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The five twin laws of gypsum (CaSO$_4$·2H$_2$O).
A theoretical comparison of the interfaces of the contact twins.

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Abstract

The relaxed interfaces of the 100, 101, 001, 201 and 101 contact twins of the gypsum crystal are theoretically examined and their perturbed structure is described. The obtained twin energies ($\gamma_{PT}$) are: 13.6, 145, 255, 826 and 848 erg cm$^{-2}$ for the 100, 001, 101, 201 and 101 laws, respectively. Hence, the five twin laws can be divided in three classes reflecting the deep difference due to the character of the face, in the sense of Hartman-Perdok, on which the contact twin is supposed to form. In fact, the Original Composition Planes of the 100, 001 and 101 laws correspond to stepped (S) faces, while those of the 201 and 101 laws correspond to kinked (K) faces. Furthermore, the highest probability of occurrence of the 100 contact twins is strictly related not only to the lowest $\gamma_{PT}$ value, but also to the fact that, among the three S faces, (100) is the only one belonging to the athermal equilibrium shape of the crystal.

1. Introduction

In recent papers the theoretical and athermal ($T = 0$K) equilibrium morphology of gypsum was evaluated¹ and a theoretical analysis was performed on the development of the stepped prisms belonging to the main [001] zone. The conclusions were strengthened by the agreement between the predicted equilibrium morphology of the [001] zone and that of the giant crystals of Naica mine,² which grew very close to the thermodynamic equilibrium.³ Recently, the theoretical structure of the 100 contact twin was investigated and the athermal twinning and adhesion energies were determined by minimizing the twin energy.⁴ Here we report on the continuation of this systematic study on the theoretical equilibrium and growth morphology of gypsum. Indeed a deeper knowledge of twinning (either contact or penetration) is required. Five laws⁵ describe gypsum twins but, as far as we know, no studies dealt with the structures of the interfaces and very few on the energies involved in their generation. As a matter of fact, the “geometry of the twins” awakened great interest since the times of Romé de l’Isle and Haüy, and was described and set systematically in the papers by Mallard⁶ and Friedel,⁷ and the
genetic mechanisms of the “growth twins” has been thoroughly treated, for the first time, as late as at the end of the Fifties, by Curien and Kern. In this paper, starting from these works, we will attempt at determining a theoretical hierarchy among the five twin laws of gypsum, confining our attention to the contact twins and we will compare the calculated twin energy with their probability of occurrence obtained from the literature data about natural and artificial crystals. In a forthcoming paper we will deal with the homologous penetration twins, assuming the 010 plane as the original composition plane (OCP), in order to compare, for each twin law, the probability of nucleation of a given twin, either by contact or by penetration.

The reader is referred to the well documented papers by Cody and Cody on the observed occurrence frequency of gypsum twins, where over six decades of gypsum literature are summarized. It is a merit of this paper to give reasons to be critic on the proposed identifications of the most frequently occurring twin laws.

As throughout more than two centuries of investigations on the morphology of gypsum twins a variety of reference frames were adopted, we put at reader’s disposal the relationships among them, in the Appendix. This is certainly not due to an excess of zeal but simply to the fact that people working on crystal morphology and those determining crystal structure often find difficulty in communicating each other.

2. The choice of the reference frame

We adopted in this paper the reference frame chosen by De Jong and Bouman, Cole and Lancucki, and Hejinen and Hartman (see Table 1). Our choice was grounded on two main reasons:

- this frame uses the smallest and the second smallest vectors of the four possible lattice vectors;
- the \( \bar{z} \) axis coincides with the morphological elongation of the crystals growing from pure aqueous solution and of the major part of natural crystals.

<table>
<thead>
<tr>
<th>( a_0 )</th>
<th>( b_0 )</th>
<th>( c_0 )</th>
<th>( \beta ) (°)</th>
<th>Space group</th>
<th>Authors</th>
</tr>
</thead>
</table>
| 5.63 | 15.15 | 6.23 | 113.83 | C2/c | De Jong and Bouman
| 5.67 | 15.201 | 6.274 | 113.91 | A2/a | Cole and Lancucki
| 5.678 | 15.213 | 6.286 | 114.08 | A2/a | Hejinen and Hartman

Table 1. Cell parameters (Å) and related space groups referred to the morphological cell by Friedel (0.372:1:0.412; \( \beta = 113.83^\circ \)).

