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# ARTICLE

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# The crystalline state of rubrene materials: intermolecular recognition, isomorphism, polymorphism, and periodic bondchain analysis of morphologies

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A survey of all crystal structures of rubrene materials in the Cambridge Structural Database is presented. Although the chemical substitution landscape is wide, hydrogen bonding functionalities are absent. Recognition motifs frequently found in crystals are the "slipped-cofacial" molecular pairing, and herringbone or purely translational 3-D propagation. Packing modes are classified in terms of structure determinants, cohesive energies of pairs of molecules in closer contact computed by the CLP atom-atom potential field. In these terms, crystal isomorphism with different chemical substitution is quantitatively assessed. Polymorphs are relatively few, perhaps due to poor solubilities that hamper crystallization screenings. True polymorphs are also identified by structure determinant patterns, and a new polymorph of the di-p nitrophenyl derivative has been prepared and characterized by X-ray diffraction. Crystal morphologies of selected rubrenes have been predicted by Hartman's Periodic Bond Chain approach using PIXEL attachment energies; there is good agreement with experimental morphologies of crystals grown by sublimation. The good results obtained by CLP and PIXEL show promise for a computationally cheap access to lattice energies and morphology prediction. In general, from our overview it looks like sensitive spots in the driving forces for rubrene packing are the 4-substitution sites at the lateral rings, with substituents of moderate steric bulk. Peripheral substitution at the tetracene core seems to be less relevant. Our survey provides a structural background fostering new ideas on the synthesis and planning of physical properties of rubrenes.

### Introduction

Organic semiconductors have been for a long time under scrutiny for the development of organic electronic devices.1,2 Among a host of molecular candidates, rubrene (5,6,11,12-tetraphenyl-tetracene) has been at the forefront of research after discovery of the high charge-carrier mobility of its orthorhombic polymorph.3,4 Although organic semiconductors have not yet reached performance levels suitable for extensive marketing, the synthesis and the characterization of rubrene derivatives are still highly attractive.<sup>5</sup>

A primary issue is the design of high yield chemical synthetic paths, but a better understanding of the solid-state properties of these organic crystals is also desirable because physical properties depend on detail of crystal packing. Intermolecular interactions determine the crystal structure, influencing charge-carrier mobility, transfer integrals and exciton diffusion length.6 Bulk chargetransport properties are related to  $\pi$ -stacking interactions, arising from orbital overlap of adjacent molecules. In the nearest-neighbor stacking arrangement of the rubrene crystal the presence of lateral phenyls forces a slippage on the long molecular axis of the tetracene core, forming a hopping pair called "slipped-cofacial"<sup>7</sup> with a distance of 3.74 Å, larger than in typical  $\pi$ -stacks<sup>8,9</sup> and in tetracene itself.<sup>7</sup> Stacking geometries are also influenced by the twisting of the tetracene core imposed by intramolecular strain. The pair propagates in rows that are the main direction of conduction<sup>10</sup> with a high value of the transfer integrals. $11$  These rows often pack in a zig-zag fashion, giving the crystal the so-called "herringbone" shape (Fig. 1).

The preparation of effective semiconducting organic materials depends on i) finding molecules with improved solubility and stability against light and/or oxygen degradation<sup>12</sup> (lack of these features is a drawback of pristine rubrene); and ii) obtaining crystal structures with planar polyacene cores and optimum  $\pi$ - $\pi$  overlap. The task is, to say the least, awkward. In fact, previous analyses of crystal packing in rubrene derivatives (see reference 5 and references therein) focussed on  $\pi$ - $\pi$  stacking distances, slip distances and angles, tetracene planarity, herringbone angles, trying to find a rationale for enhancement of transport properties through chemical modification. However, a study based on graphical tools found little or no correlation between planarity of the tetracene core and the nature of lateral substituents.13

Modification of the rubrene molecule at the rim of the tetracene core frequently results in the disruption of the herringbone motif and/or significant twisting of the tetracene backbone. Partial or

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E-mail angelo.gavezzotti@unimi.it Electronic Supplementary Information (ESI) available: an Appendix on crystallographic terminology, more crystal packing and aggregation diagrams as mentioned in the text; numerical detail of the partitioned Coulombic-dispersion energies and comparison of force fields, energy crystal graphs for PBC modeling of crystal morphologies. Deposition number 2149694 contains single crystal X-ray data of the triclinic polymorph of 5,11-bis(4-nitrophenyl)-6,12-diphenyltetracene (provisionally called here CIYNAB01). These data are provided free of charge by the joint Cambridge Structural Data Centre and Fachinformationszentrum Karlsruhe Access Structure service www.ccdc.cam.ac.uk/structures. See DOI: 10.1039/x0xx00000x



pyridine rings. (b) Some key structural features in rubrene crystals:  $\pi$ -stacking is in the see Table 1).

only two cases but with loss of the favourable rubrene packing and meta positions of the attached phenyl rings, trying to preserve rubrene.16,17 These new derivatives allowed a characterization of oxygen/light stability and charge transport properties, providing new hints about the relation between molecular architectures, crystal structures and physical properties. With introduction of para functional groups on centrosymmetric rings 5 and 11<sup>17</sup> the unmodified tetracene core preserved the structural motif of the (200) monomolecular slice, although the stacking of layers was no longer feasible in the orthorhombic system due to the increased corrugation of the (h00) surface. The strong repulsion caused by<br>Refcode identifier). A preliminary screening to assess the adaptability crowded peripheral phenyl rings can be reduced by twisting of the The 1 la) The induced was the control in the control in the most control in the m stable for molecules in solution, vapour phase and amorphous profile migs: (8) Some key structural rearries in the mean of the desired electronic material to the proposition of the solid structure is see Table 1). Solid the mean tracene or each structure of this see Table 1). Solid properties; unfortunately, this is observed in about half of the er Table 1).<br>
strachabre, the crystalling mathematic provides a planar tetracene core in discultimenty, running along specific crystalligraphic [wwv)<br>
complete terrace fluorities of the favourable rubre packing discursions intermolecular steric factors. out an partial terative care to exact met (200) suce to the case and the polymorm polymorphic stability and the mobility in t oxyger/inglin salarite that that has a different properties, propagation properties and physical properties. With introduction of *para*<br>
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structures and p

Polymorphism is also an issue. Pristine rubrene has three polymorphs (orthorhombic, monoclinic and triclinic) obtained under<br>the MiCMoS ambient conditions by crystallization from different solvents or by vacuum techniques, plus a high-pressure triclinic structure stable displacement.<sup>23</sup> The monoclinic polymorph shows no  $\pi$ -stacking with

consequential connection between molecular structure and crystal packing, at least in its essential features if not in a complete crystal structure prediction. Such studies rely on a statistical approach based on crystal and molecular geometries as found in experimental diffraction work, supplemented by a quantitative analysis of packing energies and forces. Previous attempts on various small subsets of

**a planet tetracene fluorination and the second state term in the core in the second state of the sec** mode.<sup>14,15</sup> A more promising route involves substitution at the para units); ii) during crystal growth strong intermolecular bonds are **a)**<br> **b** textural sections with the core be a method, and the extracene core and the extracene core and the extracene core and the extrace in the equal tetrace is a method, and the equal tetrace is a method, and the equa present here a survey of molecular and crystal properties of 33 tetra(ring)-substituted tetracenes with rubrene-like architecture, fully characterized by single-crystal X-ray diffraction, along with a detailed analysis of their intermolecular energies and crystal packing modes. In a different perspective to the same aim, the equilibrium and growth morphologies of crystals of orthorhombic rubrene and four monoclinic derivatives<sup>17</sup> have been modeled. Crystal morphology depends mainly on intermolecular potentials that determine anisotropic growth forces, but other external, **F**(Chemistryins Page 18 of 67<br>Journal Name<br>present here a survey of molecular and crystal properties of 33<br>tetra(ring)-substituted tetracenes with rubrene-like architecture,<br>fully characterized by single-crystal X-ray dif supersaturation, temperature, pH, the influence of solvent, of impurities or additives, and the presence of electromagnetic or mechanical fields. The theoretical crystal morphology can be derived solely from the crystal structure by Periodic Bond Chain (PBC) methods,28-30 whereby a crystal structure is seen as a 3D array of uninterrupted chains of strong bonds with the crystal's stoichiometry, running along specific crystallographic [uvw] directions. The method works under the assumptions that i) crystal growth is controlled by surface processes (incorporation of growth formed between growth units; iii) external factors play only a minor role (growth performed at low/moderate supersaturations). Obviously, a reliable method for the quantitative evaluation of the energies of these bonds must be available. supersaturation, temperature, pH, the influence of solvent, of<br>mechanical fields. The theoretical crystal morphology can be derivend<br>solely from the crystal structure by Periodic Bond Chain (PBC)<br>methods,<sup>28-30</sup> whereby a

