} & C_\xi(k)
\end{pmatrix}
\]

We point out here that only the rotation of the direction \( \mathbf{OV} \) can be cancelled, but not its length change. Indeed, the distance \( ||\mathbf{OV}(k)|| = ||\mathbf{U}_{\text{ortho}}^{\text{p-t}}(k)\mathbf{OV}|| \) calculated from the matrix (18) is not constant. The ratio of the length \( \mathbf{OV}' \) divided by its initial value \( \mathbf{OV} \) is

\[
\frac{\mathbf{OV}'}{\mathbf{OV}}(k) = \frac{\sqrt{17 - 16k^2 + 4k\sqrt{2} - 2k^2}}{\sqrt{17}}
\]

Its graph is given in Fig. 6b. This proves that, although the length \( \mathbf{OV} \) returns to its initial value when the distortion is complete, the distance is not constant during the twinning process. Consequently, although the plane \((0\bar{1}12)\) remains untilted and is eventually restored, this plane is \textit{not} fully invariant during the process. Strictly speaking, one should say that the twinning plane \((0\bar{1}12)\) is \textit{globally} invariant (untilted and restored).

The distortion matrix that lets the \((0\bar{1}12)\) plane untilted is \( \mathbf{D}_{\text{ortho}}^{\text{p-t}}(k) = \mathbf{R}(k) \mathbf{U}_{\text{ortho}}^{\text{p-t}}(k) \). The calculations show that

\[
\mathbf{D}_{\text{ortho}}^{\text{p-t}}(k) = \begin{pmatrix}
1 & 0 & 0 \\
0 & \frac{17 - 17k^2 + 2k\sqrt{2} - 2k^2}{\sqrt{17}d(k)} & \frac{3k(2\sqrt{2} - 2k^2)}{2\sqrt{34}\ d(k)} \\
0 & -\frac{3k\sqrt{1 - k^2}}{\sqrt{17}d(k)} & \frac{68 - 64k^2 + 25k\sqrt{2} - 2k^2}{4\sqrt{17}d(k)}
\end{pmatrix}
\]

with \( d(k) = \sqrt{17 - 16k^2 + 4k\sqrt{2} - 2k^2} \)

The matrix \( \mathbf{D}_{\text{ortho}}^{\text{p-t}}(k) \) is the full form of the lattice distortion during extension twinning. The displacements of the atoms at the corners of the XYZ cell (i.e. O, X, Y, Z) follow a trajectory whose equation in the basis \( \mathbf{B}_{\text{ortho}} \) is directly given by this matrix, with \( \kappa \) continuously varying from \( \kappa_e = 0 \) to \( \kappa_f = 1/3 \). The atoms M, N, Q shuffle in the XYZ cell. Their trajectories are given by the “shuffling rotation” (21) compensated of the angle \( \xi \), i.e.

\[
\mathbf{R}_{\text{shuffle}}(\eta) = \begin{pmatrix}
1 & 0 & 0 \\
0 & \cos(\eta - \xi) & -\sin(\eta - \xi) \\
0 & \sin(\eta - \xi) & \cos(\eta - \xi)
\end{pmatrix}
\]

with \( \xi \) function of \( \eta \) by equation (22).

As the distortion matrix (25) is the product of a rotation matrix by the stretch matrix (18), the volume change is the same as the one given in equation (20). The changes of the unit volume and of the distance \( \mathbf{OV} \) in \((0\bar{1}12)\) plane mean that simple shear matrices are not the most appropriate tool to describe the continuous paths of the atoms.
A movie of the transformation of $4\times4\times4$ XYZ cell is reported in Supplementary Material 4. Three snapshots taken at initial, intermediate and final states are extracted; they are given in Fig. 5. The $(0\bar{1}1\bar{2})$ plane is not tilted during the transformation. Its intra-planar distortion in the transient states due to the change of length OV (+1.4\%) is hardly perceptible.

4.2. Complete form of the distortion and orientation matrices

When the transformation is complete, the distortion matrix takes the value

$$D_{ortho}^{p-t} = D_{ortho}^{p-t}(1/3) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 16 & 3 \\ 0 & 2\sqrt{2} & 34\sqrt{2} \end{pmatrix}$$

(27)

In the hexagonal basis $B_{hex}$, it becomes, by using equation (3),

$$D_{hex}^{p-t} = \begin{pmatrix} 1 & -\frac{1}{34} & 1 \\ 0 & 16 & 2 \\ 0 & -1 & 18 \end{pmatrix}$$

(28)

In the reciprocal space, one must take the inverse of the transpose:

$$\left(D_{hex}^{p-t}\right)^* = \left(D_{hex}^{p-t}\right)^{-T} = \begin{pmatrix} 1 & 0 & 0 \\ 1 & 18 & 1 \\ \frac{1}{34} & 17 & \frac{34}{17} \\ -1 & 2 & 16 \end{pmatrix}$$

