Theoretical study on aromatic and open-shell characteristics of carbon nanobelts composed of indeno[1,2-b]fluorene units: dependence on the number of units and charge states†

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In this study, we theoretically investigate the aromatic and open-shell characteristics of carbon nanobelts (CNBs) composed of five- and six-membered rings. We have designed nanobelts composed of indeno[1,2-b]fluorene ([I]IF) units, which are referred to as [N]IF-CNB (N: the number of five-membered rings). The number of π-electrons, nπ, in neutral [N]IF-CNB is 7N, and thus depending on N and charge states, nπ can be 4N + 2 and 4N. Quantum chemical calculations on neutral [6]IF-CNB and [8]IF-CNB and dicaticonic [8]IF-CNB† have revealed that they are expected to exhibit unique aromatic and open-shell characteristics depending on nπ, there are several analogies of the electronic structures in [N]IF-CNB to those in [N] annulene. Delocalized and intermediate open-shell electronic structures of [N]IF-CNB are also useful to drastically change the third-order nonlinear optical properties. These results suggest that theoretically designed [N]IF-CNB can be attractive and challenging targets of organic synthesis for realizing novel open-shell functional conjugated macrocycles.

Introduction

Creation of π-conjugated oligomers with well-defined cyclic structures, i.e., conjugated macrocycles, has been one of the hot topics in materials science.1, 2 Owing to the cyclic topologies, the conjugation lengths of fully conjugated macrocycles are regarded as infinite, and this feature is reflected in their unique delocalized electronic structures. Such electronic structures are closely related to their geometries, stabilities, properties and related functionalities. Recent progress in experimental techniques have allowed us to realize a variety of conjugated macrocycles.1-11 One of the most important features of conjugated macrocycles is that there are inner and outer domains that are spatially separated from one another by the cyclic backbone. Since conjugated macrocycles can offer different electrostatic environments in the inner and outer domains, they are also considered to be useful as efficient materials for molecular recognition and host–guest chemistry.11-17 Not only planar, but also non-planar conjugated macrocycles with curved π-conjugation surface, such as Möbius-, saddle- and belt-shaped systems, are very interesting materials in this regard.11, 18 Even though synthesis of such non-planar conjugated macrocycles with well-defined curved structures is one of the most challenging topics in synthetic chemistry, there have been plenty of successes to realize such materials, including cycloparaphenylene (CPPs), extended porphyrinoids and carbon nanobelts (CNBs).1, 19

Aromaticity is an important chemical concept for characterizing the delocalized electronic structures of π-conjugated systems involving π-conjugation rings. According to Hückel rule for a planar π-conjugation ring, the number of π-electrons, nπ, can be the simplest index for characterizing its aromaticity.20 Aromatic compounds with (4n + 2)π electrons are known to exhibit characteristic thermodynamic stabilities and properties, whereas antiaromatic compounds with 4nπ electrons tend to be rather unstable. The concept of aromaticity and the classification of compounds by nπ are extensively used to characterize the electronic structures of various π-conjugated systems.21

Since the aromaticity concept is closely related to the electron occupancies and degeneracy of the frontier molecular orbitals (MOs), it is also connected to another important chemical concept concerning the electron occupancies and degeneracy, i.e., open-shell character.22,23 For example,
antiaromatic 4nπ-electron systems tend to have very small energy gap between the highest occupied and lowest unoccupied MOs (HOMO and LUMO). When the HOMO and LUMO characterize bonding and anti-bonding natures for a specific or effective chemical bond, such a system tends to have open-shell (diradicaloid) electronic structure in the singlet state. Indeed, most antiaromatic compounds are known to be highly reactive, since the electronic structures of open-shell systems are easily fluctuated by external physical and chemical perturbations. Such a feature of open-shell electronic structure is, on the other hand, expected to be useful for creation of novel stimuli-responsive functional materials, such as efficient nonlinear optical (NLO) materials, if one can increase their thermodynamic and kinetic stabilities by molecular and materials design strategies. In this regard, a theoretical index, diradical character \( y \) [0 (closed-shell) \( \leq y \leq 1 \) (pure diradical)], which characterizes the degree of open-shell of an electron pair, has been shown to be a useful index to design novel open-shell functional molecules with unique electronic, optical and magnetic response properties.