# Results and Discussion

### Structure screening: a normalized dataset of rubrenes

Proprium in a sole of insults (and in the signing in a sole of provided by standard procedures in a dioc modules of<br>analytical product in a sole of the module of the MiCMAS platform (see the documentation are<br>analytical above ca. 6 GPa.<sup>21</sup> The mobility in the orthornombic polymorph<sup>22</sup> Is group assignment, etc., were present. The presently determined<br>an order of magnitude greater than that of the triclinic polymorph, attuative of a nelim structures and popision in popision of the structure and point of the structure are energies. Win influention or power state in the structure in the stru signles to the mean the signles and and the dichical signles and and the signe and anomy by the components, phenomenology the signles are all 5,11,12-substitution only by ing componing properties; unfortunalized the base o Table S1 (ESI<sup>†</sup>) has a list of all the rubrene crystal structures in the Cambridge Structural Database (CSD, $31$  labeled by their six-letter of each structure to a systematic crystal packing analysis was carried out, as described below. The main features that guided the selection were a) 5,6,11,12-substitution only by ring compounds, phenyls, furans, pyridines, or thiophenes; b) available unit cell dimensions, space group and atomic coordinates for all non-hydrogen atoms; c) no unresolved disorder; d) R-factors below 7.5% (with one exception). Unreliable or absent hydrogen atom positions were corrected or provided by standard procedures in ad hoc modules of platform (see the documentation at sites.unimi.it/xtal\_chem\_group/). The MiCMoS module Crysaa was used to make sure that no undetected errors in atom positions, space Obviously, a reliable method for the quantitative evaluation of the<br>energies of these bonds must be available.<br>**Results and Discussion**<br>**Structure screening: a normalized dataset of rubrenes**<br>Table S1 (ESH) has a list of a structure of a polymorph of CIYNAB, provisionally labeled CIYNAB01, was also considered. Two resolved twins (RAGDEL, VICHAT) were included, and two partially disordered structures (RAGDAH, PIXPUJ01) were included using the major component of disorder. The polymorphs GORVIU, PIXPOD and PIXPUJ, although interesting, could not be included due to major disorder that prevents a unique analysis in our terms. The final dataset of 33 crystal structures (Table 1) can be used with confidence for further theoretical studies. A complete list of atomic coordinates including hydrogen atoms is available from the authors upon request.

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Table 1 Molecular and crystal data of rubrenes in the optimized Database. Table S1 (ESI+) has other detail and cell dimensions



[a] Identification code of the Cambridge Structural Database (CSD). [b] Molecules in unit cell. [c] Molecules in asymmetric unit. [d] Intramolecular symmetry: I inversion, A twofold axis, M mirror. [e] C-C-C-C cis torsion angle across the central C=C bond in tetracene (trans is 180-cis). [f] Crystallographic R-factor. [g] Ph=phenyl, Me=methyl, t-Bu=tert-butyl, CF<sub>3</sub>=trifluoromethyl. In PhX codes, the X substituent is in the para position unless otherwise stated. [h] Substitution at the tetracene ring. [i] Major component of disorder. [j] Resolved twin

### Force field assessment

The choice of a suitable force field is imperative in the present case and must be made with a careful maximization of the efficiency/cost ratio. The focus being on intermolecular contact, all molecules are kept in the geometry determined by X-rays so that no intramolecular force field is necessary.

Intramolecular energy terms can be neglected when discussing interaction energies between pairs of molecules, that are just tracers of the packing arrangement in the crystal. Intramolecular factors may be important for comparisons between total crystal energies; a procedure for an approximate evaluation will be described for the

case of perfluororubrene (see below). The choice of a suitable intermolecular force field must be made maximizing the efficiency/cost ratio. The MiCMoS environment offers three schemes for intermolecular potentials, listed in order of increasing accuracy: a) the AA-CLP formulation,<sup>32</sup> an atom-atom scheme of entirely empirical origin, that requires only a few seconds; b) the AA-LJC formulation,<sup>33</sup> also atom-atom but requiring atomic point-charge parameters derived from an MP2-MO wavefunction; the molecular orbital calculation is demanding for the big rubrenes, but lattice energy calculations take fractions of a second; c) the PIXEL scheme, 34 that requires an MP2-level electron density and also a considerable

amount of computing time for lattice calculations by finite integration of the various operators on the discrete wavefunction. The three methods work with default library parameters for organic repulsion terms (CLP and Pixel also include a polarization term). PI-New Journal of Chemistryins<br>
amount of computing time for lattice calculations by finite<br>
integration of the various operators on the discrete wavefunction.<br>
The three methods work with default library parameters for or energy trends and often quite similar absolute energy values. Therefore, in large scale structure comparisons the cheapest CLP method was applied, while the accurate Pixel method was used in the analysis of periodic bond chains for some representative **QQQCIG05** compounds. Table S3 (ESI<sup>+</sup>) collects the detail of partitioned lattice energies for all crystals in Table 1.

### Crystallography of rubrenes: General aspects

The sample is too small for statistics on space group frequency, but the obvious  $P2<sub>1</sub>/c$  is predominant. Depending on molecular constitution, rubrene molecules can have internal inversion, mirror or twofold axis symmetry preserved in the crystal, with 1/4, 1/2, 1, 3/2 or 2 molecules in the asymmetric unit. Although



stabilization energy is  $> 10$  kJ mol<sup>-1</sup>. (b) Energy/distance plot for the independent pairs with labeling of the symmetry operations,  $O_{mm}$ . The red rectangle encloses the short



Fig. 3 The slipped-cofacial arrangement: top, in two polymorphs of the parent compound; below: POGZIV shows a twisted tetracene core. Oxygen red, nitrogen blue, sulfur yellow.

are normal (0.68 to 0.74), and nearly all crystal structures show a 10-12-first-neighbor coordination sphere resembling a close packing of spheroids. There are no short atom-atom distances below the sum of contact radii, leaving no ground for the introduction of intermolecular "bonds". Rubrene lattice energies are dominated by dispersive factors (Table S3, ESI+), because carbon and hydrogen atoms account for 91% of the total in the database, fluorine contributing another 7%, so that no "polar" groups are present. There are no hydrogen bond donors in our database, preventing a test of competition between dispersion and hydrogen bonding.

### Structure determinants and pairing symmetries

Intermolecular structure can be conveniently analyzed by partitioning the total lattice sums into interaction energies between pairs of molecules ( $E_{mm}$ ). While the lattice energy is comparable to sublimation heats, the  $E_{mm}$ 's are purely computational quantities, having no experimental counterpart. The  $E_{mm}$ , the corresponding distance  $R_{mm}$  between molecular centers of coordinates, and the symmetry operator acting between the two partners,  $O_{mm}$ , form a triad called a structure determinant. These determinants are of great help in a quantitative description and comparison of packing modes. In a way, they are zero-dimensional precursors of the periodic bond chains. low the sum of contact radii, leaving no ground for the regretice condition of intermolecular "bonds". Rubrene lattice regries are dominated by dispersive factors (Table S3, ESI†), cause carbon and hydrogen atoms account because carbon and hydrogen atoms account for 91% of the database, fluorine contributing another 7%, so that too "polar" groups are present. There are no hydrogen bond donors in our database, preventing a test of competit between dispersion and hydrogen bonding.<br> **Structure determinants and pairing symmetries**<br>
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the 7.0-7.25 and 7.75-8.0 Å bins. In fact, 21 out of 33 crystals show  $R_{mm}$ 's of 7-8 Å either by translation (cell axis length) or by some other of structure determinants. The plot has an obvious bias to higher  $E_{mm}$ for shorter  $R_{mm}$ , meeting a lower limit at 7 Å. With few exceptions, translation and inversion are the only symmetries that allow a pairing



Fig. 4 Examples of coupling modes between pairs in the asymmetric unit. Left: carbon atoms of one molecule magenta; right: fluorine atoms blue in one molecule, green in the other  $(R_{mm}$  Å,  $E_{mm}$  kJ mol<sup>-1</sup>).

frequency between 9.5 and 11 Å, while the distribution levels at 10- 25 kJ mol<sup>-1</sup> with distances > 10 Å at second-neighbor stage, as recognition becomes less selective.

distance is called "slipped-cofacial", a parallel arrangement of the tetracenes allowing as much  $\pi$ -overlap as possible compatibly with extended gallery of such coupling patterns. This mode occurs with 7 There are the method of the station between the station of the method of the station between the station of the station of the method of the physical properties of rubrence and extended packing the stationary particle is



Fi**g. 5** P1 space groups. (a) Expansion of the SC7 motif by coordination into a fourfold **Fig. 6** (a) The SC7 + 4 Screw herringbo<br>Cage of translation related molecules (QQQCIG14). (b) Propagation into ribbons of (compare w cage of translation related molecules (QQQCIG14). (b) Propagation into ribbons of alternate inversion-related molecules (GORVUG).

This configuration is fostered almost exclusively by dispersion, although some charge-transfer contribution may be also present.

