Asymmetric rotations and dimerization driven by normal to modulated phase transition in 4-biphenylcarboxy coupled L-phenylalaninate

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Amongst the derivatives of 4-biphenylcarboxylic acid and amino acid esters, the crystal structure of 4-biphenylcarboxy-(L)-phenylalaninate is unusual owing to its monoclinic symmetry within a pseudo-orthorhombic crystal system. The distortion is described by a disparate rotational property around the chiral centers (\(\varphi_{\text{chiral}} \simeq 129^\circ\) and \(58^\circ\)) of the two molecules in the asymmetric unit. Each of these molecules comprises planar biphenyl moieties (\(\varphi_{\text{biphenyl}} = 0^\circ\)). Using temperature-dependent single-crystal X-ray diffraction experiments we show that the compound undergoes a phase transition below \(T \sim 124\, \text{K}\) that is characterized by a commensurate modulation wavevector, \(\mathbf{q} = 0, 1, 2\). The (3+1)-dimensional modulated structure at \(T = 100\, \text{K}\) suggests that the phase transition drives the biphenyl moieties towards noncoplanar conformations with significant variation of internal torsion angle (\(\varphi_{\text{biphenyl}} \leq 20^\circ\)). These intramolecular rotations lead to dimerization of the molecular stacks that are described predominantly by distortions in intermolecular tilts (\(\beta_{\text{max}} \leq 20^\circ\)) and small variations in intermolecular distances (\(\Delta d_{\text{max}} \simeq 0.05\, \text{Å}\)) between biphenyl molecules. Atypical of modulated structures and superstructures of biphenyl and other polyphenyls, the rotations of individual molecules are asymmetric (\(\Delta \varphi_{\text{biphenyl}} \approx 5^\circ\)) while \(\varphi_{\text{biphenyl}}\) of one independent molecule is two to four times larger than the other. Crystal-chemical analysis and phase relations in superspace suggest multiple competing factors involving intramolecular steric factors, intermolecular \(\text{H} \cdots \text{C} \cdots \text{C} \cdots \text{H}\) contacts and weak \(\text{C} \cdots \text{H} \cdots \text{O}\) hydrogen bonds that govern the distinctively unequal torsional properties of the molecules.

1. Introduction

Molecular biphenyl has been investigated extensively for its stability and conformation in different thermodynamic states. At ambient conditions, the differences in the conjugation states of the \(\pi\) electrons are governed primarily by the twist about the central \(\text{C} \cdots \text{C}\) single bond in the order \(40^\circ \sim 45^\circ\) in gas phase, \(20^\circ \sim 25^\circ\) in solution and \(0^\circ\) (mutually coplanar) in solid state in centrosymmetric monoclinic space group \(\text{P2}_1/\alpha\) (Bastiansen, 1949; Suzuki, 1959; Trotter, 1961; Hargreaves & Rizvi, 1962).

The planar conformation due to constraints from intramolecular interactions is energetically unfavorable and steric hindrance between the ortho hydrogen atoms is compensated for by out-of-plane dynamic disorder and in-plane displacements of those hydrogen atoms away from each other (Hargreaves & Rizvi, 1962; Casalone et al., 1968; Charbonneau & Delugeard, 1976; Charbonneau & Delugeard, 1977; Busing,
Temperature dependence of unit-cell parameters and the components of the modulation wavevector, $\sigma_1$ and $\sigma_3$.

See Table S1 in supporting information for reflections used.

| $T$ (K) | $a$ (Å) | $b$ (Å) | $c$ (Å) | $\beta$ (°) | $\sigma_1$ | $\sigma_3$ | $V$ (Å$^3$) |
|---------|---------|---------|---------|------------|-----------|----------|-----------|
| 200$^\dagger$ | 5.0560 (3) | 8.6622 (4) | 42.242 (3) | 90.349 (4) | 1850.00 (18) |
| 160 | 5.0479 (2) | 8.6330 (4) | 42.1525 (15) | 90.513 (3) | 1836.87 (13) |
| 150 | 5.0480 (7) | 8.6161 (8) | 42.1201 (11) | 90.713 (11) | 1827.6 (6) |
| 140 | 5.0484 (6) | 8.6093 (7) | 42.145 (10) | 90.720 (9) | 1826.5 (5) |
| 130 | 5.0511 (3) | 8.6014 (8) | 42.113 (11) | 90.723 (11) | 1826.9 (6) |
| 128 | 5.0446 (6) | 8.5992 (7) | 42.114 (10) | 90.720 (9) | 1826.5 (5) |
| 126 | 5.0440 (6) | 8.5978 (7) | 42.100 (10) | 90.733 (9) | 1825.7 (5) |
| 122 | 5.0443 (6) | 8.5978 (7) | 42.100 (10) | 90.733 (9) | 1825.7 (5) |
| 116 | 5.0432 (6) | 8.5926 (7) | 42.087 (9) | 90.775 (9) | 1823.6 (5) |
| 114 | 5.0422 (6) | 8.5923 (7) | 42.090 (10) | 90.787 (10) | 1823.3 (5) |

$^\dagger$ Sasmal et al. (2019a).