In order to realize novel open-shell functional molecules, it is very important to construct practical design strategies for tuning \( y \) by chemical modifications, such as inclusion of paraquinodimethane (pQM) substructures into the molecular skeleton in order to tune quinoid (closed-shell)/benzenoid (open-shell) resonance structure. For example, several fused polycyclic hydrocarbons involving plural five-membered rings can involve pQM substructures, and thus there are several real open-shell molecules with a variety of \( y \) values in this class. These systems are also known to exhibit a variety of antiaromatic characters. Indenfluorene (IF), where five- and six-membered rings are alternately fused, is one of the typical antiaromatic open-shell molecules (for possible resonance structure, see Fig. 1a). Depending on the topologies of ring-fusion, the regioisomers of IF are shown to exhibit different \( y \) values, and thus, several derivatives of IF are expected to be useful building units for creation of open-shell macromolecules. Indeed, Nobusue, Shimizu and Tobe reported synthesis of planar tetraacyclo[defjkl.pqr.wux]tetraphenylene (TCPTP, 28π electrons), which involves repeating indeno[2,1-c]fluorene substructures. Quantum chemical calculations combined with the experimental analysis revealed that the \( D_{2h} \) structure having almost pure diradical and weak tetraradical characters is considered to be a transition state between the almost closed-shell \( D_{2h} \) local minima. On the other hand, within the planar fully-fused conjugated macrocycles, increasing the number of repeating IF substructures is considered to be difficult because of the structural limitations.

In our previous study, we have theoretically investigated the relationship between the geometric features, electronic structures and third-order NLO properties of a non-planar conjugated macrocycle, i.e., a carbon nanobelt‡ composed of five- and six-membered rings. Here we call the system [6]IF-CN (Fig. 1, \( N = 6 \)), where \( N \) corresponds to the number of five-membered rings involved in [\( N \)]IF-CN. The molecular design of [6]IF-CN was inspired from those of indeno[1,2-b]fluorene and (6,6)CNB, the latter of which has been synthesized by Itami and coworkers. [\( N \)]IF-CN involves repeating indeno[1,2-b]fluorene substructures (see Fig. 1a), and owing to its belt-shape, we can reasonably increase the number of units without changing the repeating structures. Note that only even numbers of \( N \) are allowed in [\( N \)]IF-CN to make a reasonable cyclic structure of hoop.

One of interesting features of [\( N \)]IF-CN is the number of π-electrons, \( n_\pi = 7N \). In other words, \( n_\pi = 4n + 2 \) when \( N = 4m + 2 \) (\( m = 1, 2, \ldots \)), while \( n_\pi = 4n \) when \( N = 4m \), and thus unique aromatic characters are expected if they are fully π-conjugated. Actually, [6]IF-CN is expected to exhibit weak global aromatic character from the analysis of the magnetically induced ring-current (MIC). Analysis of the MO levels of [6]IF-CN has revealed that there is an analogy in the orbital degeneracy of [6] IF-CN to that of [6]annulene. Furthermore, [6]IF-CN was shown to exhibit an intermediate multiradical character owing to the presence of IF substructures, the feature of which is reflected in the enhancement of third-order NLO property of [6]IF-CN compared to that of closed-shell (6,6)CNB. Since [\( N \)] annulene is known to drastically change their structures, stabilities and properties by changing the number of π-electrons, the results of [6]IF-CN have stimulated us to further investigate \( n_\pi \)-dependence of electronic structures and physicochemical properties of [\( N \)]IF-CN. In this study, we therefore conduct several theoretical analyses in order to clarify the relationship between \( n_\pi \) (depending on the number of units and charge states), aromatic and open-shell characters, and NLO

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† According to the ref. 19, the term “carbon nanobelt” is used for a class of belt-shaped hydrocarbons composed only of six-membered rings. In this classification, belt-shaped compounds involving five-membered rings are classified into “aromatic belt”. However, here we call the present belt-shaped hydrocarbons involving five-membered rings, [\( N \)]IF-CN (i.e., IF-containing carbon nanobelt) as a counterpart of regular CNB, because they can exhibit both aromatic and antiaromatic characters depending on \( N \), and the term “aromatic belt” may cause confusion for readers about the aromatic characters of [\( N \)]IF-CN.
properties of [N]IF-CNBl. On the basis of the results, we will
discuss a novel design strategy for creation of open-shell func-
tional materials. In order to focus on the fundamental rela-
tionship between \( n_\pi \) and electronic structures (aromatic and
open-shell characters), we here discuss the results of neutral \( [6] 
\) IF-CNBl \( (n_\pi = 42; [4n + 2]_\pi) \), \( [8] \)IF-CNBl \( (n_\pi = 56; 4n^2) \) and
dication \( [8] \)-IF-CNBl \( (n_\pi = 54; [4n + 2]_\pi) \).