A definite symmetry relationship is not a prerequisite for short distance stabilization. The most stable pair in the database, -82 kJ f Chemistry ins<br>
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This configuration is fostered almost exclusively by dispersion,<br>
although some charge-transfer contribution may be also present.<br>
A definite symmetry relationship is not a prerequisite for short infrequent short distance coupling over a screw axis is MIVDOM (Fig. S4, ESI<sup>+</sup>) with an  $E_{mm}$  of -70 kJ mol<sup>-1</sup> between non-parallel tetracene units. In a few cases, 7 Å couplings with low stabilization result from f) Chemistry ns<br>
This configuration is fostered almost exclusively by dispersion,<br>
although some charge-transfer contribution may be also present.<br>
A definite symmetry relationship is not a prerequisite for short<br>
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### Coordination spheres and extended packing motifs

Example is a second common coupling mode at short intermolecular distance ("SC7" motif) is expanded by pure translation (Fig. 5a) or by<br>The most common coupling mode at short intermolecular distinguished intersion (Fig. 5b Since the reciprocal orientation of tetracene planes is so important for the physical properties of rubrenes, this feature is now taken as the leading one in the analysis of extended crystal packing. In the six  $\overline{P1}$  space groups, and in two C2/c, Z' = 1/2 space groups with the tetracene plane riding a twofold axis (see Table 1), any symmetry operation (translation, inversion and centering) can only produce a parallel alignment of tetracenes. The slipped-cofacial motif at 7 Å **EXECUTE:**<br>
This configuration is fostered almost exclusively by dispersion,<br>
although some charge-transfer contribution may be also present.<br>
A definite symmetric pair in MIVCUR (Fig. 4). An example of<br>
distance stabiliz **EXECTS**<br>
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A definite symmetry relationship is not a prerequisite for short<br>
A distance stabilization. The most stable pair in the database, -82 kl<br>
mol<sup>-1</sup>, is the asymmetric pair in MI

In other  $Z' = 1$  structures with screw and/or glide operations the 3D expansion produces a corrugated arrangement of tetracenes. A



19 20

 $21$ 22

23

classification of the structure expansion motifs includes the symmetry operators acting among nearest-neighbor molecules,<br>while a masseum of the composition is magnitude by the curels.  $\sim$  -10 while a measure of the corrugation is provided by the angles between vectors perpendicular to the average tetracene plane. Fig. S5 and Table S5 (ESI<sup>†</sup>) carry the detail of the classification. A first group includes 8 strictly similar structures with SC7 motif, an group includes 8 strictly similar structures with SC7 motif, an  $\begin{bmatrix} 2 & 30 \\ 4 & 7 \end{bmatrix}$ interplanar angle close to 60° and a fourfold cage provided by screw  $\frac{1}{2}$ <br>operators, as exemplified in Fig. 6a. A second group includes 7<br>structures whose motif is again a fourfold cage, but with 9 Å **Example 12** (Specified in Fig. 6a. A second group includes 1<br>
distribution of the structure expansion motifs includes the<br>
symmetry operators are<br>
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S5 and Table structures whose motif is again a fourfold cage, but with  $9 \text{ Å}$   $\text{H}^5$ PI(New Journal of Chemistry ins<br>
classification of the structure expansion motifs includes the<br>
symmetry operators acting among nearest-neighbor molecules,<br>
while a measure of the corrugation is provided by the angles<br>
be interplanar angles. The aggregation motif in these two groups can be classified as fourfold "herringbone-cage". Six other structures and the three structures with  $Z'$  > 1 show packing motifs with a complex  $\frac{1}{70}$ admixture of first neighbors over various symmetry elements that defies a simple classification.

![](_page_6_Figure_5.jpeg)

1). SC7 = slipped-cofacial with 7 Å translation; 4 screw, coordination by 4 screw related correspondence of the first two major determinants. The spread and molecules. The upper right part is manifold of closely related symmetries.

### Isostructurality of rubrene crystals

The purely geometrical, qualitative analysis of packing modes can be supplemented by a quantitative analysis of coordination spheres by o structure determinants, plotting molecule-molecule energies  $E_{mm}$ against distance  $R_{mm}$ . These plots constitute a unique energetic  $\sim$  -10 profile of a crystal structure, by which similarities and differences can<br>20

An exemplary case of isostructurality is seen in Fig. 7: four<br>structures with same space group and nearly identical cell<br>parameters (they are also isomorphous). All structures show the SC7<br> $\frac{3}{2}$ <br> $\frac{40}{10}$ <br>E-stacking parameters (they are also isomorphous). All structures show the SC7  $\leq$  -40 -stacking pair, but being chemically different, the interaction energy varies from 45-55 kJ mol<sup>-1</sup> in the three phenyl substituted  $-50$ compounds to 62 kJ mol<sup>-1</sup> in the thiophene derivative, stabilized by a larger Coulombic contribution. These considerations show that -60 geometrical isomorphism is not always equivalent to energetic pairing in the fourfold cage, of nearly identical stabilization in all four structures. The second coordination shell (upper right part of the plot) includes an admixture of screw and translation (T) determinants, in a common area although with minor energetic differences.

![](_page_6_Figure_10.jpeg)

Fig. 8 Energetic profiles (kJ mol<sup>-1</sup>) for a series of almost isostructural crystals (see Table 1). Tx, Ty: translation along cell edges.

compounds are a hydrocarbon (QQQCIG13), a sparsely fluorinated compound (AXIDER) and a perfluoro derivative (INELUK02). The crystal structures are similar (same space group and cell parameters) but the energetic profile of the perfluoro compound shows a much larger cohesive energy in the first determinant, due to a substantial Coulombic contribution (nearly zero in the other two structures). The second determinant is over a longer screw axis relationship, testifying a significant structure deformation due to perfluorination. **Fig. 8 g 9 10 R R R Fig. 8 Energetic profiles (k)** mot<sup>1</sup>) for a series of almost isostructural crystals (see Table 1). Tx, Ty: translation long cell edges.<br>
A second very nearly isostructural series is s **Fig. 8** Energetic profiles (ki mot<sup>2</sup>) for a series of almost isostructural crystals (see Table 1). Tx, Ty, translation along cell edges.<br>
1). Tx, Ty, translation along cell edges.<br>
2). Tx, Ty, translation along cell edg

The triclinic structures of the parent hydrocarbon (QQQCIG14), of the newly determined polymorph CIYNAB01, and (strangely enough) of the compound in which two lateral phenyls are substituted by pyridines, are isomorphous (same cell parameters). The energy minor differences in the second-neighbor coordination shell are marginally significant because the interaction energies are anyway small. Fig. 7 Energetic profiles (kJ mol<sup>-1</sup>) for a series of strictly isostructural crystals (see Table profiles in Fig. 9 prove that they are also isostructural, with a strict

![](_page_6_Figure_14.jpeg)

Fig. 9 Energetic profiles (kJ mol<sup>-1</sup>) for a series of three isostructural triclinic crystals. All pairings are by pure translation.

### Polymorphism of rubrenes

has three well described polymorphs, and our database includes two other cases of clear cut polymorphism (CIYNAB, INELUK). Other compounds (GORVIU, PIXPOD, PIXPUJ, CIYXUF) are apparently polymorphic, but one of the partners is heavily disordered; in such cases one may wonder whether the postulated polymorphism is a real material property or is rather the result of poor sample quality or handling. In any case these latter instances cannot be adapted to lattice energy analysis and were not considered.

Polymorphism can be discussed qualitatively by comparisons of space groups and cell dimensions, or by packing diagrams, but such analyses are often ambiguous and always suffer from a certain amount of subjectivity. Especially in this case, the energetic profiles differences.

![](_page_7_Figure_6.jpeg)

Fig. 10 The energetic profile (kJ mol<sup>-1</sup>) of the three polymorphs of pristine rubrene, quantifying the different packings.

particular, the monoclinic polymorph with shortest contact distance<br>
The Royal Society distinct from the other forms The<br>
second neighbor coordination sphene is also quite efferem in the<br>
second neighbor coordination sphe hydrocarbon. The stable orthorhombic polymorph and the triclinic polymorph have the same slipped-cofacial pairing by 7 Å translation (SC7), but then all other determinants are completely different: in particular, the monoclinic polymorph with shortest contact distance of 8.7 Å stands out as clearly distinct from the other forms The second neighbor coordination sphere is also quite different in the Table 2 for a structure determinion team of the structure determinion team in the structure of the profiles of the profiles of the profiles of the are three polymorphs of the are three polymorphs are three polymorphs are material phases, even though they have nearly identical densities<br>and particle as finish the AA GLD lattice associas (207, 205 and Fig. 11 shows the profiles for the CIYNAB and CIYNAB01 and packing coefficients. The AA-CLP lattice energies (207, 205 and 196 kJ mol-1 for monoclinic, orthorhombic and triclinic, respectively) are only marginally different, as is very often the case in polymorphic **Example 19 and 19** intramolecular energy differences may not be too large. Total energy differences between polymorphs should then be quite small, in agreement with the findings of quantum chemical calculations<sup>36</sup> ( $\approx$ 2  $kJ$  mol $-1$ ).

