# Polarizing agents beyond pentacene for efficient triplet dynamic nuclear polarization in glass matrices

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ABSTRACT: Triplet dynamic nuclear polarization (triplet-DNP) is a technique that can obtain high nuclear polarization under moderate conditions. However, in order to obtain practically useful polarization, large single crystals doped with a polarizing agent must be strictly oriented with respect to the magnetic field to sharpen the electron spin resonance (ESR) spectra, which is a fatal problem that prevents its application to truly useful biomolecular targets. Instead of this conventional physical approach of controlling crystal orientation, here, we propose a chemical approach, i.e., molecular design of polarizing agents; pentacene molecules, the most typical triplet-DNP polarizing agent, are modified so as to make the triplet electron distribution wider and more isotropic without loss of the triplet polarization. In fact, the modification of pentacene with thiophene moieties makes the ESR spectrum sharper and stronger than that of pentacene. To elucidate the effect of the substitutions on spin polarization and zero-field splitting parameters, which determine ESR spectrum, state-of-the-art quantum chemical calculations were performed and revealed that the direction of the spin polarization is altered by the modification with thiophene moieties and the size of D and E parameters are reduced from parent pentacene due to the partial delocalization of spin densities on the thiophene moieties. The triplet-DNP with the new polarizing agent successfully exceeds the previous highest <sup>1</sup>H polarization of glassy materials by a factor of 5. This demonstrates the feasibility of a polarizing agent that can surpass pentacene, the best polarizing agent for more than 30 years since triplet-DNP was first reported, in the unoriented state. This work provides a pathway toward practically useful high nuclear polarization of various biomolecules by triplet-DNP.

# INTRODUCTION

Nuclear magnetic resonance (NMR) and magnetic resonance imaging (MRI) are powerful analytical techniques that provide useful information in the chemical and clinical fields, but poor polarization of nuclear spins limits their sensitivity and hinders their versatility. Dynamic nuclear polarization (DNP) is groundbreaking technology that boosts the sensitivity of NMR and MRI by transferring the polarization of electron spins to nuclear spins.<sup>1–7</sup> However, sufficiently high electron spin polarization in thermal equilibrium requires harsh conditions, such as high-field (~ 7 T) and cryogenic temperatures (~1 K), preventing a wide range of practical applications.<sup>8–11</sup>

DNP using photo-excited triplet electrons (triplet-DNP) can achieve nuclear hyperpolarization even at room temperature since high electron spin polarization between triplet sub-levels is achieved regardless of temperature by the spin-selective intersystem crossing (ISC) (Figure 1a).<sup>12-18</sup> The polarization of triplet electron spins can be efficiently transferred to nuclear spins by integrated solid effect (ISE) sequence with microwave irradiation and magnetic field sweep (Figure 1b). Since the electron spin resonance (ESR) linewidth of triplet is broadened over 100 mT at X-band by hyperfine coupling and dipolar interactions between triplet electrons, only limited part of spin packets simultaneously satisfies the resonance condition for polarization transfer, so-called Hartmann–Hahn condition. By sweeping the magnetic field during microwave irradiation, spin packets in the sweep range can be used for polarization transfer. However, the magnetic field sweep width is practically limited to about 10 mT, since the field sweep must be slow enough to maintain the adiabatic condition and within the lifetime of triplet polarization.

In previous studies of triplet-DNP, the strategy has been to suppress the ESR linewidth broadening due to dipole interactions by strictly controlling the orientation of the single crystal doped with the polarizing agent relative to the magnetic field.<sup>14,15,19,20</sup> This strategy has enabled most of the electron spin packets to be utilized for polarization transfer, and nuclear spin polarization of several tens of percent has been achieved. However, this strategy is technically demanding because it requires growing large single crystals on the order of centimeters, and the orientation of the single crystals must be very strictly controlled. Furthermore, the matrix single crystals have been limited to aromatic compounds that can be doped with polarizing agents, which is a serious problem for application to important non-aromatic MRI probes such as pyruvate. Although the transfer of nuclear spin polarization to various molecules by dissolving hyperpolarized naphthalene single crystals at high concentrations has been reported, its medical applications are not easy because it has been limited to DNP in organic solvents.<sup>21</sup> Considering these difficult challenges faced by the physical strategy of controlling crystal orientation, new breakthroughs are needed to extend the application of triplet-DNP to a variety of biomolecules.