**Computational details**

All the carbon nanobelts investigated in this study are expected
to have open-shell characters in the singlet state. For such
systems, spin-unrestricted density functional theory (UDFT) are
known to sometimes give incorrect bond-length alternation
(BlA) patterns around the radical sites.\(^{30}\) Geometry optimiza-
tions followed by frequency analysis were therefore performed
at the RB3LYP/6-311G\(^*\) level of approximation, which is ex-
pected to reproduce BlA patterns for several intermediate
multiradicaloids involving five-membered rings.\(^{29}\) In this study,
because of the unique cyclic structure of \([N]IF-CNBl\), we have
carefully discussed DFT functional dependence of the opti-
mized geometry for \([N]IF-CNBl\). For details, please see the ESL.\(^{1}\)

Diradical character for the \( i \)-th natural orbital (NO) pair \( (y_i) \) was
evaluated from the occupation number of the \( i \)-th lowest-
unoccupied NO (LUNO + \( i \)). Since \( y_i \) represents the degree of
radical for a pair of two electrons in the \( i \)-th orbital, a set of \( y_i \) is
a measure of multiradical characters of the system.\(^{31,32}\) Spatial
distribution of unpaired electrons can be visualized from the
map of odd electron density distribution \( D_{y_i}^{\text{odd}}(r) \) for \( y_i \), which
is defined by the spatial distributions of the HONO-\( i \) and
LUNO+\( i \), respectively \( \{\phi_{\text{HONO}-i}(r) \) and \( \phi_{\text{LUNO}+i}(r)\) \) and \( n_{\text{LUNO}+i} \) as,\(^{32}\)

\[
D_{y_i}^{\text{odd}}(r) = n_{\text{LUNO}+i} \left( |\phi_{\text{HONO}-i}(r)|^2 + |\phi_{\text{LUNO}+i}(r)|^2 \right)
\]

where the spatial integration of \( D_{y_i}^{\text{odd}}(r) \) is related to \( y_i \) by the
following equation,

\[
y_i = \frac{1}{2} \int dr D_{y_i}^{\text{odd}}(r)
\]

If \( y_i \) and \( y_{i+1} \) are the same, the corresponding NOs for \( y_i \) and
\( y_{i+1} \) are degenerate, so that these NOs are not necessarily the
symmetry-adapted ones. We therefore need to take the sum or
the arithmetic average of \( D_{y_i}^{\text{odd}}(r) \) and \( D_{y_{i+1}}^{\text{odd}}(r) \) to illustrate the
symmetry-adapted odd-electron densities.\(^{33}\) The result of \( y_i \) is
known to depend on the employed quantum chemical calcula-
tion method.\(^{34}\) In this study, we employed a spin-unrestricted
long-range corrected exchange-correlation functional, LC-
UBLYP,\(^{35}\) using the range-separating parameter of \( \mu = 0.33 
bohr^{-1} \) along with the 6-31G* basis sets for the calculation and
analysis of diradical characters.

There are several criteria to evaluate aromatic characteristics
of a system. In order to investigate the aromaticity of the belts,
we employed gauge-including magnetically induced current
(GIMIC) method to evaluate magnetically induced current
(MIC) density.\(^{34}\) The unperturbed and magnetically-perturbed
electron densities which are needed for GIMIC calculations were
evaluated with the LC-UBLYP exchange-correlation func-
tional with a range separating parameter \( \mu = 0.33 \) bohr\(^{-1} \) and the
6-311+G** basis set. This functional was found to reproduce
the excitation energies, NLO properties such as the second
hyperpolarizability \( (\gamma) \), and magnetic shielding tensors of
a variety of open-shell singlet \( \pi \)-conjugated systems either
calculated at higher levels of theory or determined by experi-
ments.\(^{35}\) The external magnetic field was applied to the direc-
tion going through the hoop \( (z \) direction). The gauge-invariant
atomic orbital (GIAO) method was used throughout the calcu-
lation.\(^{36}\) Then, the MIC densities was evaluated by GIMIC
program. MIC density vector is evaluated at each position of
space. In order to clarify the contribution of induced current for
each bond, we evaluated bond-integrated MIC density. Here, we
set a bisection plane for two atoms \((width \times depth = 2 \ \text{Å} \times 10 \ \text{Å})\)
so that one of the edges of the plane involves the center of the
belt.

The static diagonal second hyperpolarizability tensor \( \gamma_{iii} \)
which characterizes third-order NLO property at the molecular
scale was calculated by using the finite field (FF) approach\(^{37}\)
combined with the Romberg procedure.\(^{38}\) In the Romberg
procedure, we employed the same procedure as the previous
study.\(^{39}\) For all the calculations, the 6-31+G* basis set was
adopted. The automatic Romberg differentiations were per-
formed using the T-REX program.\(^{40}\) We employed the B-
convocation for the definition of \( \gamma. \)\(^{39}\) The total energies are
evaluated by using the LC-UBLYP \( (\mu = 0.33 \) bohr\(^{-1} \) functional
with the 6-31+G* basis set. We set the molecular coordinate axis
so that \( z \)-axis is going through the hoop of the belt, which
coresponds to the short-axis direction of the belt.\(^{39}\) Then, \( x \)-axis
was set to be parallel to the line going through the centers of two
face-to-face six-membered rings on the opposite side of the belt.