3. Geometrical description of the five twin laws

In Table 2 the data concerning the geometrical identity of the five twin laws are collected. In the first row the indexing refers to the mirror planes that relate the reference frames of the parent (P) and twinned (T) crystals. With T we indicate the crystal generated from P by the twin operations. In the second one, the twinning operation is described by the \( A_2 \) axis lying in the twin plane and perpendicular to the [010] diad axis: this is an important structural feature, i.e. the row parallel to the \( A_2 \) axis is common to both P and T individuals. It is worth remembering that the operations described in the first and second row are equivalent. Actually, in gypsum, a \( h\beta\bar{m} \) mirror plane does not change the direction of the [010] diad axis, whilst any \( A_2 \) axis coincident with a [\( u\bar{b}w \)] direction,
changes the [010] direction in the [010] one; nevertheless, [010] and [010] directions are equivalent in gypsum crystal owing to the 010 glide symmetry plane.

The third row shows the areas of the 2D coincidence meshes at the P/T interfaces. The meshes are nearly rectangular, defined by the lattice vectors parallel to the twin axes and by [010], which is obviously common to all the gypsum twins.

Rows from 4 to 7 identify, for each twin law, the multiple 2D-cells that can be obtained from the superposition of the P and T lattices onto their common 010 plane. In other words, following Simon, we attempted to find if super-lattices exist, defined by lattice points that are common to both P and T individuals.

In the fourth row are shown the vectors defining the 2D-mesh on the common 010 plane of each twin:
- items i) and ii) refer to the quasi-perfect multiple cells (either monoclinic or orthorhombic), i.e. to those cells where the angular misfit between the lattices of P and T crystals are close to nil, as illustrated in Figs. 1a and b, and in the fifth row of Table 2.
- item iii) refers to the multiple cells (pseudo-cells) where the angular misfit between the lattices of P and T is significant.

In the sixth row are listed the areas of the multiple 2D-cells on the common 010 plane; finally, in the seventh row the corresponding multiplicities are given.

### Table 2. Geometrical description and growth aspect of the five twin laws of gypsum. Both areas of the common 2D-mesh and related obliquities were calculated on the base of the Cole and Lancucki cell.12

<table>
<thead>
<tr>
<th>(1) Twin plane</th>
<th>100 “Swallow tail” or “Butterfly”</th>
<th>010 “Montmartre” or “spear head” or “fer de lance”</th>
<th>001</th>
<th>201</th>
<th>101</th>
</tr>
</thead>
<tbody>
<tr>
<td>(2) Equivalent twin A2 axis</td>
<td>[001]</td>
<td>[101]</td>
<td>[100]</td>
<td>[102]</td>
<td>[101]</td>
</tr>
<tr>
<td>(3) Area of the common 2D-mesh (Å²) at the interface of the contact twin</td>
<td>[001]×[010] = 95.371</td>
<td>[010]×[010] = 99.303</td>
<td>[100]×[010] = 86.189</td>
<td>[102]×[010] = 174.599</td>
<td>[101]×[010] = 151.95</td>
</tr>
<tr>
<td>(4) Vectors of the 2D-mesh on the 010 common plane</td>
<td>i) – monoclinic [401]×[001]</td>
<td>i) – monoclinic [402]×[101]</td>
<td>ii) – orthorhombic [803]×[001]</td>
<td>ii) – orthorhombic [705]×[101]</td>
<td>iii) – pseudo-orthorhombic [301]×[001]</td>
</tr>
<tr>
<td>(5) 2D-mesh angular misfit (°)</td>
<td>1.18 (i); 0.60 (ii); 2.30 (iii)</td>
<td>0.176 (i); 0.089 (ii); 1.19 (iii)</td>
<td>2.92 (i); 2.92 (ii); 12.81 (iii)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(6) Area of the 2D-mesh (Å²) on the common 010 plane</td>
<td>(i) = 130.08</td>
<td>(i) = 195.05</td>
<td>(ii) = 260.16</td>
<td>(ii) = 390.10</td>
<td>(iii) = 97.56</td>
</tr>
<tr>
<td>(7) Index of the twin</td>
<td>4 (i); 8 (ii); 3 (iii)</td>
<td>6 (i); 12 (ii); 5 (iii)</td>
<td>2 (i); 2 (ii); 2 (iii)</td>
<td>2 (i); 2 (ii); 2 (iii)</td>
<td>2 (i); 2 (ii); 2 (iii)</td>
</tr>
</tbody>
</table>