**Figure 1.** a) Scheme of triplet-DNP. After photoexcitation of polarizing agents, spin-selective ISC produces spin-polarized triplet electrons. Triplet electron spin polarization is transferred to nuclear spin by the integrated solid effect (ISE). b) ISE sequence. The pulsed laser excitation is followed by microwave irradiation and magnetic field sweep. c) ESR simulation with various *D* and *E* values while other parameters are fixed (*g* = 2, *B* = 8.98 GHz,  $p_x = 0.4$ ,  $p_y = 0.2$ ,  $p_z = 0.2$ ).

Here, we propose a solution to the essential problem of triplet-DNP by a chemical approach of molecular design of polarizing agents instead of a physical approach of orientation control. In order to achieve triplet-DNP of various biomolecules, it is necessary to achieve high nuclear polarization in randomly oriented samples, free from the constraints of single crystals. The key parameters are the zero-field splitting (ZFS) parameters D and E, which represent the energy gaps between triplet sublevels in the absence of magnetic field and determine the linewidth of the ESR spectrum of the triplet state (Figure 1b).<sup>22-24</sup> Simulations with different values of D and E while keeping other parameters fixed show that the ESR spectrum becomes sharper as *D* and *E* become smaller in the examined range, and that there are more spin packets within 10 mT, the magnetic field sweep width of the ISE sequence (Figure 1c). D and E values are described as follows,

$$D = \frac{3}{4} \frac{\mu_0}{4\pi h} (g_e \mu_B)^2 \langle \frac{r^2 - 3z^2}{r^5} \rangle$$
$$E = \frac{3}{4} \frac{\mu_0}{4\pi h} (g_e \mu_B)^2 \langle \frac{y^2 - x^2}{r^5} \rangle$$

where *r* is the average distance between electron spins, *x*, *y*, *z* are dipole axes,  $g_e$  is the g-factor,  $\mu_B$  is the Bohr magneton, *h* is plank constant and  $\mu_0$  is Magnetic permeability in vacuum. From the above equations, the *D* and *E* values are related to the average distance between spins and the rhombicity of the zero-field splitting tensor, respectively. This implies that the molecular design with smaller *D* and *E* should make the triplet electron distribution wider and more isotropic.

As a proof of concept, based on pentacene, the most typical polarizing agent of triplet-DNP, we designed novel polarizing agents by modifying pentacene with 5-membered rings, 6,13-di(benzofuran-2-yl)pentacene (DBFP), 6.13bis(benzo[b]thiophen-2-yl)pentacene (DBTP) and 6,13di(thiophen-2-yl)pentacene (DTP) (Figure 2a). The relatively less sterically hindered furan and thiophene ring is not orthogonal to pentacene and expected to extend the conjugation, contributing to an extended and more isotropic triplet electron distribution. These new polarizing agents actually exhibit a sharper and stronger ESR signal than pentacene, as well as a higher polarization ratio. The state-of-the-art quantum chemical calculations were performed to predict the ZFS parameters and the ISC rate. The calculations of the *D* tensor based on molecular structures allows us to predict direction of the principal axis, i.e., the sign of the *D* and *E* parameters. It was confirmed that the modifications to parent pentacene enhance broadening the triplet electron distribution, providing theoretical support for the rationality of the present design guidelines, i.e., reducing the size of *D* and *E* values. Interestingly, the spin selectivity during the ISC, which is found to be governed by the vibronic effects, was proved to be altered by the modification with thiophene moieties. Triplet-DNP with novel polarizing agents exceeded the 1H polarization obtained so far in amorphous matrices by a factor of 5, demonstrating that they rationally outperform pentacene, which has been the best performance to date.