(29)

The distortion matrix in the direct space and hexagonal basis given by equation (28) has only one eigenvalue equal to 1 and an infinity of eigenvectors that are all linear combinations of the two vectors $OX = [100]_{hex}$ and $OV = [121]_{hex}$. This means that the distortion matrix (28) is a simple shear matrix on the plane $(0\bar{1}2)_{hex}$. The shear coefficient $s$ is the tangent of the angle made by the vector $n$ normal of the shear plane with its image. It is easily calculated with $n = [0, -\sqrt{3}]_{ortho}$ and its image by the matrix (27); it is $s = \frac{1}{6\sqrt{2}} \approx 0.118$, as expected from the theoretical value of shear [2]. The shear vector $s$ is

$$s = (D_{ortho}^{p-t} - I)\frac{n}{\|n\|} = \frac{1}{34} \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ -\sqrt{2} & 3 \end{pmatrix}_{ortho} = \frac{1}{2\sqrt{102}} [121]_{hex}$$

(30)

The vector $s$ makes an angle of 43.31° with the basal plane. It can be checked that the vector $s$ belongs to the “shear” plane $(0\bar{1}2)_{hex}$. The distortion matrix expressed in the reciprocal space is given by equation (29); it has only one eigenvalue equal to 1 and an infinity of eigenvectors that are all linear combinations of the two vectors $(10\bar{1})_{hex}$ and $(\bar{1}20)_{hex}$. This means that, in addition to the shear plane $(0\bar{1}2)_{hex}$, all the planes that contain the shear vector $s$ are also invariant, as expected from a shear matrix. These calculations show that the continuous approach allows us to find the classical result (a shear matrix) in the special case where only the initial and final states are considered. To our knowledge, it is the first time that a continuous analytical form of the distortion matrix is given for a deformation twinning mode in hcp metals.

The orientation of the twin is equal to that of the $(90°, a)$ twins corrected of the $3.4°$ angle, i.e. it is $(86.6°, a)$, simply noted in the paper $(86°, a)$. This result can also be obtained by calculating the coordinate transformation matrix

$$T_{hex}^{p-t} = D_{hex}^{p-t} \left(C_{hex}^{c-p}\right)^{-1}$$

(31)
with \( \mathbf{D}^{p-t} \) given by the equation (28) and \( \mathbf{C}^{r-p} \) by the equation (9). In order to obtain the rotation matrix, the matrix \( \mathbf{T}_{\text{hex}}^{p-t} \) must be composed with the mirror symmetry \( \mathbf{M}_{\text{hex}} \) across the basal plane and then be expressed in the orthonormal basis. The result is

\[
\mathbf{R}^{p-t} = \mathbf{H}_{\text{hex}} \mathbf{M}_{\text{hex}} \mathbf{T}_{\text{hex}}^{p-t} \mathbf{H}_{\text{hex}}^{-1} = \begin{pmatrix}
1 & 0 & 0 \\
0 & \frac{1}{17} & \frac{12\sqrt{2}}{17} \\
0 & \frac{12\sqrt{2}}{17} & \frac{1}{17}
\end{pmatrix}
\]

(32)

The matrix \( \mathbf{R}^{p-t} \) is indeed a rotation around the \( \mathbf{a} \)-axis of angle \( \text{ArcCos}(1/17) \approx 86.6^\circ \approx 90^\circ - \xi_p \).

5. Orientational and distortional variants

The orientational variants are the twin variants that are similarly orientated with the parent crystal. At maximum, there are as many variants as symmetries in the point group of the parent phase \( \mathcal{G} \), but often the number is lower due to the fact that, for some special orientation relationships \( \mathbf{T}^{p-t} \), some symmetries are common to the parent crystal and its twins. These common symmetries form a subgroup of \( \mathcal{G} \), called “intersection group” \( \mathcal{H} \). In the case of deformation twinning

\[
\mathcal{H} = \mathcal{G} \cap \mathbf{T}^{p-t} \mathcal{G} \left( \mathbf{T}^{p-t} \right)^{-1}
\]

(33)

The orientational twin variants \( \mathcal{I} \), can be mathematically identified to the left-cosets based on \( \mathcal{H} \), i.e. \( \mathcal{G}_{\mathcal{I}} \), and their number is given by the Lagrange’s formula; it is the order of the parent point group \( \mathcal{G} \) divided by the order of the intersection group \( \mathcal{H} \). The distinct orientations are

\[
\mathbf{T}^{p-t} \mathcal{I} = g_i \mathbf{T}^{p-t} \quad \text{with} \quad g_i \text{ a symmetry arbitrarily chosen in the coset } g_i \mathcal{H}
\]

(34)

The misorientations between the variants, called “operators”, are identified the double-cosets based on the intersection group, \( \mathcal{H} g_i \mathcal{H} \); their number is given by the Burnside’s formula. The variants and their operators form an algebraic structure called groupoid. More details on these concepts are given in Ref. [43].