Twisting between the rings has been demonstrated to regulate conductivity of single molecule biphenyl–dithiol junctions (Vonlanthen et al., 2009; Mishchenko et al., 2010; Bürkle et al., 2012; Jeong et al., 2020), tune thermopower as a function of the twist angle (Bürkle et al., 2012), degeneracy of energy states on substrates (Cranney et al., 2007) and theoretically suggest wide band gap semiconducting properties of its derivatives (Khatua et al., 2020). On the other hand, biphenyl derivatives have also been reported to influence and increase the efficiency of photophysical properties (Oniwa et al., 2013; Wei et al., 2016).

Planar biphenyl molecules in the solid state favor maximum intramolecular conjugation of $\pi$ electrons as well as increasing the probability of interactions between delocalized electrons that could aid in optimal stacking of molecules.

A coupling reaction mechanism (Seechurn et al., 2012) was successfully employed to synthesize 4-biphenylcarboxy protected amino acid esters of $\alpha$-serine, $\alpha$-tyrosine, $\alpha$-alanine, $\alpha$-leucine and $\alpha$-phenylalnine via the formation of peptide-type linker O=C–NH groups (Sasmal et al., 2019b,a).

In the solid state, the compounds crystallize either in noncentrosymmetric space group $P2_12_12_1$ or the monoclinic subgroup $P2_1$ (Sasmal et al., 2019b,a). Crystal packing in these systems is determined by $\pi\cdots\pi$ stacking between the biphenyl fragments and strong linear hydrogen bonds between the amino acid ester moieties.

We presumed that the biphenyl moieties in these chemically coupled systems could influence the bioactive amino acid esters and vice versa with respect to evolution or suppression of translational and rotational degrees of freedom in their crystal structures at some thermodynamic condition.

Reanalyzing all their crystal structures, the system of 4-biphenylcarboxy-($\alpha$)-phenylalaninate attracted our attention because the structure appeared to be similar to the $\alpha$-tyrosine analog albeit the monoclinic distortion (Table 1, Sasmal et al. 2019a) and two crystallographically-independent formula units $[Z = 2$ (Steed & Steed, 2015), Fig. 1(a)] in the crystal structure of the former.

The torsion angle about the chiral center is significantly different for the independent molecules while the remainder of the rotations are similar [Fig. 1(a), (Sasmal et al. 2019a)].

Each of these molecules consists of coplanar biphenyl moieties which are stacked along $a$ and $b$, while the amide groups are connected by intermolecular $N\cdots H\cdots O$ hydrogen bonds [Fig. 1(b), Sasmal et al. 2019a)].

In the present study, the temperature-dependent phase transition of 4-biphenylcarboxy-($\alpha$)-phenylalaninate has been investigated using single-crystal X-ray diffraction experiments. Low-temperature phase II is found to be a $2a \times b \times 2c$ superstructure of the high-temperature (phase I) structure.
The superstructure is described within the (3 + 1)D-super-space approach as a commensurately modulated structure (de Wolff, 1974; Janner & Janssen, 1977; Wagner & Schönleber, 2009; van Smaalen, 2012; Janssen et al., 2018).

Structural properties of phase I and the modulated structure have been tabulated and compiled within t-plots (t = phase of the modulation). The origin and stability of phase II is discussed in terms of intramolecular steric factors and intermolecular HC⋯CH contacts and intermolecular hydrogen bonds. It is suggested that the order parameter of the phase transition is correlated with the suppression of dynamic disorder.

2. Experimental

2.1. Temperature-dependent single-crystal X-ray diffraction

Single crystals of the compound used in this study were obtained from those reported in Sasmal et al. (2019a). The crystals were protected in oil under mild refrigeration. Single-crystal X-ray diffraction (SCXRD) experiments were performed on an Agilent SuperNova, Eos diffractometer employing Cu Kα radiation. The temperature of the crystal was maintained by an Oxford Cryosystems open flow nitrogen cryostat.

During cooling, visual inspection of diffraction images revealed weaker reflections in addition to strong reflections at low temperatures.