#### **RESULTS AND DISCUSSION**

**Time-resolved ESR measurements.** The new polarizing agents DBFP, DBTP and DTP were synthesized following the reported method,<sup>25</sup> and their purity was confirmed by NMR and elemental analysis (see Supporting Information for details). For time-resolved ESR measurements, the polarizing agents were dispersed in  $\beta$ -estradiol glass.  $\beta$ -estradiol forms a colorless glass that is stable at room temperature, making it a convenient matrix for ESR measurements.<sup>26</sup> Parent pentacene and phenyl-modified pentacene, 6,13-diphenylpentacene (DPP) were used in addition to DBFP, DBTP, and DTP for comparison (Figure 2a).

Each polarizing agent was mixed with  $\beta$ -estradiol at a concentration of 0.01 mol% and dispersed in  $\beta$ -estradiol by heating to 453 K, which is above the melting point of  $\beta$ -estradiol, followed by rapid cooling with liquid nitrogen. Molecularly dispersed state of polarizing agents in  $\beta$ -estradiol was confirmed by absorption and fluorescence spectra. The peak positions and shapes of the absorption spectra of each



**Figure 2.** a) Chemical structures of polarizing agents, pentacene, 6,13-diphenylpentacene (DPP), 6,13-di(benzofuran-2-yl)pentacene (DBFP), 6,13-bis(benzo[b]thiophen-2yl)pentacene (DBTP) and 6,13-di(thiophen-2-yl)pentacene (DTP). b) Time-resolve ESR spectra of pentacene, DPP, DBFP, DBTP and DTP doped in  $\beta$ -estradiol at room temperature. Microwave frequencies for each spectral measurement are 8.988, 8.959, 8.983, 8.960 and 8.963 GHz, respectively. Results simulated with EasySpin toolbox in Mathlab are shown as red lines. c) Principal axes of zero-field splitting interaction of pentacene triplet electrons.

polarizing agent in the  $\beta$ -estradiol glass were similar to those in the chloroform solution, and were completely different from those of the neat solid, which had a large red shift and broadening (Figure S1). When the absorption spectra were measured by changing the concentration of DBTP in the  $\beta$ -estradiol glass, the spectral shape changed above 0.1 mol%, while it was maintained below 0.01 mol% (Figure S2). Pentacene and its derivatives do not emit fluorescence due to efficient singlet fission when aggregated,<sup>27</sup> but the fluorescence spectra in the  $\beta$ -estradiol glass of each polarizing agent were similar to those in chloroform (Figure S3). From these results, we confirmed that each polarizing agent was dispersed in  $\beta$ -estradiol glass without aggregation at the concentration of 0.01 mol%.

Under pulsed laser irradiation at 527 nm, ESR spectra and signal decays of each polarizing agents in  $\beta$ -estradiol glass were measured at X-band (0.24-0.40 T) and room temperature (Figure 2b, S4). Laser intensity was set to 8.4 A for pentacene, DBFP, DBTP and DTP and 8.8 A for DPP. ESR spectra were simulated by EasySpin toolbox in Matlab<sup>28</sup> and obtained parameters were summarized in Table 1. The principal axes of ZFS interactions were same as pentacene and all signs of *D* and *E* values were assumed to be *D* > 0 and *E* < 0 (Figure 2c).<sup>29,30</sup> Compared to pentacene, DBTP and DTP exhibited much stronger and sharper ESR spectra, which is reflected in their smaller values of |D| and |E|. On the other hand, DPP showed similar ESR spectral shape and intensity and similar values of |D| and |E| as pentacene. The |D| value was even smaller in DBFP than in DBTP. The reason for these differences in ZFS parameters will be discussed later based on theoretical calculations. The ESR intensity of DBFP was smaller than that of DBTP. This may be due to the lower ISC efficiency of DBFP in addition to the smaller population difference (Table 1, S1). Interestingly, pentacene, DPP, and DBFP were polarized along the x-axis, whereas DBTP and DTP were y-polarized, indicating a clear substituent effect. Decays of the ESR signal showed that both DBTP and DTP exhibited spin-lattice relaxation times  $T_{1e}$  comparable to that of pentacene and long enough to apply the ISE sequence (Figure S4, Table S2).

**Table 1.** |D| and |E| values and relative population ratio of  $p_{x_i} p_{y_j} p_z$  of each polarizing agent in the zero-field spin state.