In order to clarify the spatial contribution of electrons to the
second hyperpolarizability tensor \( \gamma_{iii} \), we performed the \( \gamma \)
density analysis.\(^{40}\) The \( \gamma \) density \( \rho_{\text{iii}}^{(3)}(r) \) is defined as the third-
order derivative of the electron density \( \rho(r) \) with respect to the
applied electric field \( F \):\(^{40}\)

\[
\rho_{\text{iii}}^{(3)}(r) = \frac{\partial^3 \rho(r)}{\partial F^3} \bigg|_{F=0}.
\]

The \( \gamma \) density is related to the \( \gamma \) value by the following
equation,

\[
\gamma_{iii} = -\frac{1}{3!} \int dr r^i \rho_{\text{iii}}^{(3)}(r),
\]

where \( r^i \) is the \( i \)-axis component of the electron coordinate. The
\( \gamma \) density represents the field-induced third-order response of
electron density at position \( r \). A pair of positive and negative \( \gamma 
\) densities contribute to \( \gamma \) value, the sign of which is positive
(negative) when the direction of vector drawing from positive
to negative densities coincides with the positive direction of
coordinate axis, and the amplitude of which is proportional to
the distance between the positive and negative densities.\(^{40}\) All
the calculations were carried out using the Gaussian 09
package.\textsuperscript{41} DrawMol package\textsuperscript{44} was used for the visualizations of molecular modelling and spatial distributions of MOs, MIC and $\gamma$ densities.

**Results and discussion**

At first, we discuss the local minimum structures of calculated systems. In the previous study, we found that the local minimum structure of neutral [6]IF-CNB belongs to the highest possible $D_{4d}$ symmetry. In this study, we have found two local minima with no imaginary frequencies for neutral [8]IF-CNB that belong to the $C_{4v}$ and $D_4$ symmetries, respectively. Energetically more stable (about 7 kcal mol$^{-1}$) structure is found to be the local minimum with the lower $D_4$ symmetry. The symmetry reduction of the most stable local minimum structure may be similar to the case in TCPTP to some extent, where $D_{4h}$ structure is considered to be a transition state between the almost stable $D_{2h}$ local minima.\textsuperscript{26} In the case of dicaticonic [8]IF-CNB\textsuperscript{2+}, we found one local minimum structure which belongs to the highest possible $D_{4d}$ symmetry. Reduction of symmetries in [8]IF-CNB and recovery of symmetry in dicaticionic [8]IF-CNB\textsuperscript{2+} are considered to be related to $\eta_{p}$.

Fig. 2a shows the optimized bond lengths of selected CC bonds for the most stable local minimum structure of each system. Only the results of symmetrically unique bonds are given in Fig. 2a. One of the important points is the alternation pattern around the bonds 5 and 6 that belong to the five-membered ring, since it reflects the tendency of quinoid/benzenoid resonance of $pQM$ substructures involved in the belt. [6]IF-CNB and [8]IF-CNB\textsuperscript{2+} are shown to exhibit BLA-less features for bonds 5 and 6, whereas [8]IF-CNB is found to exhibit a BLA pattern for these bonds. When we focus on the BLA patterns in the six-membered ring parts, bonds 1 vs. 3 (4 vs. 7) are shown to be equivalent in [6] IF-CNB and [8]IF-CNB\textsuperscript{2+}, and thus, all the six-membered rings involved in the belt are considered to be equivalent in the sense of the resonance structure. On the other hand, these CC bonds are not equivalent in [8]IF-CNB, which means that not all the six-membered rings are equivalent. Judging from the bond lengths of 1 and 7, which are closer to 1.4 Å than those of 3 and 4, there are rather quinoid- and rather benzenoid-like six-membered rings in [8]IF-CNB, and these two-types of six-membered rings appear alternately in the belt.

In order to further examine the quinoid- and benzenoid-like characters of the geometries, we also examined distances between the vertices of adjacent five-membered rings (marked with stars in Fig. 2a). If there is a quinoid-like structure, the corresponding distance between the vertices is expected to become shorter than that for the benzenoid-like structure, \textit{i.e.}, we discuss the alternation patterns of this distance (see Fig. 2b). From the calculation results, averages of the distances are found to increase about 0.1 Å when we increase the number of

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**Fig. 2** Comparison of bond lengths of selected CC bonds (a), and of distances between the vertices of adjacent five-membered rings (b). For (b), the numbering of the position is arbitrary because of the high symmetry of each system.