Table 2. Geometrical description and growth aspect of the five twin laws of gypsum. Both areas of the common 2D-mesh and related obliquities were calculated on the base of the Cole and Lancucki cell.12
Some preliminary conclusions can be drawn from the geometrical data reported in Table 2. From the third row, one can see that three out of the five laws (100, 1̅01 and 001) imply 2D coincidence mesh areas very close each other (mean value 93.62 Å²), while the areas associated to the two remaining laws (20̅1 and 101) are respectively 1.86 and 1.62 times larger than the mean value. This geometric feature is not a chance, but it has a counterpart in term of twin energy, as we will see later on.

Fourth and fifth rows indicate that quasi-perfect 2D-supercells cannot be found (on the common 010 plane) in the case of 001, 20̅1 and 101 twin laws. This is an indication that embryos twinned following the 100 and 1̅01 twin laws can nucleate on the {010} faces, OCP of penetration twins, while in the case of 001, 20̅1 and 101 twin laws, the nucleation on the {010} faces is contrasted by a supplementary activation energy barrier, owing to the elastic strain due to the angular misfit between P and T lattices.

Summarizing the geometrical constraints, we can reasonably divide the five twin laws as follows:

- contact and penetration twins following both 100 and 1̅01 laws, can likely occur; furthermore, the 1̅01 penetration twins seem to be favored in respect to the 100 ones. Obviously one cannot predict the frequency on the sole geometric considerations;
- contact twins described by 20̅1 and 101 laws seem to be highly disadvantaged when compared to 100 and 1̅01;
- the behavior of the 001 law twins is midway between the just mentioned groups.

The considerations just drawn from Table 2, originate from the correlation between the probability of formation of an epitaxy and the multiplicity of the common 2D-cell at the epitaxial interface i.e.: the lower are the 2D-cell area and the related parametric and angular misfit, the higher the probability of epitaxy to occur. As twinning can be considered a special epitaxy between two crystals of the same species, we highlighted the relationship between the lattice twin geometry, the interfacial twin energy and the occurrence frequency of the corresponding twin law.
Figure 1. The geometry of the contact twins of gypsum viewed along the [010] direction. Twin laws: 100 (a); 101 (b); 001 (c); 201 (d); 101 (e). For each twin law the contact twins have been drawn starting from the assumption that the growth rate of both \{120\} and \{\bar{1}11\} prisms is lower than that of the \{010\} pinacoid. Contact planes are represented by dashed lines. P, T stay for parent and twinned individuals, respectively. Next to each twin morphology, the corresponding 2D-coincidence lattice on the common 010 plane, is represented. The common 2D-mesh of the coincidence lattice is also drawn. It is worth nothing that $\delta_{100} = \epsilon_{101} = 105.02^\circ$.

In the Supporting Information we reported the observed occurrence of twins, classified according to the five twin laws. Here we confine our attention to focus on some structural features of the twin planes. It is interesting to remark that all the twin planes are of the $h0l$ type i.e.: they belong to the [010] crystal zone which is not an important one from the morphology point of view. As a matter of fact:
- the [010] direction does not correspond to a periodic bond chain (PBC hereinafter), in the sense of Hartman-Perdok;\(^{17}\)
- none of the twin planes corresponds to a flat (F) face;
- the \{100\} form, that could be an original composition plane (OCP) for the contact 100 twin, in the sense of Curien and Kern,\(^8\) has been proven to belong to the athermal equilibrium shape (ES) of the crystal, even if it is a S face;\(^1\)
- both \{\bar{1}01\} and \{001\} forms represent other potential OCPs, for \{\bar{1}01\} and 001 contact twins, respectively. As we will show later on, they can belong to the ES of the crystal, even if they are
stepped forms and their surface energies ($\gamma_{101} = 1010$ and $\gamma_{001} = 825$ erg cm$^{-2}$) are much larger than that of the $\{100\}$ form ($\gamma_{100} = 616$ erg cm$^{-2}$);