	D  (MHz) <sup>*</sup>	E  (MHz) <sup>*</sup>	$p_x$	$p_y$	$p_z$
Pentacene	1360.1	47.2	0.56	0.31	0.13
DPP	1368.3	39.9	0.54	0.34	0.12
DBFP	1190.9	45.1	0.60	0.26	0.14
DBTP	1330.0	26.9	0.20	0.80	0.00
DTP	1311.5	29.2	0.20	0.78	0.02

\*Sign of *D* and *E* value are assumed to be *D* > 0 and *E* < 0.

**Calculations of spin selective ISC rates.** To elucidate a detailed mechanism of the spin polarization experimentally observed for the pentacene derivatives, we performed quantum chemical calculations. It was assumed that the spin polarization is caused by a spin selective ISC between the lowest singlet excited state ( $S_1$ ) and an energetically close triplet state ( $T_n$ ) of the molecules. To characterize the final state of the ISC, energy levels of the low-lying triplet states of an unmodified pentacene molecule were calculated at the LC-BLYP( $\mu$ =0.15)/def2-TZVP level of theory. The 0-0 energy differences from  $S_1$  state were predicted as

-1.33, -0.19, 0.77 eV for  $T_1$  (1<sup>3</sup>B<sub>2u</sub>),  $T_2$  (1<sup>3</sup>B<sub>1g</sub>), and  $T_3$  (1<sup>3</sup>B<sub>3u</sub>) states of the pentacene, and therefore the  $T_2$  state that is characterized by the HOMO  $\rightarrow$  LUMO+1 and HOMO-1  $\rightarrow$ 

LUMO singly excited state is likely to be the final state of the spin selective ISC. The energy differences between the initial ( $S_1$ ) and final ( $T_2$ ) at each minimum were predicted to be -193, -85.1, 3.96 meV for pentacene, DBTP, DBFP molecules, respectively. The prediction that the energy level of  $T_2$  state is about the same or higher than that of  $S_1$  state for the DBFP molecule is consistent with the observation in our experiments that the fluorescence quantum yield of DBFP is higher than that of the other molecules, i.e., the ISC transition is not efficient in DBFP.

The ISC transition rate from the  $S_1$  to  $T_2$  spin sublevels  $T_2^X$ ,  $T_2^Y$ , and  $T_2^Z$  states are expressed approximately

$$k_{\rm ISC}^{\mu(={\rm X},{\rm Y},{\rm Z})} = \frac{2\pi}{\hbar} \sum_{i,f} \left| \left\langle \Psi_i \middle| \widehat{H}_{\rm SOC} \middle| \Psi_f^{\mu} \right\rangle \right|^2 \delta({\rm E}_i - {\rm E}_f) \qquad (1)$$

applying the Fermi's Golden rule where  $\Psi_i$  and  $\Psi_f^{\mu}$  indicate vibronic states associated respectively with  $S_1$  and  $T_2^{\mu}$ states,  $E_i$  and  $E_f$  are their energy levels, and  $\hat{H}_{SOC}$  is the spin-orbit coupling operator. Since the potential energy surfaces of the spin sublevels are virtually degenerated, i.e., the energy differences are negligible in the prediction of the rates, the difference in the ISC rate for  $\mu$ =X, Y, Z should be caused solely by the difference in the magnitude of the SOC matrix elements (SOCMEs)  $\langle \Psi_i | \hat{H}_{SOC} | \Psi_f \rangle$ . Because the SOC-MEs of the planar molecules vanish for all the components  $\mu$  = X, Y, Z at the equilibrium geometries due to the point group symmetry, and the Heltzberg-Teller like effects, e.g., vibronic spin-orbit coupling, caused by molecular vibrations that break the planar symmetry are dominant,<sup>31,32</sup> and the SOCMEs should be corrected by adding the first-order vibronic spin-orbit coupling term as

$$V^{\mu} = \langle \Psi_{i} | \hat{H}_{\text{SOC}} | \Psi_{f}^{\mu} \rangle$$
  

$$\approx \langle \Phi_{i} | \hat{H}_{\text{SOC}} | \Phi_{f}^{\mu} \rangle |_{q=0} \langle \nu_{i} | \bar{\nu}_{f} \rangle$$
  