The distortional variants are defined similarly as the orientational variants, but by replacing the orientation relationship matrix \( \mathbf{T}^{p-t} \) by the distortion matrix \( \mathbf{D}^{p-t} \) [22]. The intersection group is

\[
\mathcal{K} = \mathcal{G} \cap \mathbf{D}^{p-t} \mathcal{G} \left( \mathbf{D}^{p-t} \right)^{-1}
\]

(35)

The distortional variants are the left-cosets based on the subgroup \( \mathcal{K} \), i.e. \( \mathcal{G}_{\mathcal{K}} \). The matrices of distinct distortions are

\[
\mathbf{D}^{p-t} \mathcal{I} = g_i \mathbf{D}_0^{r-t} (g_i)^{-1} \quad \text{with} \quad g_i \text{ a symmetry arbitrarily chosen in the coset } g_i \mathcal{K}
\]

(36)

The number of distortional variants is also given by the Lagrange’s formula. It is not always equal to the number of orientational variants (see the case of fcc-hcp transition in Ref. [22]).

The point group of hcp metals is \( \mathcal{G} = 6/mmm \); it is constituted of 24 symmetry matrices, that are made of 0, 1 or -1 when expressed in the hexagonal frame. The calculations of the extension twinning variants are direct. They were done geometrically and algebraically. The results were also confirmed by the software GenOVA [44] (Supplementary Material 5).

In the case of the (90°, \( \mathbf{a} \)) twin, the intersection group \( \mathcal{H} \) is constituted by the identity, the inversion, the 2-fold rotation around the \( \mathbf{a} \)-axis, the mirror symmetry across the basal plane, and their combinations. This geometrical result is also numerically obtained by using formula (33) with the coordinate transformation matrix \( \mathbf{T}^{p-t} \) given in equation (12). The number of twin variants is thus \( 24/8 = 3 \). For distortional variants, the...
intersection group $\mathbb{K}$ is calculated by using equation (35) with the distortion matrix (12). The calculations prove that $\mathbb{K} \equiv \mathbb{H}$, and consequently, there are also three distortional variants.

In the case of the (86°, a) extension twins, the mirror symmetry across the basal plane is not anymore a common symmetry and consequently the intersection group $\mathbb{H}$ is reduced to four symmetry matrices: the identity, the inversion, the 2-fold rotation around the a-axis and the mirror symmetry across the plane normal to the a-axis, i.e. the (2110) plane. This result is also obtained numerically by using formula (33) with the coordinate transformation matrix $T_{\text{p-t}}$ given in equation (31). The number of twin variants is thus $24/4 = 6$. Here again, the calculations prove that $\mathbb{K} \equiv \mathbb{H}$, and thus, the number of distortional variants is also equal to six. The six twin variants can be grouped into 3 pairs of low-disorientated variants. For example, a pair is constituted by the variants $t_1 = \mathbb{H}$ and $t_5 = g_5 \mathbb{H}$ (see Supplementary Material 5). These variants are linked by the operator $O_2$ which contains the rotation of $2\Delta \theta = 7.4^\circ$ around the a-axis. The distortion matrices associated with these variants are

$$D_{\text{hex}}^{\text{p-t1}} = \begin{pmatrix} 1 & -\frac{1}{34} & \frac{1}{17} \\ 0 & \frac{16}{17} & \frac{2}{17} \\ 0 & -\frac{1}{34} & \frac{17}{16} \end{pmatrix}$$

and

$$D_{\text{hex}}^{\text{p-t5}} = g_5 D_{\text{hex}}^{\text{p-t1}} g_5^{-1} = \begin{pmatrix} 1 & -\frac{1}{34} & -\frac{1}{17} \\ 0 & \frac{16}{17} & \frac{2}{17} \\ 0 & \frac{1}{34} & \frac{17}{16} \end{pmatrix}.$$  

The difference between the two matrices is small ($4/17 = 0.23$). This result can be explained by the fact that the two (86°, a) variants $t_1$ and $t_5$ are derived from the same (90°, a) variant by the same distortion matrix (12) and differ only by the sign of the correction angle. An analogy with ferroelectrics and martensitic transformations will be discussed in the next section.