Table 2 shows the energy profiles for two polymorphs that crystallize in the same space group but with quite different packing patterns. The polymorph with one full molecule in the asymmetric processes at the crystal/environment interface.<sup>39</sup> The PBC approach unit relies mostly on inversion-related neighbors, while the best works for growth at low supersaturation, without strong solutepolymorph in which the molecule sits on a crystallographic inversion

Plate of S7<br>
Dournal Name<br>
Journal Name<br> **Polymorphism of rubrenes**<br> **Polymorphism of rubrenes**<br> **Polymorphism of rubrenes**<br> **Polymorphism (2' = 1/2)** has translation and a fourfold of screw-related<br>
The parent compound (d **PI New Journal of Chemistry** ins<br>
Journal Name<br> **Polymorphism of rubrenes**<br> **The parent compound (database refcode QQQCIG, see Table 1)**<br> **Polymorphism of ubrenes**<br> **The parent compound (database refcode QQQCIG, see Tabl**  $P1$  New Journal of Chemistry n.s<br>
bournal Name<br> **Polymorphism of rubrenes**<br> **Control of state and the control of structure similarly control of the anent compound (database refrode QQQCIG, see Table 1)** molecules in its **Example 10**<br> **Example 10**<br> **Example 2014**<br> molecules in its coordination shell; determinant analysis gives a precise indication of the packing differences. INELUK was determined at lower temperature (123 vs. 173 K) but has nevertheless a lower density (1.93 vs. 2.04 g cm<sup>-3</sup>) and a lower packing coefficient (0.72 vs. 0.77): as expected, its total intermolecular energy is much less stabilizing, mainly due to a large deficiency in dispersion (close packing) terms. The tetracene core is flat in INELUK02 but twisted in INELUK, leading to a different overall conformation of the substituents. The non-bonded contact energy between pairs of lateral overlapping  $C_6F_5$  groups, evaluated by the AA-CLP potentials, is destabilizing by 8 kJ mol $^{-1}$  in INELUK02, but stabilizing by 4 kJ mol<sup>-1</sup> in INELUK: our numbers are a clear indication of tradeoff between favorable intermolecular contact and a less favorable intramolecular conformation. Absolute values may not be good enough for a reliable estimate of the relative stability of the two polymorphs, for which a quantum chemical study<sup>36</sup> predicted total energy differences of a few kJ mol<sup>-1</sup>. Although R-factors are comparable (4.5 vs. 5.6%), the unusually large lattice energy difference, together with the differences in densities and packing coefficients, suggest that INELUK is a metastable polymorph. lecular energy is much less stabilizing, mainly due to a large<br>
yei m dispersion (close packing) terms. The tetracene core is<br>
EEUIXO2 but twisted in INELUK, leading to a different overall<br>
atation of the substituents. Th ELUKO2 but twisted in INELUK, leading to a different overall<br>not of the substituents. The non-bonded contact energy<br>points of lateral overlapping  $C_6F_5$  groups, evaluated by the<br>potentials, is destabilizing by 8 kJ mol<sup></sup> ation of the substituents. The non-bonded contact energy<br>paiss of lateral overlapping  $C_6$ F<sub>S</sub> groups, evaluated by the<br>potentials, is destabilizing by 8 kJ mol<sup>-1</sup> in INELUKO2, but<br>g by 4 kJ mol<sup>-1</sup> in INELUK: our numbe pairs of lateral overlapping C<sub>G</sub>F<sub>5</sub> groups, evaluated by the<br>potentials, is destabilizing by 8 k mol<sup>-1</sup> in INELUK02, but<br>potentials, is destabilizing by 8 k mol<sup>-1</sup> in INELUK02, but<br>of between favorable intermolecular ootentials, is destabilizing by 8 kJ mol<sup>-1</sup> in INELUKO2, but<br>g by 4 kJ mol<sup>-1</sup> in INELUKO2, but<br>off between favorable intermolecular contact and a less<br>intramolecular conformation. Absolute values may not be<br>ugh for a re

Table 2 Energetic profiles of two polymorphs, INELUK ( $P2_1/n$ , Z=4) and INELUK02 ( $P2_1/c$ , Z=2) Each row is a structure determinant composed of a symmetry operator, a distance between centers of mass and a pairing energy with its Coulombic, polarization, dispersion, and repulsion components. S, Screw, T, translation, INV, inversion center

						ergy differences of a few kJ mol <sup>-1</sup> . Although R-factors are nparable (4.5 vs. 5.6%), the unusually large lattice energy erence, together with the differences in densities and packing	
	efficients, suggest that INELUK is a metastable polymorph.						
						le 2 Energetic profiles of two polymorphs, INELUK ( $P2_1/n$ , Z=4) and INELUK02 ( $P2_1/c$ , Each row is a structure determinant composed of a symmetry operator, a distance ween centers of mass and a pairing energy with its Coulombic, polarization, ersion, and repulsion components. S, Screw, T, translation, INV, inversion center	
			INELUK $P2_1/c$ Z=4				
$O_{mm}$	$R_{mm}$	Coul	pol	disp	rep	$E_{mm}$	
<b>INV</b>	8.081	$-13.5$	$-7.2$	$-73.0$	30.1	$-63.6$	
INV	8.663	$-8.3$	$-6.2$	$-57.9$	21.8	$-50.6$	
2S	10.593	$-5.7$	$-5.3$	$-48.0$	25.0	$-34.0$	
2Ty	11.149	$-6.7$	$-3.6$	$-28.9$	13.1	$-26.2$	
2S	11.618	$-4.5$	$-4.3$	-35.4	22.3	$-21.8$	
Elatt		$-34.6$	$-35.4$	$-229.3$	115.1	$-184.2$	
				INELUK02 P21/c Z=2			
2T	8.902	$-20.4$	$-6.3$	$-64.4$	31.4	$-59.8$	
4S	10.423	$-8.1$	$-5.3$	$-48.3$	25.0	$-36.7$	
2T	10.527	$-0.5$	$-4.0$	$-32.0$	13.6	$-22.9$	
	13.665	$-3.5$	-3.3	$-24.9$	15.9	$-15.8$	
4S		$-47.5$	$-38.6$	$-262.4$	133.4	$-215.1$	

![](_page_7_Picture_682.jpeg)

### Periodic bond chains analysis of crystal morphologies

PBC theory<sup>37,38</sup> has been long exploited for modeling theoretical crystal morphologies. Comparison with experimental outcomes provides useful information about the mechanism of growth **Example 19.1.1.1010 44.3 44.3 44.3 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44.4 44 Example 1998**<br> **Example 1998**<br> **Example 1998**<br> **Example 10.423** - 46.3 - 64.4 - 31.4 - 59.8<br> **Example 10.527** - 0.5 - 4.4 - 4.3 - 2.20 - 13.6 - 2.2.9<br> **Example 10.527** - 0.5 - 4.4 - 3.3 - 2.4 - 13.5 - 2.5 - 4.5 - 4.5 - 4 solvent interactions (e.g. vacuum sublimation), and when

![](_page_8_Figure_3.jpeg)

Fig. 11 Proof of the different packing of the new polymorph CIYNAB01.

faces, characterized by two or more non collinear periodic chains running parallel to the pertinent (hkl) plane. Slices of  $d_{hkl}$  thickness agreeing with extinction rules of the space group<sup>42</sup> represent layers. The growth units were defined as single molecules with a twice that of {200}, due to the lower energy of the c bond. Flat faces planar tetracene backbone as observed in the crystal structures. Attachment energies  $E_{att}^{hkl}$  for F faces were calculated with the reference molecule interacting with a half crystal exposing the (hkl) plane, applying a 25 Å cutoff for distance between molecular centroids. They were corrected for the energy required to convert<br>connected through weak **d** bonds (Fig. S6, ESI+). This face does not the tetracene backbone from the twisted conformation of vapor or solution to the planar one found in crystals, assuming an average value of 10 kJ mol<sup>-1</sup> from the 6-16 kJ mol<sup>-1</sup> range.<sup>18,20,36,43</sup>

Surface energies per unit area  $y_{hkl} = W_{hkl} / 2A_{hkl}$  were estimated for a reference molecule attached to the (hkl) plane, where  $W_{hkl}$  is

plane and  $A_{hkl}$  is the 2D unit cell area.  $W_{hkl}$  has been estimated with the same  $E_{mm}$  values used for attachment energies, with just minor approximation thanks to the fast decay of intermolecular (mainly dispersion) energies with distance. No significant reconstruction/relaxation has been observed in rubrene surfaces, 44 supporting the application of Born-Stern definition with cuts of the bulk crystal structure. Theoretical equilibrium and growth crystal morphologies have been obtained with Wulff plots<sup>45</sup> of  $\gamma_{hkl}$  and  $E_{att}^{hkl}$ , respectively. **Fig. 1.1** Fig. 3 For the same  $E_{mm}$  values used for attachment energies, with just minor approximation thanks to the fast decay of intermolecular (mainly dispersion) energies with distance. No significant energies with

Table 3 Intermolecular bonds in orthorhombic parent rubrene and flat faces, with bond  $(ESI<sup>+</sup>)$ .

