## SYNTHESES AND PROPERTIES OF BENZANNELATED DIPHENYLPENTADECAFULVENE DERIVATIVES

## Jūro OJIMA,\* Shigeyasu KURODA, † and Mitsuru KIRITA

Department of Chemistry, Faculty of Science, Toyama University, Gofuku, Toyama 930 <sup>†</sup>Department of Industrial Chemistry, Faculty of Engineering, Toyama University, Nakagawasonomachi, Takaoka 933

Synthesis of 10-methyl-16,16-diphenyl-4,5-benzo-6,8-bisdehydro- 4, 12-methyl-16,16-diphenyl-6,7-benzo-8,10-bisdehydro- 5, and 16,16-diphenyl-4,5:10,11-dibenzo-6,8-bisdehydropentadecafulvene 6 has been described. Comparison of <sup>1</sup>H-NMR spectra of these benzamelated fulvenes 4-6 with that of nonbenzamelated fulvene 3 reveals that 5,10-dimethyl-16,16-diphenyl-6,8-bisdehydropentadecafulvene 3 is atropic.

Of many methods available for synthesis of fulvene system, most involve either addition reaction followed by elimination, or condensation reaction between various reagents and cyclic compounds to elaborate an exocyclic double bond. However, we recently reported a synthesis of diphenyltrideca-(type  $\mbox{\ le m=1}\$ ), -pentadeca- (type  $\mbox{\ le m=1}\$ ), -heptadeca- ( $\mbox{\ le m=2}\$ ) by an intramolecular oxidative coupling from the corresponding cross-conjugated acyclic polyenes containing two terminal acetylene groups. This method is advantageous since the precursors of the desired fulvenes would serve as the "open chain" models necessary for investigation of ring currents. Furthermore, although the fulvenes of both types  $\mbox{\ le and }\mbox{\ le messample models}\$  and  $\mbox{\ le messample models}\$  are suggested to be atropic by an examination of their H-NMR spectra, an uncertainty was left in that judgement of tropicity of type  $\mbox{\ le messample models}\$ , since the inner ( $\mbox{\ H}^B$ ,  $\mbox{\ H}^A$ ,  $\mbox{\ H}^D$ ) protons of  $\mbox{\ le messample models}\$  are unusually low field, whereas the outer ( $\mbox{\ H}^A$ ,  $\mbox{\ H}^B$ ,  $\mbox{\ H}^C$ ,  $\mbox{\ H}^B$ ,  $\mbox{\ H}^C$ ,  $\mbox{\ H}^B$ , and methyl protons of  $\mbox{\ le messample models}\$  and  $\mbox{\$ 

It is well recognized that annelation of one or more benzenoid rings for a large-membered conjugated  $\pi$ -electron system usually causes progressive reduction of tropicities of the macrocyclic system with an increasing annelation if the position of benzenoid rings excludes the contribution of an equivalent Kekulé structure. Accordingly, we considered that an inspection on the tropicity of the benzammelated series of 3 would provide an alternative diagnostic tool for judgement of tropicity of 3 itself.

In order to get a further insight for the tropicity of  $\mathfrak{Z}$  as well as to test the scope of validity of the reaction sequence used for the syntheses of the previously reported fulvene derivatives, 2) we were interested in examining the properties of the benzammelated derivatives of  $\mathfrak{Z}$ . In this paper, we

now describe syntheses and properties of the title compounds 4-6, as well as the further examination of the properties of 3.