6. Discussion

6.1. A unique mechanism for the 90° and 86° extension twin domains

The classical “shear” model of deformation twinning is based on the lattice correspondences resulting from simple shears; but the atomic displacements are not really taken into account, and the shuffles are inferred only after determining the lattice shear. The present model reverses the order of thinking. The atomic trajectories are calculated in order to restore the crystal structure in a special orientation relationship. The magnesium atoms are assumed to be hard spheres of constant size and their displacements are chosen as low as possible. These assumptions constrain the calculations such that only one parameter, a distortion angle $\eta$, is sufficient to follow the trajectories of all the atoms during the process of extension twinning. The distortion matrix is calculated as an analytical function of the angular parameter $\eta$ only once the atomic displacements are known. First, the calculations were performed by assuming that the final parent/twin orientation is (90°, a), as observed by Liu et al. in submicron-sized Mg pillars [33][34]. In such a case, the distortion matrix is diagonal, i.e. it corresponds to a stretch deformation. However, contrarily to the interpretation based on unit cell reconstruction, our model is dispersive, as expected for a process implying rapid and collective atomic displacements. Then, we showed that the distortion matrix that lets “invariant” the (0112) plane differs from the stretch form only by a small “compensating” rotation of angle that varies continuously from 0 to of 3.4°. In other words, the diagonal distortion matrix related to Liu et al.’s observations is the symmetric matrix that can be extracted by polar decomposition from the usual twinning matrix, as already shown by Ostapovest et al. [35], in a similar way that the Bain tensor appears as a component of the lattice distortion in the fcc-bcc martensitic transformation in steels [21]. However, we don’t believe that the Liu et al.’s observations result from an average of two twinning shear on different conjugate (1012) planes, as proposed by Ostapovest et al.

We think that the observed (90°, a) twins and related stretching observed in the magnesium nano-pillars actually result from a real “natural” mechanism that is free to appear due the small size of the sample. The interface strains required for the basal/prism and prism/basal interfaces (-6% along a+2b and +6% along c) are probably more easily accommodated in the “free” submicron-sized pillar than in the bulk samples. In bulk single crystal or polycrystalline magnesium, such strains are probably too high and need to be reduced by a slight crystal rotation of 3.4° that permits to let the (1012) habit plane untilted. The distortion mechanism in the submicron pillars and in bulk samples are actually very similar because the distortion matrices (12) and (28) have close numerical values.
6.2. Analogies and expectations

To our point of view, there is a strong analogy between the extension twins in magnesium and the orientational domains in ferroelectrics. Let us explain. If, for sake of simplicity, we represent the cubic→tetragonal transition in a ferroelectric crystal by a square→rectangle distortion along the x and y-axes, four domains misoriented by 90° and 180° rotations should be formed, as illustrated in Fig. 7a. However, experience shows that the domains are actually oriented such that they share one of their two diagonals. These domains are thus actually height domains, and among them, the 90° misorientation angle is lost and replaced by 90°-2δ, with δ = ArcTan(a0/c) and a and c the lattice parameters of the tetragonal phase, as shown in Fig. 7b. The angle δ compensates the rotation of the diagonal induced by the tetragonal distortion; it is called “obliquity”. The concept of obliquity dates from Friedel’s work in 1920 [45]. It is a major parameter to define twins in minerals and ferroelectrics [46]-[48]. For example, in barium titanate, the tetragonality is close to 1% and δ is close to 0.6° [43]. The fact that the diagonal is maintained invariant is explained by strain minimization and compatibility conditions [49][50].

6.3. Experimental confirmation of the link between the (86°, a) and the (90°, a) twins

To our opinion, these features are the plastic trace left by the lattice distortion. They can be imaged a) as the consequences of the back-stresses created by the geometrically necessary dislocations (GND) generated in the surrounding matrix by the lattice distortion [52], or b) as the result of the growth of the product phase inside the surrounding parent matrix that has been rotated by the GNDs [53]. For example, continuous features were observed between the two variants in the pairs of low-misoriented Kurdjumov-Sachs (KS) variants, and the other Nishiyama-Wasserman (NW) and Pitsch variants. The 24 KS variants are formed according to a low-symmetry OR (intersection group of order 2) and the 12 NW and the 12 Pitsch variants are formed according to a higher-symmetry OR (intersection group of order 4). As we consider that deformation twinning is a displacive phase transformation, similar features are expected for the extension twins in magnesium. More specifically, it is expected that the back-stresses generated by the twinning distortions inside the surrounding matrix will create a continuous rotation between the two (low-symmetry OR) D1 and D2 variants, re-orienting them back to the initial orientation of the parent lattice and to the (high-symmetry OR) D2 variant. A continuous rotation of angle 2ζ around the a-axis is expected between the D1 and D2 variants, as schematically shown in Fig. 8b.
surrounding matrix, which explains the color gradients observed in Fig. 9d. The effect of twinning transformation is not localized at the sole parent/twin interface (Fig. 9d), and thus that the structure of disconnections at the interface is not sufficient to explain the twinning mechanism (see section 1.1).