Diffraction images collected at 150 K, 140 K and 130 K–114 K in steps of ΔT = 2 K showed that the weaker diffuse features appear at 124 K and condense into satellite reflections at 122 K (Table 1 and Fig. S1 in supporting information). The transition temperature is significantly higher than that of molecular biphenyl (Tc, biphenyl ≈ 42 K). On the other hand, related polyphenyls p-terphenyl and p-quarterphenyl undergo phase transition towards superstructure phases at much higher critical temperatures [Tc, terphenyl ≈ 190 K (Yamamura et al., 1998), Tc, quarterphenyl ≈ 233 K (Saito et al., 1985)]. Complete diffraction data were collected at T = 160 K and 100 K.

Determination of unit-cell parameters and data reductions were performed using the software suite CrysAlisPro (Rigaku Oxford Diffraction, 2019) (Tables 1 and S1).

Satellite reflections of first order (m = 1) observed below Tc could be indexed with modulation wavevector q = (σ1, 0, σ3), σ1 = σ3 = 1/2 with respect to the basic monoclinic crystal system. Here, q = 1/2(101) is perpendicular to the b axis consistent with monoclinic symmetry while in molecular biphenyl qI violates monoclinic symmetry and qII is parallel to b (Cailleau et al., 1979).

Using the plugin program NADA (Schönleber et al., 2001) in CrysAlisPro, deviations of the σ values as a function of T

Figure 1

(a) Two independent formula units of 4-biphenylcarboxy-(l)-phenylalaninate (C23H21NO3) with atomic labels of non-hydrogen atoms (a and b for molecule A and B, respectively) in phase I at T = 160 K. ϕA = -130.1°, ϕB = 56.1°; |ψA| = 32.8°, |ψB| = 30.8°; ϕA = ϕB = 0°; ψA = 36.4°, ψB = 36.2°. Viewing direction along [001]. (b) Crystal packing in phase I viewed along [111] emphasizing biphenyl stacks (AA) and (BB), along a (horizontal arrows), (ABAB), along b (zigzag vertical arrows) and intermolecular N–H⋯O bonds between amide groups (dashed orange) along [100] directions. Phenyl rings of the ester groups (transparent) are stacked only along a.
from a rational value of 0.5 were found to be within their standard uncertainties (Table 1), indicating a commensurate nature of the modulation.

Reflections at $T = 100$ K were indexed by four integers ($hklm$) using a basic monoclinic $b$-unique lattice (Tables 1 and S1) and modulation wavevector, $\mathbf{q} = (\frac{1}{2}, 0, \frac{1}{2})$ and data integration was performed. Empirical absorption correction was performed using the AbsPack program embedded in CrysAlisPro.

The ratio of the average intensities ($\langle I \rangle$) between main and satellite reflections is 13:1 and that of their average significance ($\langle \langle I \sigma(I) \rangle \rangle$) is 3:1. This indicates pronounced modulation which is characteristic of modulated molecular crystals (Schönleber & Chapuis, 2001; Schönleber et al., 2003; Dey et al., 2016, 2018; Rekis et al., 2020, 2021).

The monoclinic crystal system in addition to the reflection conditions suggest the superspace group $P2_1(\sigma_1, 0, 0, 0)$ with $\sigma_1 = \sigma_3 = \frac{1}{2}$ (Stokes et al., 2011; van Smaalen et al., 2013).

2.2. Structure refinement of the modulated structure

The crystal structure of the room-temperature phase (phase I hereon) was redetermined at 160 K using Superflip (Palatinus & Chapuis, 2007) and refined using Jana2006 and Jana2020 (Petříček et al., 2014).

Atoms were renamed with suffixes a and b for the two independent molecules A and B [Fig. 1(a)]. Anisotropic atomic displacement parameters (ADPs) of all non-hydrogen atoms were refined. Hydrogen atoms were added to carbon and nitrogen atoms using a riding model in ideal chemical geometry with constraints for isotropic ADPs [$U_{iso}(H) = 1.2U_{eq}(N)$, $U_{iso}(H) = 1.2U_{eq}(\text{aromatic})$ and $U_{iso}(H) = 1.5U_{eq}(\text{sp}$)]

Owing to the pseudorhombic crystal system, the integrated data was tested for twinning employing twofold rotation along the [100] direction as twin law. This twin law is a true symmetry element in the case of a hypothetical orthorhombic crystal system with point group symmetry 222 (Petříček et al., 2016; Nepolo, 2019). The fit of the structure model improved (compare $R_F^{\text{obs}} = 0.0463$ to 0.0408) and volume of the second component refined to 0.0240 (8) (Table S2). Finally, positions of the H atoms of NH groups and the parameter corresponding to isotropic extinction correction were refined that further improved $R_F^{\text{obs}}$ values ($R_F^{\text{obs}} = 0.0393$, Table S2 in supporting information).