- the equivalent twin axes, respectively $[001]$ and $[101]$, (i.e., the common rows of the two individuals) are PBCs of the crystals, $[001]$ being the most important one. This means that, in principle, twinned rows can form on these faces by 1D-nucleation and develop $100$ and $\bar{1}01$ twinned embryos. We note that $100$ and $\bar{1}01$ are the most frequently occurring twin laws.

4. The surfaces corresponding to the five twin laws of gypsum

When building a twinned crystal we must comply with the symmetry, stoichiometric and local neutrality constraints. We start from the point of view that atomic deposition during growth progressively fills layers $d_{nh\_nk\_nl}$ thick. The supersaturation of the mother phase plays an important role in this process determining the extent of lateral interactions between particle of the partially filled layer. We may expect that when the crystallization affinity is high such extended bond network can be frozen in a state not corresponding to an absolute minimum of the surface energy but nevertheless being locally neutral, stoichiometric and satisfying the symmetry we just observe in a twin. These configurations could be also accessed by a mechanical strain experienced by the crystal.

In kinetic terms, we can say that if a face (potentially OCP) can exists during growth at high crystallization affinity, that is in a configuration having energy higher than the stable surface face, during the life time of this state a 2D twinned nucleus may stick on the face and freeze the instantaneous configuration. Therefore a necessary, although not sufficient, condition fulfilled by an OCP is that the corresponding stable face has a surface energy not too high.

In the Supporting Information a detailed analysis of the character of the surfaces of gypsum is performed. Here, we only report the classification of the surfaces according to Hartman-Perdok: the $(100)$, $(10\bar{1})$ and $(001)$ faces have a S (stepped) character, whereas the $(20\bar{1})$ and $(101)$ faces have a K (kinked) character.

5. The structures of the contact twin interfaces and their calculated twin energies

In the following we describe how the contact twins are generated and then we characterize them. At first, a crystal slab limited by the composition plane $h0l$ is reflected about a mirror plane parallel to $h0l$. In order to set an initial configuration of the interface between the two crystals, the original and the reflected slabs are moved one relatively to the other; ions and molecules can also be displaced to avoid evident repulsions. In this way several initial configurations can be tested. The energy of the bi-crystal is then minimized under the constraint of constant volume. Two dimensional periodic boundary conditions are imposed repeating, in direction parallel to $h0l$, the smallest 2D cell common to both crystals. We did not define supercells because the parametric coincidence of the 2D lattices and the small translations required to obtain the structural continuity of the PBC network through the interface between one crystal and its reflected image. By the way, to afford calculations based on supercells was too heavy.

We used the program GDIS$^{18}$ to build the slabs and the program GULP$^{19}$ for the energy calculations. As in our previous works on gypsum 100 contact twin,$^{4}$ we used the force field proposed by Adam.$^{20}$ At variance with our previous works, we needed a lower tolerance on the values of the energy ($ftol < 10^{-6}$) and gradient ($gtol < 10^{-5}$) for optimization; this allowed to obtain a mirror symmetry of the mean displacements of the plane of atoms at the interface between the P and T crystals, but only in the case of the twins having lower energy.
The energy of the interface is the difference of energy, per unit surface area ($S$), between the twinned crystal ($E_{PT}$) and the not twinned one ($E_{untwinned}$) comprising the same number of atoms and exhibiting the same outmost layers. Then the twin energy is an interfacial energy ($\gamma_{PT}$):

$$\gamma_{PT} = \frac{E_{PT} - E_{untwinned}}{S} \quad (1)$$

5.1. The 100 contact twin

This contact twin has been extensively treated in a recently published investigation. Nevertheless, its relaxed interface is illustrated is Fig. 2, for sake of comparison with the other four twins. Here, we would like just recollect that the transition from P to T individuals, is quite soft.