$$+ \sum_{k} \partial_{q_{k}} \langle \Phi_{i} | \hat{H}_{\text{SOC}} | \Phi_{f}^{\mu} \rangle |_{q=0} \langle \nu_{i} | q_{k} | \bar{\nu}_{f} \rangle + O(q^{2})$$
(2)

where  $v_i$  and  $\overline{v}_f$  denotes the vibrational states in the initial  $(\Phi_i)$  and final  $(\Phi_f^{\mu})$  electronic states. Hereafter, we denote the derivative  $\partial_{q_k} \langle \Phi_i | \hat{H}_{\text{SOC}} | \Phi_f^{\mu} \rangle$  at the reference structure (q=0) as  $\partial_{q_k} V_0^{\mu}$ . There are many ways to evaluate Eq.(1). In this study, the time-dependent formulation33-37 implemented in ORCA5<sup>38,39</sup> was used,<sup>37,40</sup> where the ISC rates are evaluated by the Fourier transform of the time-correlation function. The approximations adopted in the method, e.g., the potential surfaces are modeled by simple harmonic oscillators, can be insufficient to make quantitative predictions for the ISC rates, but should be sufficient to predict the size relation between the spin sublevels. The calculated ISC rates in the pentacene, DBFP and DBTP molecules are shown in Table 2. The most efficient ISC transitions were predicted for  $S_1 \rightarrow T_2^X$  in the pentacene and DBFP molecules and for  $S_1 \rightarrow T_2^{\text{Y}}$  in the DBTP molecule. It is consistent with the spin polarization in T1 state observed in our ESR experiments.

To analyze what vibrational modes contribute to enhancing the vibronic spin-orbit coupling, the first-order derivative of SOCME along each mode was calculated by finite difference techniques. The largest derivative of the SOCME of

**Table 2.** Predicted ISC rates between  $S_1$  and  $T_2$  sublevels  $T_2^{\mu}$  ( $\mu$  = X, Y, Z) based on the Fermi Golden rule. Relative rate ratios are shown in parentheses.

$ns^{-1}$	$T_2^{\rm X}$	$T_2^{\mathrm{Y}}$	$T_2^Z$
Pentacene	$4.4 \times 10^{-2}$	$2.8 \times 10^{-2}$	$1.1 \times 10^{-5}$
	(0.61)	(0.38)	(0.00)
DBFP	$1.8 \times 10^{-3}$	$3.1 \times 10^{-5}$	$1.1 \times 10^{-5}$
	(0.97)	(0.02)	(0.01)
DBTP	$1.0 \times 10^{-4}$	$5.1 \times 10^{-2}$	$5.9 \times 10^{-3}$
	(0.00)	(0.89)	(0.10)

each sublevel are summarized in Table 3 and the associated normal modes are shown in Figure 3. For the pentacene molecule, it was found that the out-of-plane modes consisting of C-H and C-C bending contribute to increasing the SOCME of the  $T_2^X$  and  $T_2^Y$  components. Those out-of-plane distortions should mix the carbon  $\pi$  orbitals with  $\sigma$  bonding orbitals, i.e., the transition between carbon  $2p_z$  and  $2p_y$  or  $2p_x$  orbitals are involved in the ISC, and thus the SOCME of the  $T_2^X$  and  $T_2^Y$  should be enhanced. None of the modes contribute to increasing that of the  $T_2^Z$  components. It implies that transitions between carbon  $2p_x$  and  $2p_y$  orbitals are negligible in the ISC even with the out-of-plane distortions. Similar observation was made for the DBFP molecule.

For the DBTP molecule, the largest derivatives were found along the modes involving displacements in the benzothiophene substituents, and those modes contribute only to  $T_2^Y$ and  $T_2^Z$  components. The derivative values along those modes are much larger than that along out-of-plane distortions of the pentacene backbone, which increase the SOCME of the  $T_2^X$  and  $T_2^Y$  components. For the DBFP molecule, by contrast, the increase in the SOCME caused by the modes involving displacements of the benzofuran substituents is less than those by the out-of-plane modes of the pentacene backbone, and it results in that the transition to the  $T_2^X$  component is the most efficient.