**Fig. 3** Closed-shell contributions of the resonance structure of [8]IF-CNB (a), and the resonance structure of COT with BLA (b). The major and minor contributions of the resonance forms appear in case of lower symmetric structures.
repeating units $N$ from 6 to 8. [8]IF-CNB is shown to exhibit apparent alternations of the distances with a difference of about 0.1 Å, whereas there are no alternation patterns in the cases of [6]IF-CNB and [8]IF-CNB$^{2+}$. The average of the distances is expected to converge to a value for $N \to \infty$, which corresponds to the situation of one-dimentional polymer of repeating indeno[1,2-b]fluorenes, even though we did not confirm here whether the alternation patterns remains for $N \to \infty$ or not.

Judging from these geometric features, we suggest that closed-shell contributions of resonance structures in [8]IF-CNB can be illustrated like that given in Fig. 3a. In the previous study, we discussed the analogy of the resonance structures between [6]IF-CNB and [6]annulene (benzene). If all the six-membered rings are equivalent, like those in [6]IF-CNB, the contributions of upper and lower forms should be equivalent to each other. However, in [8]IF-CNB ($D_4$), rather quinoid-like and benzenoid-like structures appear alternately, and thus the weights of these forms in the resonance structure are not considered to be equivalent. Again, we focus on the analogy of the resonance structures between [8]IF-CNB and [8]annulene (cyclooctatetraene, COT). In COT, local minimum structure is non-planar tub-shaped structure that belong to $D_{2d}$ symmetry. From the quantum chemical calculations, two types of planar stationary point structures on the potential energy surface (PES) of COT have been predicted. The one belongs to the highest possible $D_{4h}$ symmetry with no BLA, and the other does to the $D_{2h}$ symmetry with BLA, both of which correspond to the saddle-points on the PES. For $D_{4h}$ structure, two resonance forms shown in Fig. 3b should be equivalent to each other, while for $D_{4h}$ or $D_{2d}$ structures, the weights of these two forms are not equivalent, depending on how the structure is distorted. Because of the flexible framework of COT, local minimum structure becomes non-planar, and π-electrons are relatively localized on each double bond. On the other hand, [8]IF-CNB is considered to have more rigid framework than COT owing to the fused-ring structure of the belt, and thus significant structural relaxation effect (like that in tub-shaped COT) is expected to be suppressed to some extent. From this viewpoint, even though the weights of two forms in the resonance structure given in Fig. 3a are not equivalent, the contribution of minor one may not be negligible. In such case, we can expect delocalized nature of π-electrons even in [8]IF-CNB, which would be reflected in the aromatic character of this system. On the other hand, [6]IF-CNB was shown to have intermediate multiradical characters in our previous theoretical study. We therefore need to investigate the open-shell characters of [8]IF-CNB and [8]IF-CNB$^{2+}$ and the corresponding orbital energy levels, in order to discuss their electronic structures.

In Table 1, we summarized calculation results of diradical characters ($y_i$; $i = 0, 1$ and 2). In the previous paper, we found

<table>
<thead>
<tr>
<th>System</th>
<th>$y_0$</th>
<th>$y_1$</th>
<th>$y_2$</th>
<th>$\gamma_{xxx}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>[6]IF-CNB ($D_{3d}$)</td>
<td>0.398</td>
<td>0.398</td>
<td>0.134</td>
<td>213</td>
</tr>
<tr>
<td>[8]IF-CNB ($D_4$)</td>
<td>0.526</td>
<td>0.210</td>
<td>0.210</td>
<td>698</td>
</tr>
<tr>
<td>[8]IF-CNB$^{2+}$ ($D_{2d}$)</td>
<td>0.471</td>
<td>0.458</td>
<td>0.143</td>
<td>−2691</td>
</tr>
</tbody>
</table>

Fig. 4 Frontier MO levels of [6]IF-CNB that are related to its aromatic and open-shell characters at the spin-restricted LC-RBLYP/6-31G* level.

Fig. 5 Frontier MO levels of [8]IF-CNB that are related to its aromatic and open-shell characters at the spin-restricted LC-RBLYP/6-31G* level.
that \( y_0 \) and \( y_1 \) for [6]IF-CNB become the same value owing to the degenerate nature of the corresponding HOMO/HOMO−1 and LUMO/LUMO+1 levels (see also Fig. 4). In Fig. 4–6, we therefore summarized the frontier MO levels of these systems at the spin-restricted LC-RBLYP/6-31G* level, where MOs are chosen from the viewpoints of bonding and anti-bonding pairs corresponding to \( y_i \) and of the aromatic characters. The HOMO–LUMO gaps obtained at the present level of approximation are considered to be larger than experimental HOMO–LUMO gaps of related open-shell compounds. We employed the LC-RBLYP/6-31G* in consistent to the level for evaluation of \( y_i \) (LC-UBLYP/6-31G*). [8]IF-CNB is found to have intermediate \( y_0 \) value (0.526), which is larger than \( y_0/y_1 \) values of [6]IF-CNB (0.398). In contrast to the results of [6]IF-CNB, the \( y_1 \) and \( y_2 \) values (0.210) are found to be the same in [8]IF-CNB. Indeed, the MO levels related to these diradical characters, HOMO-2/HOMO-3 and LUMO+1/LUMO+2, are shown to degenerate, respectively (see Fig. 5).