![Figure 2](image)

**Figure 2.** The contact 100 twin of gypsum after relaxation and viewed along the [010] direction. The continuity of the structure is clearly shown by the arrangement of the $SO_4$ tetrahedra across the twin interface, while the water molecules are strongly rearranged at the interface level. Grey color outlines the sequences building the PBCs, while dashed lines represent their continuation in the twinned individual (see also Fig. 3).

Indeed, the [100] PBCs running in the P crystal crossing the contact interface, deviate by $\sim 14^\circ$ and “continue” in the [101] PBCs of the T individual. In a similar way, the [101] PBCs of the P crystal cross the contact interface and “continue” in the [100] PBCs of the T individual, experiencing the same angular deflection (Fig. 2). It is worth noting that the structural order of the strong Ca–$SO_4$ bonds is slightly modified by the transition occurring between the two kinds of PBC. The symmetry relating the water molecules is only perturbed in the interface layer. Therefore, the equilibrium structure of the 100 contact twin is fairly similar to that of the normal crystal, since only hydrogen

*In relation (1) of ref. 4, ($E_b - E_{bw}$) must be changed in ($E_{bw} - E_b$).
bonds are quite distorted at the twin interface. Reasonably, this is the main reasons why the 100 contact twin energy ($\gamma_{P,T}^{00}$) acquires the very low value of 13.6 erg cm$^{-2}$.

5.2. The 001 contact twin

In Fig. 3, the [010] projection of the 001 contact twin is shown. The difference in respect to the 100 interface is striking, owing to the sharp discontinuity experienced by the PBCs across the twin interface:

i) the [001] PBC of the P individual, which is made by the stacking of iso-oriented SO$_4$ tetrahedra, continues in the [101] PBC of the T individual, where the alternating tetrahedra are mutually related by an $\approx 90^\circ$ rotation around the [010] axis. The reciprocal situation occurs for the [101] PBC of the P individual.

ii) The location of the water molecules within the $d_{200}$ slice (parallel to both [001]$_P$ and [010]$_{P,T}$ directions) is markedly different with respect to those belonging to the $d_{202}$ slice (parallel to both [101]$_P$ and [010]$_{P,T}$ directions).

Further, there is an evident singular $d_{002}$ layer, parallel to the twin interface, where the orientation of the groups SO$_4$ is intermediate between that in P and T individuals. In the bulk, the oxygen atoms linked to sulfur are arranged in four planes parallel to 001, while in the singular layer, marking the transition between the two crystals, the oxygen atoms are arranged on three planes: indeed two oxygen lay on the same plane marked by a dashed line in Fig. 3.

![Figure 3](image)

**Figure 3.** [010] projection of the 001 contact twin. A sharp discontinuity stands out in the orientation of tetrahedra within the ---SO$_4$ – Water – SO$_4$--- sequence, when the [001] and [101] PBCs belonging to P individual, cross the twin interface and continue in the T individual, as [101] and [001] PBCs, respectively. The deviation angle across the interface does not reach $5^\circ$.

In the singular layer the distance between oxygen atoms in closer contact is 6.80 Å while it is 4.46 Å in the two adjacent layers. In the bulk, the corresponding distances increase to 6.94 Å and 4.53 Å, respectively. As shown in Fig. S6, the variations of the longer O-O contacts occur over 8 tetrahedral layers, indicating damped librations of the groups SO$_4$. 
The librations are accompanied by oscillations of the distances between adjacent $d_{002}$ atomic layers in direction perpendicular to the 001 interface, as shown in Fig. S7. From the figures one sees that the layers of the water oxygens ($O_W$) move in phase with the calcium planes, while the sulfur planes are in antiphase with calcium. It is interesting to compare the amplitude of the interfacial deformation of the 100 twin with that of the 001 twin. Expressing the highest values of deformation as a percentage of the equidistance in the bulk we obtain the following values:

(i) 001 twin: $O_W$, +3.1% and -1.6%; Ca +2.5% and -1.2%;
(ii) 100 twin: $O_W$, +0.98% and -0.3%, Ca +0.43% and 0%.