The discrepancy between the experimental and theoretical predictions of the order of  $T_2^X$  and  $T_2^Z$  transition rates for the DBTP molecule, i.e., the transition rate to the  $T_2^{\rm Z}$  was largely overestimated in the FGR calculations, should be attributed to the approximations adopted in the calculations. It is quite possible that the modes that significantly increase the SOCME of the  $T_2^Z$  are coupled with the rotation of the benzothiophene substituents and cannot be described by the harmonic oscillator model nor the first-order vibronic spinorbit coupling model Eq.(2). In fact, the time-dependent method for the ISC rate could not converged in the case of DTP, which has similar structure to DBTP. In contrast, those approximations should work well for the pentacene molecule because all the modes that contribute to enhancing the ISC are out-of-plane modes and the transition should occur in the vicinity of the minimum along the modes for  $S_1$  and  $T_2$ sates. In fact, the ratio is in good agreement with the experiment.

**Calculations of ZFS parameters.** To investigate the trends of *D* and *E* values observed in the experiments, the CASSCF calculations for the  $T_1$  state were performed. Because the *D* and *E* values are parameters adopted in the phenomenolog

$\omega_k$ /cm <sup>-1</sup>	$\partial_{q_k} V_0^{\mathrm{X}} / i \mathrm{~cm}^{-1}$	$\partial_{q_k} V_0^{\mathrm{Y}}$ / <i>i</i> cm <sup>-1</sup>	$\partial_{q_k} V_0^{\mathrm{Z}} / i ~\mathrm{cm}^{-1}$
Pentacene			
902	0.358	0.002	0.000
866	0.000	0.249	0.000
1508	0.000	0.002	0.007
DBFP			
906	0.249	0.004	0.005
834	0.009	0.127	0.002
1579	0.042	0.003	0.029
DBTP			
901	0.175	0.000	0.000
1364*	0.002	0.281	0.169

**Table 3.** Derivatives of the SOCME along normal modes. Normal modes that give the large derivatives are selected for each sublevel  $T_2^X$ ,  $T_2^Y$ , and  $T_2^Z$ .

\*The mode ( $\omega$  =1364 cm^-1) provide the largest derivative for both Y and Z

ical model Hamiltonian, there are many ways to define those values from the ab initio quantum chemical calculations. In the case of triplet states, it is straightforward to determine the D and E parameters such that energy gaps of the spin Hamiltonian reproduce the ab initio calculations. Because the contributions of the SOC term to the ZFS is negligible as compared with that of the SSC term for aromatic hydrocarbons, only the latter was considered in the calculations. The active space consisting of all the  $\pi$  orbitals of the pentacene backbones, i.e., as much as CAS(22e, 22o), were adopted in the CASSCF calculations. It exceeds the capability of the computer available for the foreseeable future, and the DMRG-CASSCF theory<sup>41-43</sup> implemented in the BLOCK2 program<sup>44</sup> was used with a sufficient bond dimension (m=1024) to circumvent the limitation. The D-tensor were calculated as the matrix elements of the SSC between the non-relativistic triplet states with the different spin projections<sup>45</sup> and the *D* and *E* parameters and the orientation of the principal magnetic axis were obtained by the diagonalization. The SSC two-electron integrals were evaluated by using the RI approximation: the two- and three-center integrals were evaluated with Libcint and PySCF<sup>46-48</sup> package by using the translational invariance of the two-electron field gradient integrals.

Table 4 shows the calculated *D* and *E* parameters for the pentacene derivatives. One of the advantages of predicting the *D*-tensor by *ab initio* calculations is that the direction of the principal axes in the molecular axis system, i.e., sign of the *D* and *E* parameters, can also be directly determined. The signs were predicted to be D > 0 and E < 0, i.e., the energy ordering is  $T_2^X > T_2^Y > T_2^Z$ . The signs of the parameters were identical to those reported by the previous experimental study for pentacene molecules in a *p*-terphenyl host crystal,<sup>49</sup> while those have not been reported for the other



**Figure 3.** Normalized displacement vector of the modes shown in Table 3. The modes along which the derivatives of the SOCME are less than 0.1, i.e., not significant, are omitted.