Similar to the case of [6]IF-CNB, we can suggest an analogy of the frontier MO levels of [8]IF-CNB to those of COT. Within the Hückel approximation, COT with BLA-less regular octagon \( D_{4h} \) structure is known to have degenerate HOMO/LUMO levels. When the symmetry of COT reduces to \( D_{2h} \), the HOMO–LUMO gap is expected to increase owing to the structural distortion, whereas the degenerate level structure for other \( \pi \)-orbitals is considered to be preserved.4 The increase of HOMO–LUMO gap due to the reduction of symmetry is related to the decrease of \( y_0 \) value. Actually, metastable structure of [8]IF-CNB with higher \( C_{4v} \) symmetry is found to have \( y_0 = 1 \) (see Table S2†). Similar tendency was also discussed in the \( D_{4h} \) (\( y_0 \sim 1 \)) and \( D_{2h} \) (\( y_0 \sim 0.1 \)) structures of TCPTP,28 but in the present case, \( y_0 \) value of the most stable local minimum structure of [8]IF-CNB with reduced \( D_{4} \) symmetry still exhibit intermediate value. This may originate in the structural features of [N]IF-CNB, namely, indeno[1,2-\( b \)] fluorene substructures are involved in the smoothly curved \( \pi \)-surface in the belt. When we remove two electrons from the HOMO of [8]IF-CNB, this MO becomes unoccupied level. In dicaticionic [8]IF-CNB\(^{2+}\), \( y_0 \) and \( y_1 \) values are found to be very close to each other, which is considered to reflect the degenerate HOMO/HOMO−1 and LUMO/LUMO+1 levels (see Fig. 6). The reason why \( y_0 \) and \( y_1 \) differ very slightly may be related to the theoretical and numerical treatments of evaluating \( y \). In the case of [8]IF-CNB\(^{2+}\), \( y_0 \) and \( y_1 \) values (0.471 and 0.458, respectively) are found to be in the intermediate region.

In order to investigate how and where the unpaired electrons distribute, we illustrate the maps of odd-electron densities for each system (see Fig. 7a). Note that we only plot the odd-electron densities from the contributions of \( y_0 \) ([8]IF-CNB) or average of \( y_0 \) and \( y_1 \) ([6]IF-CNB and [8]IF-CNB\(^{2+}\)). In all the systems, odd-electron densities are found to distribute primarily on the vertices of five-membered rings. Owing to the high symmetries of these systems (even for [8]IF-CNB), the amplitude of odd-electron densities on each vertex is found to be equivalent. In Fig. 7b, we also show the net Hirshfeld charge of [8]IF-CNB\(^{2+}\) in order to investigate the distributions of positive charges. In

![Fig. 6 Frontier MO levels of [8]IF-CNB\(^{2+}\) that are related to its aromatic and open-shell characters at the spin-restricted LC-RBLYP/6-31G* level.](image1)

![Fig. 7 Odd-electron density distributions (a) and net Hirshfeld charges where the contributions from hydrogen atoms are summed into heavy atoms (b).](image2)
contrast to the odd-electron density maps, positive charges are found to be distributed on the six-membered rings as well as the vertices of five-membered rings. Such delocalized features of positive charges in [8]IF-CNB\(^{2+}\) may be reflected in the response properties of this system.

These unique geometric features and electronic structures of [\(\pi\)]IF-CNB are expected to be reflected in their aromatic characters. Among several criteria for evaluating aromatic characters, we here employed the GIMIC method, which is a powerful tool to qualitatively and (semi-)quantitatively understand the aromatic nature of conjugated systems. In our previous paper, we have found from the analysis of GIMIC that magnetically-induced current primarily originating in the \(\pi\)-electrons can flow over the belt along the curved \(\pi\)-surface, and as a result of the balance between the paratropic and diatropic contributions, [6]IF-CNB is expected to exhibit global aromatic nature, even though the amplitudes of MIC density are weak compared to those of benzene (see also Fig. 8).\(^2\) Fig. 8 shows the side views of the maps of bond-integrated MIC densities for all the calculated systems. Note that, in Fig. 8, left- and right-arrows for the bond-integrated MIC density correspond to the diatropic (aromatic) and paratropic (antiaromatic) contributions in the induced ring currents, respectively. The values are the amplitudes of bond-integrated MIC density in the unit of nA T\(^{-1}\). Note that, the value of bond-integrated MIC density for benzene at the same calculation level is more than 10 nA T\(^{-1}\). From the maps, the main circuit of the flow of MIC is considered to be that along the repeating \(p\)QM substructure. In contrast to the results of [6]IF-CNB, paratropic contributions are found to become predominant in the ring current in [8]IF-CNB. Judging from the amplitudes of bond-integrated MIC densities (0.2–1.6 nA T\(^{-1}\)) and the direction of flow, [8]IF-CNB is expected to exhibit weak global antiaromatic character. In the case of the dicaticionic [8]IF-CNB\(^{2+}\), directions and amplitudes of bond-integrated MIC densities are found to become inverted for bonds of \(p\)QM substructures, the feature of which suggests that [8]IF-CNB\(^{2+}\) is expected to exhibit weak global aromatic character like [6]IF-CNB. The tendencies of aromaticity analyzed by the GIMIC method are in consistent with those predicted from \(n\)\(^p\).