<b>ARTICLE</b>																<b>Journal Name</b>
0 -10 $-20$ $\mathsf{E}_{\mathsf{mm}}$ / kJ mol <sup>-1</sup> $-30$ $-40$ -50 -60	т		$\mathbf T$ 4Screw			W all T	Ш	<b>• CIYNAB</b> CIYNAB01			dispersion) respectively. $(ESI+)$ .		plane and $A_{hkl}$ is the 2D unit cell area. $W_{hkl}$ has been estimated with the same $E_{mm}$ values used for attachment energies, with just minor approximation thanks to the fast decay of intermolecular (mainly energies with reconstruction/relaxation has been observed in rubrene surfaces, 44 supporting the application of Born-Stern definition with cuts of the bulk crystal structure. Theoretical equilibrium and growth crystal morphologies have been obtained with Wulff plots <sup>45</sup> of $\gamma_{hkl}$ and $E_{att}^{hkl}$ , Table 3 Intermolecular bonds in orthorhombic parent rubrene and flat faces, with bond contributions to slice energy. Sections of the energy crystal graph are reported in Fig. S6	distance.	No	significant
-70 6	7	8	9	10	11	12	13	14 15	16	17		Bond <sup>[a]</sup>	Symm. op. <sup>[b]</sup>	Distance	Bond energy	
Fig. 11 Proof of the different packing of the new polymorph CIYNAB01.					$R_{mm}$ / Å							a	$(x, 1 + y, z)$	7.173	$-64.8$	
												b	$(-x, \frac{1}{2}-y, \frac{1}{2}+z)$	7.975	$-45.9$	
intermolecular interactions in the growing crystal are dominated by fast decaying dispersive forces <sup>39-42</sup> as the case of rubrenes. The												c	$(\frac{1}{2}+x, \frac{1}{2}+y, z)$	13.866	$-17.8$	
underlying theory assumes that the time needed for the formation												d	$(\frac{1}{2} - X, -y, \frac{1}{2} + Z)$	15.171	$-6.5$	
of a crystal bond decreases with increasing bond energy, leading to																
the kinetic equation $R^{hkl}$ $\propto$ $E_{att}^{hkl}$ for flat (F) faces, where $R^{hkl}$ is the growth rate of the (hkl) face and $E_{att}^{hkl}$ is the energy released per												Flat face	Bonds in slice <sup>c</sup>	$E_{att}$ <sup>d</sup>	$y_{hkl}$ <sup>e</sup>	
growth unit when a slice of d <sub>hkl</sub> thickness is attached to the surface. <sup>38</sup>												${200}$	$a + 2b$	$-39.7$	80.8	
Therefore, the boundaries of crystal polyhedra are slowly growing faces, mainly flat (F) faces whose attachment energies are usually												${002}$	$a + 2c$	$-94.8$	90.6	
smaller than those of stepped or kinked faces. <sup>37,40</sup>												${111}$	$b + c + d$	$-120.6$	98.7	
									We analyze here in these aspects a set of structures, the			${202}$	$a + d$	$-121.3$	100.2	
orthorhombic parent rubrene (QQQCIG15) and the isomorphous monoclinic derivatives (CIYNAB, CIYXUF, CIYYAM, PIXPUJ01, see												${020}$	2d	$-174.2$	80.1	
Table 1). The growth morphologies of these crystals have been evaluated according to PBC methods <sup>28,29,40</sup> using intermolecular interactions (called bonds) between first neighbors, whose energy is given by $E_{mm}$ values, derived from the more accurate force field formulation of the PIXEL approach, calculated between a reference molecule and the surrounding molecules within a cutoff of 25 Å. Bonds whose $E_{mm}$ was less than $kT \approx 4$ kJ mol <sup>-1</sup> (at the actual crystal growth temperature of 500-550 K) were discarded, leaving just four relevant bonds in the selected crystal structures. faces, characterized by two or more non collinear periodic chains running parallel to the pertinent (hkl) plane. Slices of $d_{hkl}$ thickness agreeing with extinction rules of the space group <sup>42</sup> represent physically sound molecular profiles of surfaces and natural growth									The PBC analysis allows to identify crystal planes representing F				[a] Bond between the reference and a neighbor molecule. [b] Symmetry operation connecting reference and neighbor at the listed distance between centers of mass. [c] A slice is a set of layers with overall thickness dhki. [d] Attachment energy corrected for twisted to flat tetracene backbone. [e] Surface energy, erg cm <sup>-2</sup> . Distances in Å, energies in kJ mol <sup>-1</sup> . Table 3 lists molecule-molecule energies $E_{mm}$ defining bonds for orthorhombic rubrene. Crystal graphs of intermolecular bonds are reported in Fig. S6 (ESI <sup>+</sup> ). The two strongest bonds <b>a</b> and <b>b</b> lie in the (200) slice and generate a robust growth layer with the slipped- cofacial motif. Within this slice the growth units are incorporated at the end of bond chains, resulting in layer-by-layer growth with stable and flat surface profiles. Experimental evidence from AFM imaging of rubrene {100} surfaces shows accordingly monomolecular (200) steps. <sup>46-48</sup> The attachment energy of the $\{002\}$ surface is more than			

Ereas The total Society and the selection of the present of the selection of the selection of the matter of the selection of the selecti growth temperature of 500-550 K) were discarded, leaving just four<br>reported in Fig. S6 (ESI+). The two strongest bonds a and **b** lie in the The PBC analysis allows to identify crystal planes representing F<br>cofacial motif. Within this slice the growth units are incorporated at grown in the Uniter of the means a state of the state of surface and the state of surfaces and physically preserved in the contents of considers of considers of consideration and solve y growth molecular profiles of the m orthorhombic rubrene. Crystal graphs of intermolecular bonds are (200) slice and generate a robust growth layer with the slipped**cofacial coff**  $(26\pi)$ ,  $\sqrt{3}$ ,  $\pi$  are the growth units are incorporated at the growth units are incorporated at the growth units are incorporated by the slice the growth connecting reference and a neighbor molecule. the end of bond chains, resulting in layer-by-layer growth with stable and flat surface profiles. Experimental evidence from AFM imaging of rubrene {100} surfaces shows accordingly monomolecular (200) steps.46-48 The attachment energy of the {002} surface is more than that of  $202$  at  $+2c$   $-94.8$  90.6<br>
(111)  $b+c+d$  -120.6 98.7<br>
(202)  $a+d$  -121.3 100.2<br>
(220)  $2d$  -174.2 80.1<br>
(a) Bond between the reference and a neighbor at the listed distance between centers of mass.<br>
[c] A slice is {111} and {202} are further down the ranking of  $E_{att}^{hkl}$  due to the presence of weaker **d** bonds. Interestingly, the {202} surfaces is similar to a stepped surface with [010] rows (the direction of the strongest bond and PBC) delimited by {200} and {002} facets (020) 2d -174.2 80.1<br>
(a) Bond between the reference and a neighbor molecule. [b] Symmetry operation<br>
connecting reference and neighbor at he listed distance between centers of mass.<br>
E[c] A slice is a set of layers with appear in the theoretical growth morphology. Similarly, the high  $E_{att}^{hkl}$ of face {020}, based only on d bonds, makes it disappear from the growth morphology while it is important in the equilibrium shape. Table 3 lists molecule-molecule energies  $E_{mm}$  defining bonds for orthorhombic rubrene. Crystal graphs of intermolecular bonds are reported in Fig. S6 (ESI<sup>+</sup>). The two strongest bonds a and **b** lie in the line of 200) s

the separation work required to split an infinite crystal along the (hkl) the lowest  $E_{att}^{hkl}$  and  $\gamma_{hkl}$  values (Fig. 12). As reported In summary, equilibrium, and even more strongly, growth morphology of orthorhombic rubrene is dominated by {200} surfaces

 $\mathbf{1}$ 

less rich in crystal faces than the equilibrium one and displays a higher anisotropy, definitely tabular, determined by the robust slice similar  $\gamma_{hkl}$ , resulting in more isotropic equilibrium crystal habits.

![](_page_9_Figure_4.jpeg)

The four monoclinic rubrene derivatives are isomorphous despite different functional groups on rings 5/11 or the presence of thienyl instead of phenyl rings (Table 1). Table 4 shows the results of PBC analysis. All these structures display (100) slices with the slippedcofacial motif found in the (200) slice of orthorhombic rubrene and Share the same four strong bonds. The equilibrium of and displays and the same four strong bonds. The overall picture of intermolecular paid the same of the same  $[010]$ ,  $[001]$ ,  $\langle 011 \rangle$ ,  $\langle 201 \rangle$  and  $\langle 211 \rangle$  and five flat faces:  $\{100\}$ ,  $[001]$ ,  $[001]$ ,  $\langle 011 \rangle$ ,  $\langle 201 \rangle$  and  $\langle 211 \rangle$  and five flat faces:  $\{100\}$ ,  ${10\overline{2}}$ , {011}, {002} and {111}. An example of the crystal graph Shared Physics attinge in more isotropic equilibrium crystal habits.<br>
Fig. 21 Equilibrium<br>
Terms are expected by PC analys Color codine [100] - webwe, for derivative in Fig. 13 Equilibrium crystal morphology of themselves CIYNAB. Surface the term of the term of the cytatic method in the cytatic m

Table 4 Intermolecular bonds for isomorphous monoclinic rubrenes (see Table 1 for refcodes) and corresponding flat faces. See Table 3 for units and the meaning of symbols.

![](_page_9_Picture_589.jpeg)

Bonds a and b are in all aspects analogous to orthorhombic rubrene giving rise to the most stable (h00) slice (of  $d_{100}$  thickness in **CIYYAM** the monoclinic structures). The (h00) slice energies are in the range -150/-160 kJ mol<sup>-1</sup> out of a lattice energy of -200/-225 kJ mol<sup>-1</sup>. Bonds c and d arise from molecular pairs different from those in  $V$ ellow,  $\{10\bar{2}\}$ -cyan,  $\{011\}$ -orange,  $\{001\}$ -green,  $\{11\bar{1}\}$  absent.