The synthesis was accomplished by the reaction sequence similar to that adopted for compounds of types 1 and 2, 2) and we have obtained the objective fulvenes in moderate yields. Thus, the reaction of 1-(o-ethynylphenyl)-9-methyl-1,4,6,8-undecatetraen-10-yn-3-one (11), 4b) 1-(o-ethynylphenyl)-9-methyl-1,3,6,8-undecatetraen-10-yn-5-one (12), 4b) and 1,7-bis (o-ethynylphenyl)-1,4,6-heptatrien-3-one (13), 4b) with diphenylketene<sup>5)</sup> formed in situ from diphenylacetyl chloride and triethylamine in dry benzene at room temperature for 20-25 h afforded 11-(o-ethynylphenyl)-3-methyl-9-(diphenylmethylene)-3,5,7,10-undecatetraen-1-yne (8, yellow needles, mp. 134-135°C, 38%), 6) 11-(o-ethynylphenyl)-3-methyl-7-(diphenylmethylene)-3,5,8,10-undecatetraen-1-yne (9, yellow needles, mp. 143-144°C, 34%), and 1,7-bis (o-ethynylphenyl)-3- (diphenylmethylene)-1,4,6-heptatriene (10, pale yellow needles, mp. 170-171°C, 54%), respectively. Oxidative coupling of 8, 9, and 10 with anhydrous copper (II) acetate in pyridine and dry ether at  $50^{\circ}$ C oxidative coupling of 8, 9, and 10 with anhydrous copper (II) acetate in pyridine and dry ether at  $50^{\circ}$ C oxidative coupling of 8, 9, and 10 with anhydrous copper (II) acetate in pyridine and dry ether at  $50^{\circ}$ C oxidative coupling of 8, 9, and 10 bisdehydro- (5, yellow needles, mp. 177-178°C, 46%), and 16,16-diphenyl-4,5:10,11-dibenzo-6,8-bisdehydro- (5, yellow needles, mp. 177-178°C, 46%), and 16,16-diphenyl-4,5:10,11-dibenzo-6,8-bisdehydropentadecafulvene (6, pale yellow needles, mp. 175-176°C, 68%), respectively.

The  $^1H$ -NMR spectral data of these fulvenes 3-6 as well as those of the acyclic compounds 7-10 are

$3-10$ in CDC1 <sub>3</sub> at 200 MHz ( $\tau$ -Values)	$H^{C}$ , $H^{D}$ , $H^{E}$ , $CH_3$ Ph H -C=CH	3.33 3.67	2.40 3.57 8.20 2.7–2.9	-0.93 -0.10 +0.13	3.64 3.25 3.64 8.06 2.47–2.84 6.66	.94) 3.52 8.16 (2.60–2.94)	3.59 (2.37–3.24) 8.05 (2.37–3.24) 6.65, 6.70	(2.61–2.86) 8.15 (2.61–2.86)	3.06 (2.36–2.84) (2.36–2.84) 6.62, 6.68	3.09 2.60-2.91	Fig. 1. The <sup>1</sup> H-NMR spectra of fulvenes in CD listed in Table 1. Individual assignments were made basis of multiplicities and coupling constants, and clarified by the decoupling experiments where possib necessary.  Comparison of the <sup>1</sup> H-NMR chemical shifts of the various protons of the benzamelated fulvenes 4–6 wi those of the corresponding acyclic compounds 8–10, r spectively, indicates that the fulvenes 4–6 are atro
s of the Compounds	H <sup>B</sup> ' H <sup>C</sup>	3.31 3.67	4.15 3.60	+0.84 -0.07	3.09	4.11	2.37-3.24) 3.63	(3.36	2.95	3.59	since no significant upfield and downfield shift due to outer and inner protons, respectively, are observed as compared with those of the corresponding acyclic composition. This is readily seen from Figure 1 which shows the <sup>1</sup> H-1 spectra of 3-6 at room temperature. In the spectra of benzamelated fulvenes 4-6, compared with that of 3, the high- and low-field shifts of the resonances of the outent and inner protons, respectively, are not observed. However, as exemplified in the spectrum of the monobenzamelated fulvene 4, the inner H <sup>B</sup> and the outer H <sup>B</sup> , H <sup>C</sup> , H <sup>E</sup> protonate at almost the same region as those of nonbenzamelated fulvene 3. As to only the inner H <sup>B</sup> proton, the similar tendency is true even in the spectra of the benzamelated fulvenes 5 and 6. Thus, decreasing of both the high- and low-field shifts due to the outer and improtons, respectively, is not clearly observed on passing protons, respectively, is not clearly observed on passing protons.
emical Shifts	HB			•	l		2.95	2.47	2.51		
The <sup>1</sup> H-NMR Chemical	н <sup>А</sup> н <sup>А'</sup>					.00 (2.60–2.94)					
Table 1. I	Compd.	(a) 3.		0- (2-2)√	i	3.00	3,	.5 (3.	3.	3,	from nonbenzammelated 3 to dibenzammelated 6 via mone benzammelated fulvenes 4 and 5. Since a decrease in by by benzammelation was found to be true for the corresponding 14π-electron, diatropic bisdehydro[15]annulenone server