The delocalized and intermediate open-shell characters of these systems suggest unique physico-chemical properties, such as, enhancement of third-order NLO properties. Actually, [6]IF-CNB was found to exhibit enhanced third-order NLO properties compared to its closed-shell counterpart [(6,6) CNB].\(^3\) In Table 1, we also summarized static \(\gamma_{\text{xxxx}}\) of these systems. In Fig. 9, we plot the \(\gamma_{\text{xxxx}}\) densities. The \(\gamma_{\text{xxxx}}\) value of

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**Fig. 8** Maps of bond-integrated MIC densities. Orange left-arrows and blue right-arrows represent diatropic and paratropic contributions when the magnetic field \(B\) is applied along the present direction, respectively.

**Fig. 9** Maps of \(\gamma_{\text{xxxx}}\) densities. Yellow/blue meshes represent the isosurfaces of \(\gamma_{\text{xxxx}}\) density with the contour values of \(\pm 200\) a.u., respectively.
intermediate open-shell characters. Drastic changes of $\gamma_{xxxx}$ can be found in dicationic [8]IF-CNB$^{2+}$ ($-2691 \times 10^3$ a.u.). The amplitude of $\gamma_{xxxx}$ for [8]IF-CNB$^{2+}$ is 3.86 times as large as that of [8]IF-CNB. Furthermore, the sign of $\gamma_{xxxx}$ in [8]IF-CNB$^{2+}$ is predicted to be negative at the present level of approximation, which can be visually illustrated by the inverted direction of third-order polarization in the $\gamma_{xxxx}$ density map (see Fig. 9). Such a negative longitudinal $\gamma$ value can sometimes be observed in charged soliton or charged radical species.\textsuperscript{44-46}

In order to investigate the origins of negative $\gamma_{xxxx}$ value of [8]IF-CNB$^{2+}$, we evaluated the static polarizabilities ($\alpha_{xx}$) for these systems at the same level as that for $\gamma_{xxxx}$ (see Table 1). On the basis of the sum-over-states expression for the symmetric systems within the three levels (0, $n$ and $m$), the $\gamma_{xxxx}$ value is approximately expressed as:\textsuperscript{44}

$$\gamma_{xxxx} = \gamma^I + \gamma^{III-2} = -4\left(\frac{\mu_{0x}^2}{E_{0x}}\right)^3 + 4\left(\frac{\mu_{nx}^2}{E_{0x}}\right)^2 - 4\left(\frac{\mu_{0x}^2}{E_{0x}}\right)^2,$$

(1)

where $\mu_{0x}$ and $E_{0x} = E_n - E_0$ are the $x$-axis component of transition dipole moment between the states 0 (ground state) and $n$ (excited state), and excitation energy for state $n$, respectively. Since excitation energy is positive, the first term $\gamma^I$, which is called type-II term, contributes negatively to total $\gamma_{xxxx}$ within the present approximation. In order to analyze the origin of negative $\gamma_{xxxx}$, from the sum-over-states expression, qualitative and (semi-)quantitative excited state calculations are needed. Such a calculation for the present large-sized delocalized multiradicaloids and its charged species at the same accuracy demand significant computational effort. On the other hand, $\gamma^I$ is related to $\alpha_{xx}$ as:\textsuperscript{44}

$$\gamma^I = \frac{(\alpha_{xx})^2}{E_{0x}} = \frac{(\alpha_{xx})^2}{2(\mu_{0x})^2}.$$

(2)