6.4. Prediction of the formation of extension twins for negative Schmid factors

The calculations showed that there is a volume variation during extension twinning (Fig. 3). Since the distance OV also changes (Fig. 6b), the twinning plane can’t be let fully invariant during the transformation. It is true that when the transformation finishes, i) the volume comes back to its initial value, ii) the twinning plane is fully restored, and iii) the distortion matrix becomes a simple shear matrix; however, the details of the mechanism between the initial and final state shouldn’t be ignored; twinning is not shearing. What is the consequence? The Schmid factor used to calculate the resolved shear stress on the twinning plane is not adapted anymore to predict the twin formation. Is it possible to substitute the Schmid’s law by another one? Could it explain some of the abnormalities observed in the formation of the extension twins (see section 1.2)?

It is reasonable to think that the volume change observed during the lattice distortion creates an energy barrier that should be overcome to form the twins. This idea was also suggested for the fcc-bcc martensite transformations [21]. If this assumption is correct, it would mean that instead of using the matrix of complete transformation, i.e. the simple shear matrix (27), one should use the intermediate matrix corresponding to the maximum volume change. Since this matrix is not a simple shear matrix a criterion substituting the Schmid’s law should be found. We propose to come back to the general formula giving the interaction work \( W \) of a unit volume of a material that deforms, by mechanical twinning or phase transformation, inside an external stress field \( \mathbf{\Gamma} \). The scalar \( W \) is given by the Frobenius inner product

\[
W = \mathbf{\Gamma}_{ij} \cdot \mathbf{E}_{ij}
\]  

It is the integral form of the infinitesimal work \( dW = \mathbf{\Gamma}_{ij} \cdot d\mathbf{E}_{ij} \) for uncorrelated fields \( \mathbf{\Gamma} \) and \( \mathbf{E} \), sometimes called “virtual work” in the continuum mechanics textbooks. The work \( W \) is performed by the external stress during the transformation; it is used to deform the surrounding environment in which the twin forms. A high value of interaction work means a high probability of transformation, and negative value should correspond to an impossibility of transformation. The interaction energy is the opposite of \( W \); it should be added to the usual form of energy that is calculated to predict a phase transformation. Maximizing the interaction work is equivalent to minimizing the interaction energy. As proved in Supplementary Material 6, the interaction work is proportional to the Schmid factor in the case of a simple shear, and to the Patel and Cohen criterion [54] in the case of invariant plane strain deformation. Let us illustrate how equation (38) can be used for extension twinning, first by using the final (shear) distortion matrix, and then the intermediate distortion matrix.

The numerical value of the final distortion matrix is given in the basis \( \mathbf{B}_{\text{ortho}} \) by equation (27):

\[
\mathbf{D}_{f}^{\text{p-t}} = \mathbf{D}_{\text{ortho}}^{\text{p-t}} \left( \frac{1}{3} \right) \approx \begin{pmatrix}
1 & 0 & 0 \\
0 & 0.9412 & 0.0624 \\
0 & -0.0555 & 1.0588
\end{pmatrix}
\]  

Now, we consider a mechanical test on an hcp crystal that was rotated by an angle \( \theta \) around the direction \( \mathbf{n} \) normal to (0\overline{1}12) plane, and tilted by an angle \( \phi \) around the \( \mathbf{x} \)-axis. The rotation matrices axis are noted \( \mathbf{R}_{n}(\theta) \) and \( \mathbf{R}_{x}(\phi) \), respectively. As showed in the equation (E4) of Supplementary Material 6, the distortion matrix of the rotated-tilted parent crystal, expressed in the orthonormal basis \( (\mathbf{x}, \mathbf{y}, \mathbf{z}) \) linked to the mechanical test becomes

\[
\mathbf{D}_{f}^{\text{p-t}}(\phi, \theta) = \mathbf{R}_{x}(\phi) \mathbf{R}_{n}(\theta) \cdot \mathbf{D}_{f}^{\text{p-t}} \cdot (\mathbf{R}_{x}(\phi) \mathbf{R}_{n}(\theta))^{-1}
\]  

The twinning deformation matrix is \( \mathbf{E}_{f}^{\text{p-t}}(\phi, \theta) = \mathbf{D}_{f}^{\text{p-t}}(\phi, \theta) - \mathbf{I} \). According to equation (38), its interaction work with a stress field \( \mathbf{\Gamma} \) is

\[
W_{f}(\phi, \theta) = \mathbf{\Gamma}_{ij} \cdot \mathbf{E}_{ij}^{\text{p-t}}(\phi, \theta)
\]  