These amplitudes are, in some cases, even higher than those of the layers at the surface of the slab (e.g. Fig. S7a), at variance with our finding in the case of the twin 100. We observe only very small translations and oscillations of atoms in directions laying in the OCP.

The 001 contact twin energy ($\gamma_{001}^{001} = 145 \text{ erg cm}^{-2}$) is an order of magnitude higher than that of the 100 twin.

5.3. The 10̅ contact twin

The [010] projection of the 10̅ contact twin is represented in Fig. 4a. The discontinuity encountered by the PBCs [1̅00] and [001] across the twin interface is strictly analogous to that just described for the 001 contact twin, as follows from the comparison between Fig. 3 and 4a. At variance with the 001 contact twin, the angle of deviation of the PBCs across the interface does reach ~8.9°. This clearly suggests that the transition from P to T individual should be even more expensive than that of 001 contact twin law.

Moreover, there is also in this case a singular $d_{203}$ layer, parallel to the twin interface, where the sulfate tetrahedra rotate to a position intermediate between those in the P and T crystals. The coordination of the two water molecules in the singular layer changes as well: one molecule follows the repeat sequence in the P crystal while the other the sequence of water in the T crystal. Figs. 4b and c show the non symmetrical displacements of Calcium and Sulfur atoms close to the 101 twin interface.

Typically, the amplitude of the rotations of the $SO_4$ groups and $H_2O$ molecules and the atomic translations at the interface between the two individual are high and the transition from the P to the T structure involves the movements of all particles over several layers. Summing up, it is not surprising that the twin energy associated to the number of deeply perturbed layers on the two sides of the twin interface increases, in this case, to the value of 255 erg cm$^{-2}$. 
Figure 4. (a) [010] projection of the 101 contact twin. The PBCs [100]ₚ and [001]ₚ cross the interface of the contact twin encountering a structural discontinuity analogous to that of the 001 twin law. The deviation angle across the interface increases, in this case, to ~8.9°. The mean distances between the Ca (b) and sulfate (c) layers are not symmetricaly distributed around the 101 twinned interface.

5.4. The 201 and the 101 contact twins

The last two contact twins will be treated together owing to the common features of their twin interfaces. No trace remains, indeed, of the local symmetry that characterizes the 100 and 001 interfaces and, to a lesser extent, the 101 contact twin. This can be clearly seen in Figs. 5a and 5b, where the interface layers are expanded and organized in singular 2D structures different from the layers in the bulk.

Figs. 5a and 5b show the structure of the 201 and 101 contact twins. One can notice the expansion of the interfacial regions in direction normal to the twin composition plane and the complex correlated rotations of the SO₄ tetrahedra and water molecules. All atoms experience translations parallel and perpendicular to the ideal 201 plane and, consequently, the structural disorder extends, in the 201 twin, over five layers of thickness ½ d_{201}. In the 101 twin, the SO₄ tetrahedra rotate to such an extent that a layer is formed where a plane of oxygen atoms is sub-parallel to 101. In this case high angle rotations of the tetrahedra and of the water molecules occur over seven layers of thickness ½ d_{101}.

The number of PBCs crossing the interface is the same for both twins (i.e., PBCs [100], [001] and [101]), but the angular deflection experienced in the transition from the P to the T crystal, is clearly larger in the 101 twin. In particular, the structural disorder and the lack of local symmetry at the interfacial level have a dramatic effect on the twin energy that practically increases of an order of magnitude with respect to γ_{PT}^{100}. As a matter of fact, γ_{PT}^{201} = 826 and γ_{PT}^{101} = 848 erg cm⁻², respectively.
6. Conclusions

The essential features of the five contact twins of gypsum are summarized in Table 3. For each geometrical law, the character of the original composition plane and the twin energy are indicated.