The crystal structure reproduced the values for intramolecular rotations reported those for the structure at $T = 200$ K [$\psi_{\text{chir}} = \psi_1$ (hereon) and $\psi$ in Fig. 1(a)]. In addition, we also observe that the coplanar biphenyl rings are significantly rotated with respect to the amide groups [at $T = 200$ K: $\psi_2 = 32.8^\circ$ and 31.2$^\circ$ (Sasmal et al., 2019a) and at $T = 160$ K in Fig. 1(a)] which also remains invariant as a function of temperature.

The modulated structure of phase II at 100 K was refined using Jana2006 and Jana2020. Fractional coordinates of all atoms from the crystal structure at $T = 160$ K were used in the starting model while retaining the same riding model geometry for hydrogen atoms as in phase I and the average structure was refined as main reflections. In successive steps, an incommensurate (IC) model described by one harmonic wave for displacive modulation describing the atomic modulation functions (AMFs) and basic parameters for anisotropic ADPs for non-hydrogen atoms was refined against main and satellite reflections that resulted in good fit to the diffraction pattern ($R_F^{\text{obs}} = 0.0425$). However, ADPs of four non-hydrogen atoms were found to be non-positive definite.

Since the components of $\mathbf{q}$ ($\sigma_1$ and $\sigma_3$) are rational, three commensurately modulated structures were pursued by fixing the initial phase of the modulation to values $t_0 = 0$, $\frac{1}{2}$ and $\frac{1}{3}$, respectively. While the former two $t_0$ values describe monoclinic space group $B2_1$ symmetry for the equivalent 3D $2a \times b \times 2c$ superstructure, the third corresponds to triclinic $B1$ symmetry. The commensurately modulated structure (C) model corresponding to $t_0 = \frac{1}{2}$ resulted in the best fit to the diffraction data ($R_F^{\text{obs}} = 0.0426$) including ADPs of all atoms positive definite.

As the atomic modulation functions (AMFs) have sinusoidal character, the residual values are similar to the IC model (Fig. 2, Figs. S2–S4 and Table S2 in supporting information). However, the C model at $t_0 = \frac{1}{3}$ is described with either cosine or sine waves for the AMFs (equal to number of...
refinable fractional coordinates in the equivalent superstructure) reducing significantly the number of refinable parameters as compared with the IC model (compare \( N_e = 649 \) with \( N_{IC} = 811 \), further tests in supporting information).

The final C model was further improved by refining the parameter corresponding to isotropic extinction correction and AMFs and positions of hydrogen atoms of NH groups (\( R_F^{obs} = 0.0419 \), Table S2). The refined twin volume in phase II reproduced the value similar to that in phase I \([T = 100 \text{ K}, \text{twvol}_2 = 0.0242 (7) \text{ in Table S2}]. \) Presumably, the crystal possesses pseudo-merohedral growth twins.

3. Results and discussion

3.1. Structural phase transition and unequal distortion of molecules

In the present case, the monoclinic symmetry is retained below \( T_c \) unlike monoclinic to triclinic distortion at the disorder–order phase transition of p-terphenyl (Rice et al., 2013) and p-quarterphenyl (Baudour et al., 1978).

In the final commensurately modulated structure model with \( l_0 = \frac{1}{2} \), sections corresponding to \( t = \frac{1}{2} \) and \( \frac{1}{2} \) (Figs. 3 and S5) are physically relevant. These sections represent the atomic coordinates in the equivalent twofold superstructure in 3D (Figs. 4, S6 and S7).

Crystal structures of phase I and phase II have group–subgroup relations and the doubling of the \( a \) and \( c \) axes describes the additional \( B \)-centering of the superstructure in phase II.

The superstructure derived using Jana2006 comprises four molecules in the asymmetric unit \((Z = 4)\); two each corresponding to molecules \( A \) and \( B \) of phase I (Fig. 4).

The covalent bond distances are similar for the independent set of molecules and are practically unaffected by modulation Table S7).

In the present study, discussion is based on the modulated structure in order to establish unique relations between phase I and phase II (Rekis et al., 2021; Chapuis, 2020; Ramakrishnan et al., 2019; Dey et al., 2016; Noohinejad et al., 2015; Scho¨nleber, 2011; Schönleber et al., 2003).