The graph of interaction work \( W_{f}(\phi, \theta) \) in the case of a tensile test along the \( \mathbf{z} \)-axis is given in Fig. 10a. As expected, it is similar to the graph obtained for a simple shear (Fig.6.3); the only difference is that the
interaction work is maximum for a shear plane tilted at $\phi = 45^\circ$ in the case of FigS6.3, whereas it is maximum for a crystal tilted at $\phi = 2^\circ$ for extension twinning. This is due to the fact that the (0112) shear plane is already tilted 43° far from the basal plane in a crystal positioned horizontally, which means that a tilt of $\phi_2 = 2^\circ$ is sufficient to place the shear plane at 45° far from the z-axis. According to Fig. 10a the extension twins should form only for positive interaction work, i.e. for tilt angles $\phi_2 - \frac{\pi}{4} < \phi < \phi_2 + \frac{\pi}{4}$ i.e. in the range of tilt angle $\phi \in [-43^\circ, 47^\circ]$.

What happens if the intermediate distortion matrix is used in place of the final (shear) matrix? The numerical values of the intermediate distortion matrix corresponding to maximal volume change is given in the basis $B_{ortho}$ by equation (25) with $\kappa = \kappa_1$:

$$D_i^{p+t} = D_{ortho}^{p+t}(\kappa_1) \approx \begin{pmatrix} 1 & 0 & 0 \\ 0 & 0.9852 & 0.0308 \\ 0 & -0.0290 & 1.0449 \end{pmatrix}$$

(42)

This matrix is used to calculate the intermediate distortion matrix $D_i^{p+t}(\phi, \theta)$ in a parent crystal tilted around the $x$-axis by an angle $\phi$, and rotated around the $n$-axis by an angle $\theta$, as in equation (40). The intermediate deformation matrix is $E_i^{p+t}(\phi, \theta) = D_i^{p+t}(\phi, \theta) - I$. This matrix, instead of $E_i^{p+t}(\phi, \theta)$, is now used in equation (41) to calculate the interaction work $W_i(\phi, \theta)$ at the intermediate state. The graph of $W_i(\phi, \theta)$ is given in Fig. 10b. It is clear that the region of positive work $W_i$ is extended in comparison with $W_t$. For example, for $\theta = 0$, the interaction work $W_i$ is positive in the range of tilt angles $\phi \in [-59^\circ, 59^\circ]$, whereas $W_t$ is positive only in the range $[-43^\circ, 47^\circ]$. In other words, the use of the intermediate distortion matrix allows the predictions of formation of extension twins for some domains of orientations where the classical Schmid factor is negative. The domains $(\phi, \theta)$ where $W_i \geq 0$ and $W_t < 0$ are shown in the graph of Fig. 10c. It gives the orientations where our model predicts extension twinning whereas classical shear theory does not. Since the experiments are often realized by tilting the parent crystal around the $z$-axis instead of the $n$-axis (because $n$ depends on which of the six twinning variants is formed), we have plotted in the Supplementary Material 7 the same graphs as those of Fig. 10d, but calculated by substituting the rotation $R_{n}(\theta)$ by the rotation $R_{s}(\theta)$. The range $\phi \in [-59^\circ, 59^\circ]$ predicted by our model is exactly the range of formation of extension twins deduced by Čapek et al. [30] after extrapolating a series of neutron diffraction measurements on polycrystalline magnesium samples deformed by tensile tests at different strains (see FigS6a of Ref. [30]). The agreement with some experimental data is promising. However, besides this encouraging sign, we should admit that there are many experiments reported in literature that we can’t yet explain.

7. Conclusion

The model presented in the paper is based on a displacive paradigm. Deformation twinning is considered as a martensitic transformation; the atoms move collectively by a homogenous lattice distortion. The work presented here differs, however, from the usual “shear” theory. As in our previous crystallographic studies on martensitic transformations [21][22] that showed that the shear matrices should be replaced by angular-distortive matrices to evaluate the lattice distortion during the transformation, a similar approach was applied to treat the case of the $\{10\bar{1}2\}$ extension twins in magnesium. Instead of determining the simple shear matrices that restore the lattice and then guessing the trajectories (shuffles) of the atoms that are not at the lattice nodes, we first calculated the atom trajectories assuming that they are hard spheres of constant size, and then we calculated the analytical expression of the lattice distortion. The advantage is that it is sure that the atoms do not interpenetrate each other, i.e. the energy barrier remains reasonable along the transformation path. During the distortion, the unit volume increases up to 3% and the length of the “diagonal” direction in the twinning plane (direction OV) increases up to 1.4%, both before coming back to their initial values when the transformation is complete. These values are too high to be accommodated fully elastically by relaxing the hard-sphere assumption. The $\{10\bar{1}2\}$ twinning plane is not fully invariant during the distortion; it is just untilted and fully restored when the distortion is finished.