![](_page_9_Figure_10.jpeg)

Fig. 13 Equilibrium crystal morphology of isomorphous monoclinic rubrene derivatives

orthorhombic rubrene because the monoclinic structure allows to accommodate the rougher (h00) interface of protruding para substituents (CIYNAB, CIYXUF, CIYYAM) or smaller thienyl rings (PIXPUJ01).

The reduced cohesion in the crystal of fluorinated derivative ascribed to the well-known, scarce propensity of hard fluorine for intermolecular stabilization, in contrast with some literature claims of the contrary.

a  $(x, 1+y, z)$  -56.8 -57.4 -56.3 -66.0 absence of the  $\{11\}$  form in the growth shape while  $\{001\}$  is The overall picture of intermolecular interactions is summarized in the theoretical equilibrium and growth morphologies. The decreased surface energy for {100} in the trifluoromethyl derivative produces thinner crystals along the reciprocal direction  $a^*$  with an **Fig. 13 Equilibrium crystal morphology of isomorpholos monoclinic rubrene derivatives<br>
erline, Cristol, Cristol, NeXPUD as predicted by PBC analysis. Clore coding: [100]-<br>
vellow, [102] - cyan, (011) - orange, (001) - gre EXECT INTERT CONTROVER CONTROVER CONTROVER CONTROVER CONTROVER CONTROVER CONTROVER CONTROLLY (1023 - cyan, (011) - orange, (001) - green, 1117) - blue.<br>
Delow, 11023 - cyan, (011) - orange, (001) - green, 1117) - blue.<br>
O** tabular habit on {100}, the clearer example being the trifluoromethyl case. Another feature common to these monoclinic crystals is the

![](_page_9_Figure_15.jpeg)

Fig. 14 Growth crystal morphology of isomorphous monoclinic rubrene derivatives

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negligible in the nitro-rubrene (CIYNAB) and absent in the trifluoromethyl-rubrene (CIYXUF).

Comparison among predicted and experimental growth crystal morphologies of rubrene derivatives grown in a moderate vacuum can be appreciated in Fig. 15. Further examples of vacuum grown crystals of rubrene are available in several papers.48,49 For rubrene and derivatives CIYNAB, CIYXUF and CIYYAM agreement among PBC morphology and experimental ones is satisfactory, considering that significant variability in crystal shapes even under the same experimental conditions is always observed. The strong effect of the trifluoromethyl groups in CIYXUF previously highlighted is clearly demonstrated by the growth of thin tabular crystals (Fig. 15c).

![](_page_10_Figure_6.jpeg)

Fig. 15 a-d) Experimental morphologies for crystals of rubrene derivatives grown by vacuum sublimation. In e) are shown crystals of the thienyl derivative grown from hexane/ethyl acetate 9:1 vol/vol. f) sketch of the observed crystal morphology in e) using the color coding of Fig. 14. Crystalline needles shown in c) belong to a concomitant triclinic polymorph.<sup>50</sup>

The thienyl-rubrene PIXPUJ01 could not be vacuum crystallized due to its thermal instability. Crystallization from hexane/ethyl acetate 9:1 vol/vol leads to [010] elongated crystals with {100},  ${10\overline{2}}$  and {011} bounding faces. The influence of the solvent mixture and of several byproducts in the final reaction mixture<sup>17</sup> acting as impurities are likely responsible for the modification of the crystal morphology. It is worth mentioning that the monoclinic polymorph

of di-p-nitrophenylrubrene CIYNAB has been obtained also by very slow isothermal evaporation of an acetone solution with a morphology indistinguishable from that of Fig. 15b. Faster isothermal evaporation of an acetone solutions of di-pnitrophenylrubrene afforded a solvated species.<sup>50</sup> Examples of the effect of solvent nature upon crystal morphology of pristine rubrene can be found in previous work.<sup>51,52</sup>

### Summary and final remarks

The available crystallography of rubrene materials has been thoroughly reviewed through deposited X-ray determinations, with the aim of establishing a database of fully reliable crystal structures for theoretical studies. The final result was 33 crystal structures double-checked and normalized with respect to hydrogen-atom positions. All the structures include at least one highly stabilizing molecular pair at a center-of-mass distance of 7-8 Å. The database has been analyzed systematically in terms of structural patterns, with a statistical assessment of the frequency of the main aggregation modes. The slipped-cofacial pairing arrangement is almost ubiquitous, being an obvious coupling ("encroaching") mode given the irregular molecular shape: this mode permits as much tetracene stacking as possible, indispensable for promoting transport. Its propagation in three-dimensional monoclinic networks often goes through the herringbone pattern, of which two variants have been found, one with shortest translation of 7 Å and tetracene inter-planar angle of 60°, and another with shortest translation of 9 Å and highly variable inter-planar angles. Triclinic structures occur with a perforce parallel arrangement of tetracene cores. These common features are not exclusive, as examples of other close pairing and propagation modes have been found. Somewhat unexpectedly, in a few cases even asymmetric pairing of two molecules in Z' = 2 crystals leads to very high stabilization without intervention of a symmetry operation. Systematization has proceeded to a point, but only sparse clues have appeared as concerns the dependence of packing modes on substituent chemistry, or on molecular conformation (twisting of the tetracene core).

Isostructurality has been analyzed by lattice energy calculations and plots of pairing energy profiles, that reveal at a glance similarities and differences in crystal packing over and above a sometimes dubious comparison of cell parameters. A classic example of isostructurality (and isomorphism) appears for three parasubstituted (NO<sub>2</sub>, CF<sub>3</sub> and CN) lateral phenyls, and also surprisingly, a di-thiophene derivative. Three triclinic structures, the parent hydrocarbon, the di-p-nitrophenyl derivative, and a dipyridine derivative, are also isomorphous. These findings suggest that crystal packing is a driving force that supersedes or at least attenuates the effects of chemical substitution.

Polymorphism is only sparse in the database, the scarce solubility and stability of rubrenes hampering or preventing extensive screenings. Our survey accepts only three cases of undisputable determination: first, the parent hydrocarbon, for which three well determined polymorphs have been deposited, so many presumably because it has been subjected to wider screening efforts; then, the perfluoro compound with two polymorphs, presumably for the same reasons; although our energy calculations strongly suggest that one polymorph is metastable or even unstable.

 $\mathbf{1}$  $\overline{2}$  $\overline{3}$  $\overline{4}$ 5 6  $\overline{7}$ 8 9  $10$  $11$  $12$ 

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We report here the structure of a newly found polymorph of the di-p-nitro derivative, obtained after substantial screening: the improved stability toward oxidation of di-p-nitrophenylrubrene allowed crystallization attempts with different solvents affording the monoclinic polymorph or acetone and dichloromethane solvate crystals. The triclinic polymorph has been obtained only by vacuum sublimation with a steeper temperature gradient compared to conditions leading to the monoclinic structure. The comparison of polymorphic energy profiles provides an immediate and univocal proof of the diversity of the crystal packings, the prerequisite for acceptance of a polymorph as a legitimate new material.

13 A thorough analysis of crystal structures in terms of interaction 14 energies is also fruitful for reliable modeling of crystal 15 morphologies.<sup>28</sup> Good agreement between calculated and observed 16 crystal morphologies has been demonstrated for acenes<sup>40,41</sup> and 17 oligothiophenes.<sup>42</sup> This approach naturally provides the 18 19 crystallographic planes and cuts of surfaces reasonably involved in crystal growth processes. Other simpler (and much less demanding) 20 techniques generally provide heavily biased and/or uncomplete  $21$ results.<sup>40</sup> The growth morphologies obtained in the present study<br>from analysis of crystal structures via the BBC approach compare 1 J. E. Anthony, Angew. Chem. Int. Ed. 2007, 46, 2-34. 22 from analysis of crystal structures via the PBC approach compare 1 J.E. Anthony, Angew. Chem. Int. Ed. 2007, 4<br>19. Setisfacts with another compare from the process The radio 2 A. Yassar, Polym. Sci. Ser. C, 2014, 56, 4-19  $23$ satisfactorily with crystals grown from the vapor. The only  $\frac{2}{3}$  N. Podzorov, V. M. Pudalov and M. E. Gershenson, Appl. Phys. 24 morphology differing appreciably from the predicted one involves 25 the thienyl derivative. In that case a complex mixture of byproducts 4 V. Podzorov, S. E. Sysoev, E. Loginova, V. M. Pudalov and M. 26 (separated with difficulty and incompletely from the main product E. Gershenson, Appl. Phys. Lett. 2003, 82, 3504-3506.<br>with column chromatography followed by fractional crystallizations) 5 M. L. Clapham, E. C. Murphy and 27 with column chromatography followed by fractional crystallizations) 28 29 with molecular structures related to the thienyl rubrene<sup>17</sup> are likely<br>  $6T$ . K. Mullenbach, K. A. McGarry, W. A. Luhman, C. J. Douglas responsible for modification of the crystal morphology.53 30