molecules as far as we know, the signs predicted are necessary for reproducing the EAEAEA pattern of the experimental ESR spectra for DBTP and DTP in Figure 2b. The DBFP, DBTP, DTP, DPP, and pentacene molecules have larger D parameters in that order, which is in agreement with the experiments except for DPP of which |D| is larger by 8.2 MHz than pentacene in the experiment. It was found that the DBFP, which exhibits the smallest D value, has the smallest dihedral angle between the pentacene backbone and the substituents planes, and the spin density is more delocalized on the substituents than the DPP, DTP, and DBTP (see Figure S14). Although the DPP, DTP, and DBTP have similar size of the dihedral angle, D value of the DTP and DBTP are clearly smaller than that of the DPP, which could reflect the effect of the M-shell electrons of sulfur atoms. Note that the spin-spin interaction is a two-electron interaction and it is not necessary explained solely by the oneelectron property, spin density. In fact, it was reported that the mean-field approximation to the spin-spin interaction causes a non-negligible error for ZFS parameters.<sup>50</sup>

As observed in the experiments, the |E| values for the pentacene derivatives were predicted to be very small as less than 50 MHz, though the size relationship between the molecules were not identical between the experiments and *ab*  *initio* calculation. The energy gap between  $T_2^X$  and  $T_2^Y$  sublevels is so small that there remains the possibility that the accuracy of the prediction is not fully converged within that range, e.g., the size of the active space and bond dimension etc. Furthermore, the fitting made for experimental spectrum can involve errors when the signal is not enough clear.

**Table 4.** Calculated *D* and *E* parameters obtained by the DMRG-CASSCF theory and dihedral angles between the pentacene backbones and the planar moieties.

	$D (\mathrm{MHz})^*$	$E (MHz)^*$	Dihedral angle
Pentacene	1171.4	-44.5	-
DPP	1143.1	-28.7	69 °
DBFP	1101.3	-17.8	52 °
DBTP	1129.0	-21.1	71 °
DTP	1125.2	-19.9	69 °

\*The signs D > 0 and E < 0 indicate that the energy ordering of the spin sublevels is  $T_2^X > T_2^Y > T_2^Z$  where the molecular inplane long and short axes and the out-of-plane axis correspond to the X, Y, Z principal axes

The theoretical analysis indicates that molecular vibrations that break the planar symmetry and cause a mixing of carbon  $\sigma$  bonding orbitals with  $\pi$  orbitals determine the spinselectivity in the ISC of the pentacene molecule. The spinselectivity is largely changed by the substituent modifications; the substituent effect is so large that the direction of the polarization has altered between pentacene (long-axis) and DBTP (short-axis) molecules, though it seems that we should go beyond the harmonic oscillator model or the firstorder vibronic spin-orbit coupling model for predicting the rate quantitatively for DBTP molecule. The ZFS parameters D and E are also changed by the modification with substituents. Because the spin-selectivity in the ISC and the ZFS parameters are determined by the SOC and SSC terms, respectively, those properties are expected to be controlled separately by substituent modifications.

Triplet-DNP experiments. To evaluate the potential of the new polarizing agent, the <sup>1</sup>H NMR signal intensity after triplet-DNP was compared between DTP and pentacene. Since  $\beta$ -estradiol has a short <sup>1</sup>H spin-lattice relaxation time  $T_1$  of only a few seconds, o-terphenyl (OTP) was employed instead. OTP has been used as a model glass matrix for triplet-DNP because of its long  $T_1$  of about 60 s (Figure S7).<sup>51</sup> The polarizing agents, DTP or pentacene, were doped in OTP using the same procedure except that the heating temperature was changed to 333 K. Triplet-DNP experiments were performed with OTP in the glassy state by ISE sequence under a magnetic field of 0.66 T for pentacene and 0.65 T for DTP at 120 K. After photoexcitation by irradiation of 527 nm pulsed laser, polarization was transferred from electron spins to nuclear spins by microwave irradiation (17.6 GHz) under magnetic field sweep (10 mT). After repeating this