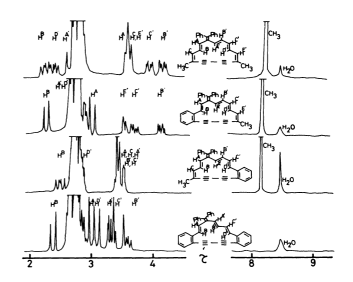


Fig. 1. The <sup>1</sup>H-NMR spectra of fulvenes in CDCl<sub>3</sub>

Comparison of the <sup>1</sup>H-NMR chemical shifts of the various protons of the benzamelated fulvenes 4-6 with those of the corresponding acyclic compounds 8-10, respectively, indicates that the fulvenes 4-6 are atropic, since no significant upfield and downfield shift due to the outer and inner protons, respectively, are observed as compared with those of the corresponding acyclic compounds. This is readily seen from Figure 1 which shows the 1H-NMR spectra of 3-6 at room temperature. In the spectra of the benzammelated fulvenes 4-6, compared with that of 3, the high- and low-field shifts of the resonances of the outer and inner protons, respectively, are not observed. However, as exemplified in the spectrum of the monobenzammelated fulvene 4, the inner HB and the outer HB', HC', HE' protons resonate at almost the same region as those of nonbenzannelated fulvene 3. As to only the inner H<sup>B</sup> proton, the similar tendency is true even in the spectra of the benzammelated fulvenes 5 and 6. Thus, decreasing of both the high- and low-field shifts due to the outer and inner protons, respectively, is not clearly observed on passing from nonbenzammelated  $\mbox{\em 3}$  to dibenzammelated  $\mbox{\em 6}$   $\emph{via}$  monobenzannelated fulvenes 4 and 5. Since a decrease in tropicity by benzammelation was found to be true for the corresponding

which should have geometry similar to these fulvene systems, this observation obtained from the  $^{1}\text{H-NMR}$  spectra of the fulvenes together with an examination of the comparison of the chemical shifts of the corresponding acyclic model 7,  $^{2}$  leads us to believe that an appearance of the resonances due to the outer and inner protons at a rather high- and low-field, respectively, in the spectrum of 3 does not reflect a paramagnetic ring current induced in  $16\pi$ -electron, cross-conjugated system of 3. The  $^{1}\text{H-NMR}$  spectral pattern observed for 3 must be a reflection of the fact that the anisotropies of two phenyl and diacetylene moieties are at work. Thus, it is concluded that the fulvene 3 is not paratropic, but is atropic.

In addition, in rather surprisingly contrast to the cases of 3-5, the dibenzannelated fulvene does not exist in a conformation 6a, but in 6 at room temperature. 8) This follows from the fact that the value of  $J_{B',C'}$  is 10 Hz, pointing to an s-trans relationship of the  $CH^{B'}$ ,  $CH^{C'}$  bond.

Since the bisdehydro[15]annulenone ring has been found to have very high planarity of the molecular skeleton among the similar type bisdehydroannulenone systems  $^{9)}$  and the corresponding dibenzammelated bisdehydro[15]annulenone was shown to exist in a conformation 14, changing from an oxygen atom of the annulenone to bulky phenyl groups at  $C_{16}$ -position of the fulvene must arouse an unlikely conformation 6. One possible explanation for this result is that in cases of 3-5, the steric hindrance experienced between the two phenyl groups and the neighboring olefinic protons can be relieved over the whole of a rather flexible, nonbenzammelated or monobenzammelated molecular skeleton. On the other hand, the molecular skeleton of 6, which is made to less flexible system than those of 3-5 by the fusion of two benzene rings, can not relieve the hindrance, causing the conformational change. The conformation of 6 is structually relevalent to that of the corresponding bisdehydro[15]annulenone 15 carrying a methyl group at  $C_{14}$ -position.

## References

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  H3C = 5a = 1.4

  H3C
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The details on this point will be reported elsewhere.