The presence of strongly bound (h00) layers in all five analyzed 7 D. A. da Silva Filho, E. G. Kim and J. L. Brédas, Adv. Mater.  $31$ structures arises from the slipped-cofacial arrangement adopted 2005, 17, 1072-1076.<br>with only small differences. This feature is associated with the 8 M. D. Curtis, J. Cao and J. W. Kampf, J. Am. Chem. Soc. 2004, 32 with only small differences. This feature is associated with the 8 33 strongest periodic bond chains and determines low values for surface<br>
g C. Janiak, J. Chem. Soc., Dalton Trans. 2000, 3885-3896. 34 energy and attachment energy, respectively determining the 10 V.C. Sundar, J. Zaumseil, V. Podzorov, E. Menard, R. L. Willett, 35 36 equilibrium and growth shape. The similarities in the overall pattern of intermolecular bonds and, hence, of periodic bond chains explain 37 the limited variations of observed growth morphologies. These 38 correspond to crystals defined by Hartman's flat faces characterized 39 correspond to crystals defined by Hartman's nat races characterized 12 E. Fumagalli, L. Raimondo, L. Silvestri, M. Moret, A. Sassella<br>by the lowest attachment energies among all possible and M. Campione. Chem. Mater. 2011. 40 41 crystallographic planes. Finally, as already noted in other organic 13 S. Bergantin and M. Moret, Cryst. Growth Des. 2012, 12, 6035semiconductor crystals, growth morphologies are more anisotropic 6041.<br>than equilibrium ones <sup>40-42</sup> with different degrees of tabular shanes 14 D. Braga, A. Jaafari, L. Miozzo, M. Moret, S. Rizzato, A. Papagni 42 than equilibrium ones, <sup>40-42</sup> with different degrees of tabular shapes 43 where the largest surfaces comprise structural layers based on the and A. Tassar, Lur. J. Org. Chem. 2011, 4100-4105.<br>15 Y. Sakamoto, T. Suzuki, J. Org. Chem. 2017, 82, 8111-8116. 44 slipped-cofacial motif. The corresponding low values for the 16 K.A. McGarry, W. Xie, C. Sutton, C. Risko, Y. Wu, V. G. Young, 45 attachment energy make these crystallographic planes dominate the 46 crystal morphology. 47

48 potentials show promise for access to lattice energies and 49 morphology prediction without recourse to computationally 18 C. Sutton, M. S. Marshall, C. D. Sherrill, C. Risko and J. L. 50 intensive quantum chemical methods. As concerns the materials  $51$ science involved, in general, from our overview it looks like sensitive 52 spots of the driving forces for rubrene packing are the 4-substitution<br> $\frac{2005}{95}$ , 166602.<br>20 A. S. Paraskar, A. R. Reddy, A. Patra, Y. H. Wijsboom, O. 53 sites at the lateral rings, with substituents of moderate steric bulk. 54 Peripheral substitution at the tetracene core seems to be less 55 influential in steering the packing and hence the physical properties. 21 S. Bergantin, M. Moret, G. Buth and D. P. A. Fabbiani, J. 56 Our survey provides a structural background that could generate Phys.Chem. C 2014, 118, 13476-13483.<br>new ideas directing the synthesis of new derivatives and provide 22 O.D.Jurchescu, A. Meetsma and T.T. Palstra, Acta Crys 57 new ideas directing the synthesis of new derivatives and provide 58 hints for the development of enhanced transport properties. 59

Massimo Moret (principal investigator) provided the framework for the research project and is mainly responsible for the PBC part. Angelo Gavezzotti developed the methodology for the crystallographic database studies. Both authors concur in the preparation and presentation of the manuscript. **Author Contributions**<br>
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erystallogra **Author Contributions**<br>
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Angelo Gavezzotti developed the methodology for the<br>
crystallogra **Example 12**<br>
Massimo Moret (principal investigator) provided the framework for<br>
the research project and is mainly responsible for the PBC part.<br>
Angelo Gavezzotti developed the methodology for the<br>
crystallographic datab the research project and is mainly responsible for the PBC part.<br>Angelo Gavezzotti developed the methodology for the<br>erystallographic database studies. Both authors concur in the<br>preparation and presentation of the manuscr

### Conflicts of interest

Author Contributions

There are no conflicts to declare.

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Electronic Supplementary Information (ESI) for New Journal of Chemistry

The crystalline state of rubrene materials: intermolecular recognition, isomorphism, polymorphism, and periodic bond-chain analysis of morphologies

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# Appendix: crystallographic terminology

Throughout the article, the following notations/terms are extensively used:

[uvw] crystallographic direction. Any direction in a Bravais lattice is identified by a set of N indices (N dimensionality of the space), written in square brackets, called the direction indices. In three-dimensional space, the direction passing through the origin and the New Journal of Chemistry<br> **Appendix:** crystallographic terminology<br>
Throughout the article, the following notations/terms are extensively used:<br> **[uwv]** crystallographic direction. Any direction in a Bravais lattice is ide identify the same direction as [uvw] observed from the opposite side.

<uvw> set of symmetry related crystallographic directions according to space group symmetry

(hkl) crystallographic plane identified by its Miller indices

{hkl} set of symmetry related crystallographic planes (form) according to space group symmetry

**isomorphous crystals**: two crystals are said to be isomorphous if (a) both have the same **Appendix: crystallographic terminology**<br>
Throughout the article, the following notations/terms are extensively used:<br> **[uvw]** crystallographic direction. Any direction in a Bravais lattice is identified by a set of N<br>
in are the same except for a replacement of one or more atoms in one structure with different types of atoms in the other (diadochy), such as heavy atoms, or the presence of one or more additional atoms in one of them (isomorphous addition). Isomorphous crystals can form solid solutions

**isostructural crystals**: Two crystals are said to be *isostructural* if they have the same structure, but not necessarily the same cell dimensions nor the same chemical composition, and with a 'comparable' variability in the atomic coordinates to that of the cell dimensions and chemical composition. One also speaks of isostructural series. The term isotypic is synonymous with isostructural.

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![](_page_15_Figure_3.jpeg)

Figure S1. RAGCUA, short distance coupling with low stabilization due to parallel tetracene cores without offset. TEFDUG, close packing is restored by the insertion of another filler in the gap. These structures are outliers in the top left part of Figure 2b. Fluorine is green.

![](_page_15_Figure_5.jpeg)

Figure S2. Molecule-Molecule energy (kJ/mol) vs. center of mass distance for the subset of slipped-cofacial pairings out of Figure 2b. No correlation

![](_page_16_Picture_2.jpeg)

![](_page_17_Figure_2.jpeg)

Figure S3. A gallery of slipped-cofacial arrangements with CSD refcodes (see Table 1).

![](_page_18_Picture_2.jpeg)

 $E$ (mm) = -70 kJ mol<sup>-1</sup>.

![](_page_18_Figure_4.jpeg)

Figure S5. Plot of angles between tetracene planes in corrugated (herringbone) propagation motifs. The T7+4S cage motif (Figure 6a) has consistently a 60° angle.

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![](_page_19_Figure_2.jpeg)

 $\mathbf{1}$ 

![](_page_20_Figure_2.jpeg)

 $\mathbf{1}$  $\overline{a}$  $\overline{\mathbf{3}}$  $\overline{\mathcal{L}}$ 

![](_page_21_Figure_2.jpeg)

Figure S6. Energy crystal graphs in rubrene (QQQCIG05). Intermolecular bonds (see Table 3 in the manuscript) are color

![](_page_22_Figure_2.jpeg)

![](_page_23_Figure_2.jpeg)

![](_page_24_Figure_2.jpeg)

Table 4 in the manuscript) are color coded as: bond a yellow, bond b green, bond c cyan, bond c purple.

![](_page_25_Picture_685.jpeg)

 $\overline{\mathbf{1}}$ 

TABLE S1. List of all the rubrene crystal structures in the Cambridge Structural Database. From left to right: Identification code of the Cambridge<br>Structural Database (CSD), Crystallographic R-factor. Spc group, Motocolo **TABLE S1.** List of all the rubrene crystal structures in the Cambridge Structural Database. From left to right: Identification code of the Cambridge<br>Structural Database (CSD), Crystallographic Refactor, Spc group, Molecu collection (K), Cell parameters (Šand degrees), Unit cell volume (Ū), Reduced cell parameters (Å and degrees), Reduced cell volume (Ū).  $^{3}$ ).