<table>
<thead>
<tr>
<th>Twin law</th>
<th>Character of the OCP</th>
<th>$\gamma_{PT}$ (erg cm$^{-2}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>S</td>
<td>13.6</td>
</tr>
<tr>
<td>001</td>
<td>S</td>
<td>145</td>
</tr>
<tr>
<td>101</td>
<td>S</td>
<td>255</td>
</tr>
<tr>
<td>201</td>
<td>K</td>
<td>826</td>
</tr>
<tr>
<td>010</td>
<td>K</td>
<td>848</td>
</tr>
</tbody>
</table>

Table 3. Features of the five twin laws of gypsum. Twin energy values ($\gamma_{PT}$) reflect the deep difference due to the character of the face on which the contact twin is supposed to form.

The five twins can be grouped in three classes, according to the magnitude of the twin energy: the 100 law, the couple 001 and 101 and, finally, the couple made by the 201 and 101 laws. Some comments further justify this classification.

a) The 100 law can be assumed as the touchstone for the other laws. The 100 surface shares the S character with the 001 and 101, but its twin energy is an order of magnitude lower. As the nucleation frequency of twins (either 2D or 3D) exponentially depends on the corresponding activation energy (which, in turn, is linearly related to the twin energy), the probability of formation of both 001 and 101 contact twins should be by far lower than that of the 100. Many factors contribute to determine the peculiarity of the 100 contact law:
i) among the \{100\}, \{001\} and \{10\overline{1}\} S forms the \{100\}, is the only one belonging to the [010] zone, that enter the athermal equilibrium shape of the crystal, owing to the low value of its specific surface energy;

ii) the 100 twin interface, among the five investigated, is the only one that is crossed by the [101] and [001] PBCs without change of the PBC structure;

iii) as a consequence, the \{100\} surfaces are the only ones where a 2D nucleation can occur both for normally oriented nuclei and for twinned ones.

b) The investigation on the 001 law deserves a novel feature: in spite of the usual tendency to neglect the 001 plane as a possible OCP, the formation of 001 contact twins seems to be favored with respect to the 10\overline{1} ones because the ratio $\gamma_{TT}^{001} / \gamma_{TT}^{100}$ is about 1.76. This should not be surprising as the PBCs crossing both the interfaces undergo the same structural change, but the angle of deviation at the 10\overline{1} interface is close to twice the value measured in the 001 twin. Such difference reflects the slight asymmetry characterizing the 10\overline{1} contact interface when compared to the highly symmetrical 001 one (see Figs. S7, 4b and 4c).

A second considerable finding is the difference $(\gamma_{TT}^{001} - \gamma_{TT}^{100}) \approx 131$ erg cm$^{-2}$. This large value establishes, on a theoretical base, a hierarchy between two contact twin laws and it accounts for the experimental results obtained by Kern and Rehn$^{21}$ who observed that the occurrence frequency of the 100 contact and penetration twins is more or less the same in growth from pure aqueous solution, while 10\overline{1} contact twins were never observed, under the same temperature and supersaturation conditions.

c) The high twin energy of the remaining laws, i.e. the $\overline{2}01$ and 101, can be understood referring to the Hartman’s model of the growth twins.$^{22}$ As mentioned above, the $\overline{2}01$ and 101 interfaces differ from all the preceding ones by the degree of structural disorder and by the sharp discontinuity of the structures of the PBCs when crossing the twin interfaces. Moreover, the $\overline{2}01$ and 101 twin energy increases with the angles of misalignment of the respective PBCs. This behaviour is a consequence of the K character of the \{201\} and \{101\} forms. In other words, a kinked (K) face, which is the best example of a crystal surface instability, does not have surface sites for the formation of 2D islands or 1D steps (either normally oriented or twinned). Then the conditions necessary to the twin formation cannot be fulfilled by a K face, unless the adsorption of foreign substances determines either a K$\rightarrow$S or a K$\rightarrow$F transitions.