The modulated structure suggests that the phase transition is dominated by evolution of internal torsional degrees of freedom \((\varphi^d > 0^\circ)\) within the biphenyl moieties [Fig. 3(a)]. The twists about the central C–C bond are significantly different for the two molecules where the torsional modulation of \( A \) are 2–4 times larger than those of \( B \) (dihedral angles \( |\varphi^B_A| = 15.6^\circ, 20.5^\circ \) and \( |\varphi^B_B| = 4.1^\circ, 9.3^\circ \)). These distortions are described by highly anisotropic AMFs \((\alpha)\) along the three basis vectors where the maximum amplitude are along \( b \) for the carbon atoms of biphenyl (Fig. 2 and Table S4). Notably, the rotations in the present structure are significantly larger than those reported for molecular biphenyl \([\varphi \simeq \pm 5.5^\circ] \) (Petricek et al., 1985; Baudour & Sanquer, 1983]). These values are smaller than those in the low-temperature superstructure of \( p \)-terphenyl and \( p \)-quarterphenyl \([\text{maximum } \varphi_{\text{terphenyl, quarterphenyl}} \simeq 23^\circ] \) (Rice et al., 2013; Baudour et al., 1976, 1978)].

Figure 3
\( t \)-plots of intramolecular rotations of molecule \( A \) (blue) and \( B \) (red) as well as intermolecular tilts and distances between stacks of 4-biphenylcarboxy-(-)-phenylalaninate. (a) Dihedral angle \( |\varphi| \) represents internal torsion within the biphenyl rings, (b) dihedral angle \( |\varphi| \) represents the torsion angle between the inner ring of the biphenyl and the amide groups, (c) \( |\varphi| \) represents the torsion angle of the amide groups with respect to the –COOCH\(_3\) groups and (d) \( |\varphi| \) represents the torsion angle of –COOCH\(_3\) groups with respect to amide groups. (e) \|\theta_{ABBB}\|\) represents tilt between biphenyl rings of \( A \) and \( A^d \) (red), and of \( B \) and \( B^d \) (red) and \( \theta_{ABBB} \) (dashed-dotted black curve) represent tilt between inner aromatic rings of biphenyl (bonded to amide groups) of \( A \) and outer ring of \( B \) and vice versa. (f) Intermolecular distances (\( \delta \)) between biphenyl rings of \( A \) and \( A^d \) (blue), between those of \( B \) and \( B^d \) (red) and between those of \( A \) and \( B \) (black). Horizontal dashed lines represent those angles and distances in phase I \( (|\varphi| = |\theta_{ABBB}| = 0^\circ) \). Vertical dashed lines indicate \( t \) values corresponding to angles and distances in the 3D superstructure. Symmetry code (ii): \( x + 1, y, z, t \).

Figure 4
Comparison of structures in phase I and phase II across the phase transition highlighting the effect of internal torsion \((\varphi^d)\) within biphenyl on the stacking arrangements along \( a \). The tilt between the biphenyl stacks, \( \theta_{ABBB} \) are different for the inner rings (bonded to amide rings) and the outer rings. Corresponding values of \( \varphi^d \) and \( \theta \) are in Fig. 3(e) and those for C–H···O and C–H···H–C in Table 2. See full unit cells in Figs. S6 and S7. Viewed along [001].
The nature of structural changes in the present system and molecular biphenyl below the phase transition temperature is different to \( p-(n > 2) \)-phenyl systems described by the property that in the later cases two disordered conformations of the molecules freeze by superstructure formation and breaking the monoclinic symmetry of their high-temperature phase.

A distinctive property of the present modulated structure is the unequal modulation for the two different moieties where \( u_{biphenyl} > u_{phenylalaninate} \) (Table S4). Compared to \( |\varphi_A| \), the distortions in torsion angles \( \varphi \) are lesser and those in \( \psi \) are very small and virtually equal for both \( A \) and \( B \) \( |\varphi_A| = -133.7^\circ, -127.6^\circ; |\varphi_B| = 55^\circ, 56.2^\circ; |\psi_A| = 36.8^\circ, 37.7^\circ; |\psi_B| = 36^\circ, 36.6^\circ \) in Figs. 3(c) and 3(d).

A possible reason for the weaker modulations of the atoms around the chiral centers is the directional strong intermolecular \( N-H \cdots O \) bonds makes large intramolecular rotations unfavorable.

Note that the observed changes in the rotations of \( \varphi \) of molecule \( A \) [compare \( \varphi_A = 39.1^\circ, 25.8^\circ \) with \( \varphi_B = 32.9^\circ, 28.9^\circ \) in Fig. 3(b)] are predominantly described by strong modulations of the molecule’s biphenyl moiety. The asymmetry in rotations of individual molecules (\( \Delta \varphi \approx 5^\circ \)) is determined by the disparate bonding environments of the biphenyl moieties where the inner rings are that are covalently bonded to amide groups while the outer interact weakly via \( C-H \cdots H-C \) interactions with the phenyl rings of phenylalaninate groups (Fig. 4). Subsequently, the unequal values at the relevant \( t \)-sections of \( \varphi \) are correlated with those of \( \varphi \) [compare Figs. 3(a) and 3(b)].