Refcode	Rfac	Space Group	z	Z'	T.	a	b	c	α	β	γ	Volume	a	b	c	$\alpha$	Reduced   Reduced   Reduced   Reduced   Reduced   Reduced		Reduced Volume
<b>AXIDER</b>	6.35	P21/c	$\overline{c}$	0.5	293	9.0298	9.909	16.217	90	90.227	90	1451	9.0298	9.9085	16.217	90	90.227	90	1451
<b>CIYNAB</b>	6.73	P21/c	$\overline{2}$	0.5	153	15.098	7.171	14.249	90 <sub>1</sub>	100.616	90	1516.2	7.1706	14.2489	15.0983	100.616	90	90	1516
<b>CIYXUF</b>	6.59	P21/c	2	0.5	123	15.978	7.276	13.981	90 I	102.701	90	1585.7	7.2762	13.9814	15.9782	102.701	90	90	1586
<b>CIYYAM</b>	6.65	P21/c	$\overline{2}$	0.5	120	14.913	7.115	14.426	90	98.099	90	1515.4	7.1151	14.4263	14.9125	98.099	90	90	1515
<b>GORVEQ</b>	7.77	P212121	4		93	9.599	13.29	25.015	90	90	90	3190.5	9.599	13.287	25.015	90	90	90	3190
<b>GORVIU</b>	4.59	$P-1$	3	1.5	93	10.553	11.34	16.568	83.481	89.45	64.275	1772.9	10.553	11.34	16.568	83.481	89.45	64.275	1773
GORVIU01	4.6	P21/c	4		93	14.386	8.452	21.794	90	116.994	90	2361.2	8.452	14.386	19.9328	103.0291	90	90	2361
<b>GORVUG</b>	4.5	$P-1$	$\overline{c}$		93	8.5037	13.24	14.277	62.458	80.781	74.581	1373.1	8.5037	13.244	14.277	62.458	80.781	74.581	1373
<b>INELUK</b>	5.62	P21/n	4		123	16.665	11.15	19.192	90	90.213	90	3565.8	11.1489	16.6651	19.1918	90.213	90	90	3566
INELUK01	6.87	$P-1$	6	3	123	12.529	17.82	24.734	90.202	101.264	90.695	5414	12.5285	17.8163	24.7339	90.202	101.264	90.695	5414
INELUK02	4.5	P21/c	$\overline{2}$	0.5	173	8.9017	10.53	17.992	90	90.408	90	1686	8.9017	10.527	17.992	90	90.408	90	1686
INELUK03	7.34	P21/n	4		173	16.635	11.13	19.245	90	90.581	90	3564.1	11.1334	16.635	19.245	90.581	90	90	3564
<b>KUSDEJ</b>	5.27	Cmca	4	0.25	173	34.197	7.158	14.052	90	90	90	3439.5	7.1577	14.052	17.469	90	101.8218	90	1720
<b>MIVCUR</b>	4.75	C2/c	12	1.5	173	41.257	10.15	24.038	90	96.739	90	9993	10.1464	21.2432	24.038	96.5431	90	103.8166	4997
<b>MIVDAY</b>	4.31	P21	$\overline{2}$		173	12.75	9.414	14.503	90	99.576	90	1716.4	9.4141	12.7495	14.5027	99.576	90	90	1716
<b>MIVDEC</b>	4.21	Pna <sub>21</sub>	4		173	15.205	13.9	14.31	90	90	90	3024.6	13.901	14.3104	15.2045	90	90	90	3025
<b>MIVDIG</b>	9.52	Pbcm	4	0.5	123	7.1443	14.05	34.143	90	90	90	3427.4	7.1443	14.051	34.143	90	90	90	3427
<b>MIVDOM</b>	7.61	Pbcm	4	0.5	123	7.5317	14.63	32.424	90	90	90	3573.7	7.5317	14.634	32.424	90	90	90	3574
<b>MIVDUS</b>	4.59	Pnma	4	0.5	123	7.1162	31.18	14.208	90	90	90	3152.3	7.1162	14.2075	31.179	90	90	90	3152
PIFHIW	7.42	Pnma	4	0.5	292	14.158	35.39	7.2215	90	90	90	3618.3	7.2215	14.158	35.39	90	90	90	3618
PIFHOC	9.77	P21/c	4		292	23.527	9.028	17.764	90	95.928	90	3752.8	9.0277	17.764	23.527	95.928	90	90	3753
<b>PIXPOD</b>	4.3	P21	$\overline{2}$		173	11.401	8.994	13.989	90	113.59	90	1314.6	8.9944	11.401	13.989	113.59	90	90	1315
PIXPOD01	5.01	P21/c	Δ		93	13.601	9.248	20.718	90	92.771	90	2602.9	9.2479	13.601	20.718	92.771	90	90	2603
<b>PIXPUJ</b>	6.49	P21/c		0.5	173	10.173	8.915	14.751	90	95.17	90	1332.3	8.9148	10.173	14.751	95.17	90	90	1332
PIXPUJ01	6.87	P21/c	2	0.5	120	13.568	7.014	14.302	90 <sub>1</sub>	103.954	90	1320.9	7.0143	13.5679	14.302	103.954	90	90	1321
POGZIV	6.91	P21/c	8	$\overline{c}$	294	22.036	8.803	30.252		90 104.921	90	5670.8	8.8034	22.036	30.252	104.921	90	90	5671

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![](_page_26_Picture_738.jpeg)

43<br>44<br>45<br>46<br>47

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![](_page_27_Picture_394.jpeg)

60

Table S2. Comparison of the CLP, LJC and Pixel force fields for some molecule-molecule energies in some isostructural rubrene derivatives. For each entry:

# New Journal of Chemistry<br> **Table S2.** Comparison of the CLP, LJC and Pixel force fields for some molecule-molecule<br>
energies in some isostructural rubrene derivatives. For each entry:<br>
Indecule-molecule distance (A), symme New Journal of Chemistry<br> **Table S2.** Comparison of the CLP, LJC and Pixel force fields for some menergies in some isostructural rubrene derivatives. For each entry:<br>
molecule-molecule distance (A), symmetry operation numb New Journal of Chemistry<br> **Table S2.** Comparison of the CLP, LJC and Pixel force fields for some menergies in some isostructural rubrene derivatives. For each entry:<br>
molecule-molecule distance (Å), symmetry operation numb symmetry operation New Journal of Chemistry<br> **able S2.** Comparison of the CLP, LJC and Pixel force fields for some molecule-molecule<br>
nergies in some isostructural rubrene derivatives. For each entry:<br>
nolecule-molecule distance (A), symmetr New Journal of Chemistry<br> **able S2.** Comparison of the CLP, LJC and Pixel force fields for some molecule-molecule<br>
mergies in some isostructural rubrene derivatives. For each entry:<br>
nolecule-molecule distance (A), symmetr CIYXUF 7.880 2 0.00 -0.50 -0.50<br>
-6.7 0.0 -73.9 27.4 -53.2<br>
-17.21 -7.26 -64.40 38.08 -50.79<br>
-6.1 -4.7 -58.8 19 -6.7 0.0 -73.9 27.4 -53.2  $-17.21$   $-7.26$   $-64.40$   $38.08$   $-50.79$  $-6.1$   $-4.7$   $-58.8$   $18.6$   $-51.0$  7.276 1 0.00 -1.00 0.00 8.2 0.0 -109.9 41.4 -60.4 -3.88 -10.47 -97.94 54.91 -57.38 -3.88 −10.47 −97.94 54.91 −57.38<br>7.7 −5.2 −87.9 29.7 −55.7  $CIVVAM$  8.043 2.0.00 0.50 0.50

molecule-molecule distance (Å), symmetry operation number

![](_page_28_Picture_271.jpeg)

 $\mathbf{1}$  $\overline{2}$  $\overline{\mathbf{3}}$  $\overline{\mathcal{A}}$ 5  $\boldsymbol{6}$  $\overline{7}$  $\bf 8$ 

Table S3. Energy (CLP) and density data for the crystal structures in the database.

For each entry:<br>first line: CSD refcode Cell vol  $(A^3)$  density (g cm<sup>-3</sup>) packing factor Fable S3. Energy (CLP) and density data for the crystal structures in the database.<br>
For each entry:<br>
first line: CSD refcode Cell vol  $(A^3)$  density (g cm<sup>-3</sup>) packing factor<br>
second line: CSD refcode Ecoul Epol Edisp Er

![](_page_29_Picture_265.jpeg)

 $\overline{3}$  $\overline{\mathbf{3}}$  $3<sup>0</sup>$  $\overline{4}$  $\overline{\mathcal{L}}$  $\overline{4}$  $\overline{4}$  $\overline{4}$  $\overline{4}$  $\overline{4}$  $\mathbf{A}$  $\overline{4}$  $\overline{4}$  $\overline{5}$  $\sqrt{5}$  $5<sub>i</sub>$  $5<sub>i</sub>$ 

![](_page_30_Picture_187.jpeg)

 $\mathbf{1}$ 

**Table S4.** A summary of pairwise molecular couplings with the slipped cofacial configuration.<br>Distances in Å, energies (AA-CLP) in kJ mol<sup>-1</sup>. T, translation, Inv inversion, other operations New Journal of Chemistry<br> **Table S4.** A summary of pairwise molecular couplings with the slipped cofacial configuration.<br>
Distances in Å, energies (AA-CLP) in kJ mol<sup>-1</sup>. T, translation, Inv inversion, other operations<br>
or with coordinate transformation. New Journal of Chemistry<br>
Table S4. A summary of pairwise molecular couplings with the slipped cofacial configuration.<br>
Distances in A, energies (AA-CLP) in kJ mol<sup>-1</sup>. T, translation, Inv inversion, other operations<br>
with

![](_page_31_Picture_201.jpeg)

Table S5. Extended motifs in the packing of rubrenes. Contacts with highest molecule-molecule contact energies. ournal of Chemistry<br>
of rubrenes. Contacts with highest molecule-molecule<br>
notice; Operator = T, I, S<br>
= inversion, T = translation, S = screw, G = glide<br>
crew related molecules

Codes: SC7 (O): slipped cofacial at 7 Å distance; Operator = T, I, S

4S: four identical screw operations, 2S: two screw related molecules

![](_page_32_Picture_263.jpeg)

 $\mathbf{1}$  $\overline{2}$   $\overline{1}$ 

![](_page_33_Picture_94.jpeg)

![](_page_33_Picture_95.jpeg)