Therefore, even though the sign of full sum-over-states expression of $\gamma_{xxxx}$ is determined by the balance of negative and positive terms ($\gamma^I$ and $\gamma^{III-2}$), the amplitude of $\alpha_{xx}$ may give information to understand the origin of negative $\gamma_{xxxx}$. The $\alpha_{xx}$ value of [8]IF-CNB$^{2+}$ (1600 a.u.) is found to be about 1.54 times as large as that of [8]IF-CNB (1040 a.u.) (1.54$^2$ = 2.37). Large $\alpha_{xx}$ value is usually related to the small excitation energy for optically-allowed excited states and the large amplitudes of transition moments for these states. Even though there are several possible reasons, it is speculated that there is a possibility of existing such excited states in the dicationic [8]IF-CNB$^{2+}$. Similar tendencies of reduction of excitation energy and negative $\gamma$ were also predicted in the charged radical species like pentalene and s-indacene, the electronic structures of which are called symmetric resonance structures with invertible polarization (SRIP).\textsuperscript{45,46} Since we can illustrate several equivalent canonical forms in the resonance structures where radicals and positive charges appear on different sites, there may be a connection to the mechanism of SRIP. As another possible explanation, we may suggest interference effects of virtual excitation pathways appearing in $\gamma^{III-2}$ of the full sum-over-states expression. Further analysis of excited states for such large-sized open-shell systems will be an interesting topic from the viewpoint of developing methods of quantum chemistry and controlling molecular functionalities, and thus studies toward this direction is still ongoing by using the multi-reference multi-state electron correlation methods.

**Conclusions**

In this study, we have investigated the geometric features, electronic structures and third-order NLO properties of [6]IF-CNB, [8]IF-CNB and [8]IF-CNB$^{2+}$, in order to clarify the relationship between $n_\pi$, aromatic and open-shell characters, and physico-chemical properties of theoretically designed [N]IF-CNB. [N]IF-CNBs, which involve repeating indeno[1,2-b]fluorene substructures on the curved $\pi$-surface, are shown to have highly symmetric local minimum structures depending on $n_\pi$. The most stable local minimum structures of [6]IF-CNB and [8] IF-CNB$^{2+}$ with $n_\pi = 4n + 2$ are found to belong to highest possible symmetry with BLA-less nature of characteristic bonds. All the five- and six-membered rings in these systems are equivalent to each other, suggesting their delocalized electronic structures over the belt. In contrast, the most stable local minimum structure of [8]IF-CNB with $n_\pi = 4n$ is shown to belong to reduced $D_2$ symmetry. Finite alternation patterns for the specific CC bonds and for the distance between the vertices of adjacent five-membered rings suggest that there are quinoid- and benzenoid-like pQM substructures that appear alternately in the belt. Owing to the existence of pQM substructures, all these systems are expected to have intermediate open-shell characters. It is found that these systems have degenerate frontier MO levels depending on $n_\pi$, the feature of which is reflected in (nearly) degenerate $y_1$ values. Aromatic characters of [8]IF-CNB and its dicaticonal state evaluated by the GIMIC method are in consistent with those predicted from $n_\pi$. Therefore, aromaticity classifications by $n_\pi$ are expected to be applicable for [N]IF-CNB with larger $N$ and their two-electron oxidized species. For the third-order NLO properties, two-electron oxidation of [8]IF-CNB is expected to enhance drastically the amplitude of $\gamma_{xxxx}$ value, whereas the sign of $\gamma_{xxxx}$ is inverted. We speculated that the negative value of $\gamma_{xxxx}$ of [8]IF-CNB$^{2+}$ is related to the enhancement of polarizability based on SRIP mechanism, which is associated with lowering of the excitation energies and raising of the transition moment amplitudes of the optically-allowed excited states, even though further analysis of excited states by using highly accurate quantum chemical calculations are needed in order to clarify the mechanisms.

Even though there are plenty of barriers to realize such fully conjugated open-shell macrocycles, such as ring-strain during the aromatization of the belt, requirements of bulky
substituents at the vertices of five-membered rings to increase kinetic stabilities, etc., the present theoretical results suggest that \( [N] \)IF-CNbands related materials are attractive targets of organic synthesis for realizing novel open-shell multifunctional conjugated macrocycles. For example, \( [N] \)IF-CNbands may also be able to offer different electrostatic environments in inner and outer spatial domain, the feature of which can be useful for host-guest chemistry, like \( [N] \)CPPs.\(^{12-17}\) Very recently, synthesis of methylene-bridged \([6]CPP\) has been reported by Segawa, Itami and coworkers.\(^{18}\) Furthermore, some well-designed open-shell conjugated molecules are known to form unique molecular aggregates where covalent-like intermolecular interactions exist. Since variation of \( n_\pi \) is expected to drastically change the structures and properties of \( [N] \)IF-CNbands, electron transfer between host and guests, which may be triggered during the trapping of guests or by external stimuli, will be an interesting strategy for establishing novel dynamic open-shell functional materials and their control schemes. Studies toward such directions as well as further investigation on the size and charged-/spin-state dependences of the structures and properties of \( [N] \)IF-CNbands and related systems are ongoing project in our group.

**Conflicts of interest**

There are no conflicts to declare.

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**Notes and references**