Consequently, only $\overline{2}01$ and 101 penetration twins should be observed in growth from pure aqueous solution, since the water adsorption is not sufficient to reduce the surface energy by such an amount to allow the formation of contact twin possible. As a matter of fact, the adhesion energy ($\beta_{adh}^{solution}$) due to the solvent adsorption cannot exceed the value of $2\gamma_{water} \approx 150$ erg cm$^{-2}$, as it follows from Young’s and Dupré’s formulae.

Summing up, in this paper we analyzed, in the light of the Hartman-Perdok theory, the twin geometry and the surface properties of the original composition planes and found that the structural features well correspond to the relative stability of the contact twins obtained from their energy.

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**Supporting Information Available.** The occurrence frequency of the five twin laws of gypsum; the character (in the sense of Hartman-Perdok) of the different surface profiles of the (100), (101),
(001), (01̅1) and (101) faces of gypsum; the variations of the O-O distances and layer thickness as a function of the number of layers in the (001) twinned slab. This material is available free of charge via the Internet at http://pubs.acs.org.

References


Appendix

The first study on the growth twins of gypsum is due to the Kern’s School. A first experimental paper, dealing with the gypsum growth from pure aqueous solutions is by Kern and Rehn.\textsuperscript{21} The interpretation of the results on the growth twins was integrated by Simon\textsuperscript{15} who analyzed the mechanisms of formation of both contact and penetration twins. In these works, a reference was made to the morphological frame of Des Cloizeaux\textsuperscript{23} that was used in two different versions: i) Kern and Rehn\textsuperscript{21} adopted the original frame by Des Cloizeaux\textsuperscript{23} which reads:

\[
a:b:c = 0.68994 : 1 : 0.41241; \beta = 80^\circ 42'
\]

according to Dana,\textsuperscript{24} and indexed the two main twin laws of gypsum as (100) and (\textbar{}01).

ii) Simon,\textsuperscript{15} reversing the \(z\) axis, used another frame:

\[
a:b:c = 0.6910:1:0.4145; \beta = 98^\circ 52'
\]

and indexed the main twin laws of gypsum as (100) and (101), respectively.

These two choices were synthesized by Simon\textsuperscript{15} adopting the frame by De Jong and Bouman which is related, within the limits of the experimental errors, to the frame by Gossner:\textsuperscript{25}

\[
a_0 (\text{Å}) = 10.47 ; b_0 (\text{Å}) = 15.15; c_0 (\text{Å}) = 6.28; \beta = 98^\circ 52'
\]

which is, in turn, coherent with the item ii).

Hence the relations between the indexes of the faces in the two reference frames: Des Cloizeaux (DC) and De Jong-Bouman (DJB) depend on the relations between the vectors:

\[
(\tau_1)'_{DC} = (-2 \tau_1 - \tau_3)_{DJB}; (\tau_2')_{DC} = (-\tau_2)_{DJB}; (\tau_3')_{DC} = (\tau_3)_{DJB}
\]

\[
\{h'k'l'\} \text{ Des Cloizeaux} = \begin{pmatrix} -2 & 0 & -1 \\ 0 & -1 & 0 \\ 0 & 0 & 1 \end{pmatrix} \{hkl\} \text{ De Jong - Bouman}
\]

Another way of indexing the most frequently occurring twins was adopted by A.M. Cody and R.D. Cody,\textsuperscript{9,10,26,27} who referred to the morphological F cell of Dana.\textsuperscript{24} Care must be taken in working with the morphologies described in their papers, since some ambiguities arise from the face indexing.
The five twin laws of gypsum (CaSO$_4$·2H$_2$O).  
A theoretical comparison of the interfaces of the contact twins.

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**Synopsis**

The relaxed interfaces of the 100, $\bar{1}01$, 001, 20 $\bar{1}$ and 101 contact twins of the gypsum crystal are theoretically examined and their perturbed structure is described. The obtained twin energies are: 13.6, 145, 255, 826 and 848 erg cm$^{-2}$ for the 100, 001, $\bar{1}01$, 20 $\bar{1}$ and 101 laws, respectively. Hence, the five twin laws can be divided in three classes reflecting the deep difference due to the character of the face, in the sense of Hartman-Perdok, on which the contact twin is supposed to form.