ISE sequence at a frequency of 500 Hz for a certain time, the sample was shuttled into an NMR coil within 1 s and <sup>1</sup>H NMR spectra were measured. (See Supporting Information for details). The <sup>1</sup>H polarization and enhancement factor was calculated by comparing the <sup>1</sup>H NMR signal intensity of ethanol at 296 K and hyperpolarized signal after triplet-DNP at 120 K (Figure S8). Figure 4 shows the buildup curves of <sup>1</sup>H spin polarization by varying the time of triplet-DNP. Experimental parameters such as concentration of the polarizing agent and the magnetic field were optimized based on the intensity of the enhanced NMR signal (Figure S9, S10). <sup>1</sup>H polarization and enhancement factor of OTP after 5 min of triplet-DNP reached 0.85% and 1,540 times, respectively, with DTP, which was about 4 times larger than that with pentacene (0.21% and 370 times). While the enhancement factor obtained by triplet-DNP also depends on other parameters of the polarizing agent, such as electron spin-lattice relaxation time  $T_{1e}$ , triplet lifetime  $\tau_{T}$  and triplet quantum yield  $\Phi_{\rm T}$  but these parameters did not show significant differences for the examined polarizing agents (Figure S4, S5, S6, Table S1, S2). Hence, the novel polarizing agent DTP gave significantly larger <sup>1</sup>H polarization than the best performing pentacene of the past mainly because DTP shows the stronger and sharper ESR than pentacene and can generate more spin packets within the magnetic field sweep range. This was achieved by the rational molecular design of polarizing agents that simultaneously achieves spin-spin interaction control to reduce |D| and |E| values by making the triplet distribution more extended and more isotropic, and spin-orbit coupling control with high sublevel selectivity in ISC.



**Figure 4.** Build up curve of <sup>1</sup>H signal intensity of OTP doped with pentacene (green line) and DTP (blue line), and partially deuterated OTP (OTP : [D14]OTP = 10 : 90 wt%) doped with DTP (red line).

Final polarization can be improved by reducing the density of <sup>1</sup>H spins to increase the spin-lattice relaxation time  $T_1$ since the accumulation of polarization by triplet-DNP competes with the spin-lattice relaxation. By changing the glass matrix from OTP to a mixture of OTP and fully-deuterated [D14]OTP (OTP : [D14]OTP := 10 : 90 wt%), the <sup>1</sup>H  $T_1$  value was elongated from 1 min to 3 min at 0.65 T and 120 K (Figure S7). Remarkably, this elongated  $T_1$  resulted in the further increase of <sup>1</sup>H polarization and enhancement factor to 7.8% and 14,000 times, respectively (Figure 4), which is 5 times higher than the previous highest polarization of 1.5% in amorphous system by triplet-DNP. $^{51}$ 

# CONCLUSION

In conclusion, we have proposed a rational design guideline for polarizing agent as a chemical approach to overcome the intrinsic problem of conventional triplet-DNP, which requires the use of big single crystals with strictly controlled orientation in order to obtain high polarization enhancement. Based on pentacene, the most representative polarizing agent of triplet-DNP, we have succeeded in reducing the ZFS parameters |D| and |E| while maintaining the high polarization of triplet by extending the conjugation through introducing the thiophene ring. This means that it is possible to simultaneously control the spin-spin interaction of the triplet that determines |D| and |E| and the spin-orbit coupling related to spin polarization. Theoretical calculations support for the rationality of the present design guidelines, i.e., by extending the  $\pi$  conjugation to make the distribution of spin density wider and more isotropic, the size of D and E values are reduced and the electron spin polarization is increased at the same time. The power of this chemical approach was demonstrated by exceeding by a factor of 5 the highest <sup>1</sup>H polarization ever achieved by triplet-DNP in amorphous system. This indicates that a polarizing agent that exceeds the performance of pentacene, which has been considered the best over the last 30 years since triplet-DNP was first proposed, is feasible. We believe that further molecular design improvements based on the current discovery will lead to the practical application of triplet-DNP, achieving nuclear polarization of more than 10% for a wide variety of biomolecules under moderate conditions independent of molecular orientation.

# ASSOCIATED CONTENT

#### Supporting Information.

Experimental details including materials, synthesis and measurement setup, UV-vis absorption and fluorescence spectra, time-resolved absorption decays, fluorescence quantum yields, time-resolved ESR and triplet-DNP measurements, calculation details, calculated normal modes and spin density distributions.

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#### Author Contributions

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The authors declare no competing financial interest.

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