The calculations also show that the stretch distortion deduced by Liu et al.[33] from their observations of (90°, a) twinning domains in a submicron-sized samples, is a component of the distortion matrix associated with the usual (86°, a) extension twins in bulk samples. The atom trajectories are very similar in the two cases and differ only by a continuous rotation angle varying between 0 and 3.4°. It can be concluded that the Liu et al.’s
observations are in agreement with the displacive nature of mechanical twinning, and there is reason to consider them as a sign for a reconstructive mechanism. An analogy with ferroelectrics makes us think that the stretch distortion associated with the (90°, a) domains could be a “spontaneous” distortion that is free to occur in nano-objects, and that the usual distortion associated with the (86°, a) twins could be a “constrained” one that occurs in bulk samples. The 3.4° compensating angle can then be viewed as an obliquity angle. From the lattice distortion model and analogies with martensitic transformations, an orientation continuity is expected between the two low-misorientated (86°, a) twin variants. This expectation was confirmed by an EBSD map acquired on a magnesium single crystal.

As extension twinning differs from a shear mechanism in its details, a criterion that generalizes Schmid’s law was used. It is based on the interaction work between an external stress field and the deformation field generated by the twinning distortion. The “virtual work” criterion allows the prediction of extension twin formation for parent crystal orientations associated with negative Schmid factors. More precisely, the model predicts the formation of extension twins during uniaxial tensile tests with parent crystals tilted in the range [−59°, 59°] whereas the usual shear model predicts extension twins only in the range [−43°, 47°]. The fact that a critical tilt angle of 59° was already experimentally reported in the literature is very encouraging. The deviation of more than 12° in the predictions made by the two models proves that it should be experimentally possible to distinguish the correct one. The calculations also show that small obliquity angles can have a huge impact on the habit planes and are worth being taken into account into the predictions.

The model in the present state is macroscopic; it is not able to give details on how the distortion is microscopically accommodated. The arrays of dislocations induced by the distortion at the interface (disconnections) and in the surrounding matrix (disclinations) are not yet predicted. The exact nature of these dislocations, the way they are distributed, and their effect in the formation of basal/prismatic ledges at the interface will be the subject of future investigations; we have no doubt that disconnection theory will then play a key role in the calculations.

Note:
A second paper treating the case of the \{10\overline{1}1\} contraction twins with the same approach is available on Arxiv [55]. Other papers are also in preparation for other twinning modes in hcp metals.

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Fig. 1. Hexagonal lattice of the parent crystal with its associated orthonormal basis \((x, y, z)\), with \(x = a_p, y = a_p + 2b_p, \) and \(z = c_p\). Some positions of the Mg atoms are labeled in order to describe the atomic displacement during the extension twinning process. The angle \(\eta\) is used as the unique parameter of the lattice distortion.

Fig. 2. Schematic representation of the atomic displacements during extension twinning viewed on different planes of the parent crystal: a) on OXY = (0001)\(_p\), b) on OXZ = (01\(\overline{1}\)0)\(_p\), c) on OYZ = (2\(\overline{1}\)1\(\overline{0}\))\(_p\) planes. The parent crystal is in blue and its resulting extension twin is in salmon. All the atomic displacements are functions of a unique parameter, i.e. the angle \(\eta\) of rotation of the M atom around the OX axis. The vector \(OX = a\) remains invariant. (d) Schemes explaining the change of the distances OY and OZ: as the atom M moves far from plane (OXY), the distance OY decreases, and as the atom N moves toward the plane (OXZ), the distance OZ, initially OZ = c, increases to eventually become OZ' = a when the distortion is complete. The (0001)\(_p\) and (01\(\overline{1}\)0)\(_p\) planes are transformed into the (01\(\overline{1}\)0)\(_t\) and (0001)\(_t\) planes, respectively. During the distortion process, the hard-sphere packing assumption imposes that \(MO = MX = MY = MS = NO = NX = NT = NT = OX = \) atomic diameter.
Fig. 3. Change of volume ratio $V'/V$ during extension twinning, as function of the parameter $\kappa = \sin(\eta)$, varying from $\kappa_s = 0$ (start) to $\kappa_f = 1/3$ (finish).

Fig. 4. 3D view of the distortion of a $(86^\circ, a)$ twins with a hexagonal Bravais unit cell. (a) Initial hcp cell ($\eta = 0$), with the $(0001)_p$ plane horizontal and $(01\bar{1}0)_p$ plane vertical, (b) intermediate state ($\eta = 9^\circ$), and (c) final state ($\eta = 20^\circ$). The final state is a restored hcp structure, but with the basal and prismatic planes interchanged in comparison with (a). The displacements between the initial and final states are marked by the black arrows.