It is also observed that the variation in \( \varphi \) is greater for molecule \( A \) than that of \( B \) [compare \( \Delta \varphi \approx 6^\circ \) with \( \Delta \varphi \approx 1^\circ \) in Fig. 3(c)]. The origin of disparate distortions in intramolecular rotations is explained in §3.2.

In the modulated structure, the biphenyl moieties in (AA)\(_n\) and (BB)\(_n\) stacks which are parallel (\( \theta = 0^\circ \)) in phase I are tilted with respect to each other [Fig. 3(e)]. These tilts (\( \theta_{AA/BB} \)) are of the order of the internal twists (\( \varphi \)) of the independent biphenyl moieties \( \theta_{AA} = 19.5^\circ \) and \( 16.6^\circ \); \( \theta_{BB} = 5^\circ \) and \( 7.2^\circ \) for inner and outer rings of biphenyl, respectively, compare Fig. 3(e) with Fig. 3(a)]. The orientation between the biphenyl moieties within the (ABAB)\(_n\) stacks also vary with \( \Delta \theta_{AA/BB} \approx 12^\circ \) where the value is intermediate to \( \varphi_A \) and \( \varphi_B \) [compare Fig. 3(e) with Fig. 3(a)]. In addition, intermolecular distances between the biphenyl moieties within the stacks at the two \( t \)-sections are different and vary up to \( \Delta d_{AA/BB} \approx 0.05 \) Å and \( \Delta d_{AB/BB} \approx 0.02 \) Å [Fig. 3(f)].

Overall distortions in \( d_{AA} \) and \( d_{BB} \) are nearly equal although \( \theta_{AA} \) is greater than \( \theta_{BB} \). These variations in \( d \) may arise to compensate for the mutual rotations of aromatic rings within the stacks. For example, the comparison \( \theta_{AA} \) for outer rings > \( \theta_{AA} \) for inner rings as opposed to \( \theta_{BB} \) for outer rings < \( \theta_{BB} \) for inner rings could explain the complimentary variations of (AA)\(_n\) and (BB)\(_n\) [sets of distances (Å) in \( t = (1/2) \): \( d_{AA} = (5.01, 5.06) \); \( d_{BB} = (5.06, 5.02) \) in Fig. 3(f)]. It could therefore be argued that the dimerization of biphenyl molecular stacks below \( T_c \) is predominantly governed by distortion described by molecular rotations rather than intermolecular distances. On the other hand, variation of intermolecular distances between aromatic rings of 1-phenylalaninate are similar to those of the biphenyls albeit the interstack rotations of the former are significantly smaller (\( \theta < 3^\circ \), Fig. S5 in supporting information).

### Table 2
Comparison of nonbonded hydrogen-\( \cdot \)acceptor and hydrogen-\( \cdot \)hydrogen distances (Å) involved in hydrogen bonds and steric factors in phase I (\( T_1 \)) and phase II (\( T_2 \)).

| Interaction class | Atom labels   | Phase | Distance (Å) |
|------------------|--------------|-------|--------------|
| N-H-O            | H1n1a-\cdots-O3a | I     | 2.00         |
|                  | H1n1b-\cdots-O3b | II    | 1.97, 1.95   |
| C-H-O            | H1c10a-\cdots-O2a | II    | 2.72         |
|                  | H1c10b-\cdots-O2b | II    | 2.78         |
|                  | H1c14a-\cdots-H1c19a | II    | 1.98         |
|                  | H1c14b-\cdots-H1c19b | II    | 1.98         |
|                  | H1c16a-\cdots-H1c23a | II    | 2.03         |
|                  | H1c16b-\cdots-H1c23b | II    | 2.02, 2.06   |
|                  | H1c13a-\cdots-O3a | II    | 2.59         |
|                  | H1c13b-\cdots-O3b | II    | 2.59         |
|                  | H1c3a-\cdots-O3a | II    | 2.37         |
|                  | H1c3b-\cdots-O3b | II    | 3.65         |
| Inter H-H        | H1c21a-\cdots-H1c7b | I     | 2.59         |
|                  | H1c21b-\cdots-H1c7a | II    | 2.45         |

3.2. Competitive forces governing modulations

Structural studies in the 3D phase of molecular biphenyl have suggested that the ortho-hydrogen atoms are displaced away in the plane of the rings to minimize steric hindrance (Trotter, 1961; Hargreaves & Rizvi, 1962; Charbonneau & Delugeard, 1976). On the other hand, dynamic disorder predominantly governed by torsional vibrations around the long molecular axis (Petricek et al., 1985) is predicted to balance the planar conformation of biphenyl favorable for crystal packing (Lenstra et al., 1994).

As short as 1.98 Å in phase I (Table 2), these contacts are shorter than the predicted values for twice van der Waals radius for hydrogen \( r = 1.1-1.2 \) Å (Rowland & Taylor, 1996; Alvarez, 2013). In the modulated structure, we observe that the distances between the ortho-hydrogen atoms are marginally but consistently larger than those in phase I (Table 2) that could suggest that the torsional modulations aid in minimization of the presumed steric hindrance below \( T_c \) (Dey et al., 2022, 2018).
A peculiar property of the modulated structure under discussion is the significant difference in the torsional amplitude of φ3 of the independent molecules. This aspect cannot be explained solely based on the intramolecular steric factors. Analysis of the crystal packing shows that each of these independent biphenyl moieties maintains close intermolecular CH⋯HC contacts with the phenyl rings of L-phenylalaninate in AB and BA fashion (Fig. 4).

These distances are significantly longer (intermolecular dH⋯H ≥ 2.4 Å, Table 2) compared with the intramolecular H⋯H distances. On the other hand, the aromatic rings of L-phenylalaninate interact with adjacent oxygen atoms of −COOCH3 via C−H⋯O hydrogen bonds (Fig. 4 and Table 2). These hydrogen bonds are weaker (Desiraju & Steiner, 2001) but highly directional [\( \angle(C−H⋯O) = 159−164^\circ \)] with very little variation in the distances.

Interestingly, those H⋯H distances involving biphenyl moieties of molecule B are consistently smaller than those of molecule A in both phases (Table 2). We argue that in the presence of both the van der Waals interactions and weak C⋯H⋯O bonds, the larger distortions of A is favored by weaker CH⋯HC interactions while that is suppressed in B.

The variations in \( \phi^2 \) and asymmetry in \( \phi^3 \) can be explained with respect to the intramolecular nonbonded nearest distances between the hydrogen atoms bonded to C13 atoms (Fig. 1) of inner rings of biphenyl and the oxygen atoms O3 of the amide groups. The distances are short (d_{H13−O3} = 2.59 Å in phase I, Table 2) and are in the range of the sum of the van der Waals radii of oxygen and hydrogens \( [r_H = 1.1−1.2, r_O = 1.4−1.56 \text{ Å}] \) (Rowland & Taylor, 1996; Alvarez, 2013). In the modulated structure, the larger distortions of \( \phi^3 \) requires the amide groups to rotate with respect to the inner phenyl ring to optimize these C⋯H⋯O contacts to avoid steric effects (compare d_{H13−O3} = 2.66 Å, 2.54 Å in Table 2 with \( \phi^2 = 39.1^\circ, 25.8^\circ \) in Fig. 3). Although the variation in \( \phi^3 \) is smaller, the positive correlation that is larger rotations with greater C⋯H⋯O distances and vice versa are observed (compare d_{H13−O3} = 2.61 Å, 2.56 Å in Table 2 with \( \phi^3 = 32.9^\circ, 28.9^\circ \) in Fig. 3). Alternatively, it could be argued that the asymmetric distortion of \( \phi^3 \) within individual molecules is a result of the constraint of the amide groups, where rotations are hindered when the C⋯H⋯O contacts are short and favored when those are longer. Therefore, the four different values of intramolecular torsion \( \phi^3 \) within the biphenyl moieties are distinctly governed by intramolecular nonbonded H⋯H and H⋯O; and intermolecular nonbonded H⋯H contacts and weak C⋯H⋯O hydrogen bonds.

The independent molecules A and B differ from each other with respect to their torsion around the chiral center namely \( \phi^3 = (−130.1^\circ \text{ and } 56.1^\circ \) for A and B, respectively, in phase I). The difference in twist angles has remarkable effects on the molecular conformation with respect to the distance between the O3 atoms of the amide groups and hydrogen atoms of the chiral centers C3 carbon atoms (d_{H13−O3} = 2.37 Å, 3.65 Å for A and B, respectively, in phase I, Table 2). In phase II, the modulations of L-phenylalaninate moieties are weaker than that of the biphenyl groups possibly due to the presence of strong intermolecular N⋯H⋯O hydrogen bonds. However, the variations in \( \phi^2 \) ( ≥6°) as compared with \( \phi^3 \) ( ≥1°) are greater which could arise to optimize significantly shorter H⋯O nonbonded contacts of A as compared with B (d_{H1−O3} = 2.38 Å, 3.65 Å for A and B, respectively, in phase II, Table 2).