Fig. 5. 3D view of extension twinning with a crystal made with $4x4x4$ XYZ cells. (a) Initial hcp cell ($\eta = 0$), (b) intermediate state ($\eta = 9^\circ$), and (c) final state ($\eta = 20^\circ$). The basal and prismatic planes interchanged during the distortion. The $(0\bar{1}12)$ plane is marked by the black-lined section. In the case of the $(90^\circ, a)$ twins, the basal and prismatic planes are untilted, and the $(0\bar{1}12)$ plane is tilted by $3.4^\circ$. In the case of the $(86^\circ, a)$ twins, the $(0\bar{1}12)$ plane remain untilted, and the basal and prismatic planes are tilted by $3.4^\circ$ (case shown here). The difference between the two twinning modes is hardly perceptible. Another view is given in the graphical abstract.
Fig. 6. Change of the direction OV during twinning distortion. (a) Schematic view on the plane OYZ = (2\overline{1}10)\textsubscript{p} of the tilt \( \xi \) of OV around the a-axis. (b) Evolution of the ratio of distances \( OV'/OV \) proving that even if the tilt \( \xi \) is corrected, the (0\overline{1}12) plane cannot be maintained fully invariant.

Fig. 7. 2D representation of the orientational domains in a ferroelectric crystal created by a cubic→tetragonal phase transition. (a) Domains of “spontaneous” distortion with the polarization vectors (arrows) along the high-symmetry x and y-axes. (b) Domains experimentally observed, with domain walls on the (1,1) and (1,-1) planes. A small rotation of the spontaneous domains by an angle \( \delta \), called obliquity, is required to respect the compatibility conditions at the interfaces. There are 4 variants in case (a) and 8 in case (b).
Fig. 8. Concomitant formation of a pair of low-misoriented (86°, a) variants. (a) In analogy with ferroelectrics, the variants $t_1$ and $t_2$ are associated to an obliquity angle of $\xi = -3.4^\circ$ and $\xi = 3.4^\circ$, and named $D_z^-$ and $D_z^+$, respectively. Their habit plane is $(0\bar{1}12)$ and $(0\bar{1}12)$, respectively. (b) In analogy with previous works on martensitic transformations, it is expected that the back-stresses generated by the dislocations created by the twinning distortion in the surrounding matrix induce a continuous rotation between the two variants $D_z^-$ and $D_z^+$. Both have the tendency to be re-orientated back to the initial lattice orientation (before distortion), which corresponds to the intermediate (90°, a) variant $D_2$. A continuous rotation of angle $2\xi$ around the a-axis is expected to link the variants $D_z^-$ and $D_z^+$. 
Fig. 9. EBSD map in longitudinal cross-section of a magnesium single-crystal deformed at 100°C. (a) Orientation maps in Euler colors. The parent crystal is in green, the twins are in red and blue. (b) <001>, <100> and {012} pole figures. They show that the red and blue crystals are low-misoriented (86°, a) twin variants. The directions normal to the traces of habit planes and reported in the {012} pole figure show that the habit plane of the blue twins is (01\bar{1}2) and that of the red twin is (01\bar{1}2). The continuous orientation gradient expected from the model is confirmed by the circular red-blue arc around the common a-axis encircled in the <100> pole figure. The middle of the arc corresponds to the (90°, a) twin. (c) Orientation gradients in the extension twins in the range [0,10°], (d) orientation gradients in the surrounding parent matrix in the range [0,10°]. (e) Misorientation profile along the bold arrow marked in (c). (f) Rotation axes associated with the low-angle misorientations (rotation angle between 2° and 10°) plotted in the inverse pole figure.
Fig. 10. Interaction work (in MPa) during extension twinning in a tensile stress field oriented along the z-axis of a parent crystal tilted by an angle $\phi$ around the a-axis and rotated by an angle $\theta$ around the n-axis (normal to the twinning plane). a) Interaction work $W_f$ calculated with the complete distortion (shear) matrix. $W_f$ is proportional to the usual Schmid factor. b) Interaction work $W_i$ calculated with the intermediate distortion matrix corresponding to the maximum volume change. c) Graph showing in yellow the orientations ($\phi, \theta$) where the condition $W_i > 0$ & $W_f < 0$ is true, i.e. where the criterion based on $W_i$ could explain the twin formation despite negative Schmid factors (“anomalous” twins). The axes in the graphs are in radians. d) Schematic view of the orientation of the parent crystal. The segments delineating the domains $W_i > 0$ & $W_f < 0$ for $\theta = 0$ are marked by the grey crosses in a) and b), respectively.