As noted in §2.2 the average intensities of the satellite reflections are an order of magnitude smaller than the main reflections. The order parameter of the phase transition might be expressed as proportional to the amplitude of the modulation which is approximately proportional to square root of the intensities of the satellite reflections (van Smaalen, 2005). The pronounced AMFs of the biphenyl moieties describing the predominant distortions in the crystal structure are accompanied by suppression of dynamic disorder in phase II as compared with phase I. For example, the carbon atoms at ortho (C14, C16, C19, C23) and meta (C13, C17, C20, C22) positions are strongly displaced [Fig. 5(b), Tables S4 and S5]. Subsequently, the ADPs are significantly reduced as compared with phase I [Fig. 5(a), Table S5]. Notably, the decrease of the ADPs \( (u_{eq}) \) from \( T = 160 \text{ K} \) to \( T = 100 \text{ K} \) is larger for those of molecule A than those for B in conjunction with the fact that overall the ADPs are smaller or similar for the former as opposed to phase I, while the square of the amplitude of modulations \( (\mu^2) \) are greater for A than those for B [compare Figs. 5(b) with 5(a)].

The decrease in ADPs of atoms of L-phenylalaninate moieties of A are greater than B from \( T = 160 \text{ K} \) to \( T = 100 \text{ K} \) [Fig. 5(c)]. Unlike the biphenyl moieties, such apparent switch-over of the ADPs is not observed where the overall values of A are greater than those of B in both phases I and II, while the squares of the AMFs in II are larger for A than those for B.

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**Figure 5**

Scatter plots of equivalent value of anisotropic ADPs \( (u_{eq}) \) and square of the amplitude of modulations \( (\mu^2) \) of the carbon atoms of biphenyl moieties (C12 through to C23) and non-hydrogen atoms of L-phenylalaninate moieties [see Fig. 1(a)] of molecules A (blue) and B (red). (a) and (c) \( u_{eq} \) of the biphenyl carbon atoms and L-phenylalaninate non-hydrogen atoms, respectively, at \( T = 160 \text{ K} \) (open circles) and at \( T = 100 \text{ K} \) (full circles). (b) and (d) \( \mu^2 \) (diamonds) of the corresponding biphenyl carbon atoms and L-phenylalaninate non-hydrogen atoms of molecules A and B, respectively, at \( T = 100 \text{ K} \). See Tables S5 and S6.
possibly arises to optimize the short nonbonded H···O angles in the former than the later. This hypothesis is supported by the small but observable distortion of \( \phi_A \) (\( \sim 6^\circ \)) as compared with \( \phi_B \) (\( \sim 1^\circ \)) while those for \( \psi \) are significantly smaller (\( \sim 1^\circ \)) for both molecules in phase II.

4. Conclusions

The 2\( a \times b \times 2c \) superstructure of 4-biphenylcarboxy-(l-)-phenylalaninate at \( T = 100 \) K has been successfully described as a commensurately modulated structure within (3+1)D superspace with superspace group symmetry \( P2_1(\sigma_1)P6_3\). The single crystal to single crystal phase transition below \( T = 124 \) K drives the 3D structure directly to a locked-in twofold superstructure accompanied by significant distortion of torsional rotations within biphenyl away from coplanarity that is also a property of incommensurately modulated structure of biphenyl but with amplitudes four times smaller than the present system.

The phase transition temperature is significantly higher than that in biphenyl yet significantly lower than for \( p \)-terphenyl and \( p \)-quarterphenyl. Consistent with the \( T_c \), the maximum amplitude of torsion is also intermediate and in the order \( \phi_{\text{quarterphenyl}} > \phi_{\text{terphenyl}} > \phi_{4\text{-biphenylcarboxy-l-phenylalaninate}} > \phi_{\text{biphenyl}} \).

Topologically separated, conformations of both the weaker C—H···O bonds and stronger N—H···O bonds are rigid and that underlines their role in stabilizing the crystal packing in both phases. A unique property of the present polyphenyl coupled amino acid ester is the distinctively unequal torsional amplitude (\( \phi_A > \phi_B \)) within the independent molecules which is governed by multiple level of competitions involving unequal van der Waals constraints in the presence of weak hydrogen bonds between the biphenyl and l-phenylalaninate moieties while the asymmetry of \( \phi \) is determined by intramolecular nonbonded constraints between phenyl rings and amide groups.

The unusual nature of the phase transition is rationalised by the fact that unequal intramolecular distortions of the two molecules are complemented by unequal suppression of the dynamic disorder of their atoms below \( T_c \).

The present investigation of the phase transition in the biphenylcarboxy coupled amino acid ester system also shows consequences for crystal packing where significant distortions in conjugations represented by mutual intramolecular rotations between homo aromatic groups as well as with aliphatic amide groups results in modulated \( \pi \cdots \pi \) stacking arrangements (\( \theta_{AABB} > 0^\circ \)) of phenyl groups but preserves the conformations of intermolecular directional N—H···O hydrogen bonds of the high-temperature structure.

5. Related literature

The following references are cited in the supporting information: Becker & Coppens (1974), Coelho (2003, 2018), Petříček et al. (2014).

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