Synthetic studies in quest of Platonic hydrocarbon dodecahedrane

GOVERDHAN MEHTA*, K RAJA REDDY and MANGALAM S NAIR

School of Chemistry, University of Hyderabad, Hyderabad 500 134, India

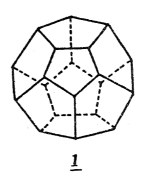
Abstract. In pursuit of Platonic hydrocarbon dodecahedrane 1, a retrosynthetic theme indicated in scheme 1, was formulated. The precursor tetraquinanedione synthon $\underline{5}$ was first designed through a photo-thermal olefin metathesis approach. The tetraquinanedione 5 was further elaborated to exo, exo-tetraquinane diester 15 through carbonyl homologation, oxidation, esterification sequence, scheme 5. Bis-cyclopentannulation of exo, exo-diester $\underline{15}$ by Greene methodology delivered a functionalised C_{20} -hexaquinane $\underline{44}$, having exo-annulated cyclopentane rings. Cyclopentane inversion was achieved by a set of reactions involving enone generation, double bond isomerisation and hydrogenation to give spheroidal (C_{2v})-C₂₀-hexaquinanedione-diester 47, the penultimate precursor of dodecahedrane 1. Several interesting transformations and rearrangements of polyquinanes are also described.

Keywords. Dodecahedrane; C_{2v}-tetraquinane dione; cis-Hexaquinane; bis-cyclopentannulation; cyclopentanone inversion.

Dodecahedrane $(1)^+$, a $C_{20}H_{20}$ hydrocarbon of I_h symmetry is one of the most structurally complex and aesthetically appealing polyquinanes known. The challenge of synthesising this platonic hydrocarbon of spheroidal shape, that arises through its twelve constituent cis, syn fused five-membered rings, has elicited a lot of interest from synthetic chemists around the world (Mehta 1978; Eaton 1979; Paquette 1979, 1984). Two outstanding efforts from the research groups led by Paquette and Prinzbach have resulted in the synthesis of dodecahedrane in 1982 and 1987, respectively (Ternansky et al 1982; Paquette et al 1983, 1987; Fessner et al 1987). Apart from these, it is known that synthetic efforts towards dodecahedrane were pursued or are currently being pursued in various other laboratories (Schleyer 1957; Woodward et al 1964; Jacobson 1967; Repic 1976; Eaton et al 1977, 1984; Deslongchamps and Soucy 1981; Roberts and Shoham 1981; McKervey et al 1981; Monego 1982; Baldwin and Beckwith 1983; Mehta and Nair 1983, 1985; Baldwin et al 1984; Carcellar et al 1986; D G Farnum and T A Monego - unpublished results; G Mehta and K R Reddy-unpublished results). Concurrently, the fascinating structural features of $\underline{1}$ have also been investigated by several theoretical chemists (Schulman et al 1975; Schulman and Disch 1978; Ermer 1977; Dixon et al 1981; Baum et al 1982). Synthetic efforts towards 1 were initiated in our laboratory in early 1981, when no successful approach to this challenging hydrocarbon was known. A perusal of the synthetic approaches that were in practice convinced us

*For correspondence

⁺IUPAC nomenclature: Undecacyclo[9.9.0.0^{2,9}.0^{3,7}.0^{4,20}.0^{5,18}.0^{6,16}.0^{8,15}.0^{10,14}.0^{12,19}.0^{13,17}leicosane.



that a good chance of success lay in a strategy leaning heavily on symmetrization that could inherently overcome some of the problems caused by steric strain and non-bonded interactions. With this in view, a retrosynthetic analysis of $\underline{1}$ was carried out as depicted in scheme 1, which identified the symmetrical, highly functionalized C_{20} -hexaquinane $\underline{2}$ as the pivotal pretarget and the C_{12} -tetraquinane dione $\underline{5}$ as the key starting synthon. Hexaquinane $\underline{2}$, which retains a spheroidal contour has all the 20 carbon atoms as well as two strategically placed $-CX_2$ type functionalities for molecular stitching through a four-fold intramolecular displacement reaction on $\underline{2}$ (scheme 2) to furnish a secondodecahedranedione $\underline{6}$. This

$$\frac{1}{2}$$

$$\frac{1}{2}$$

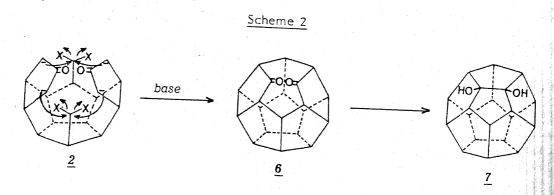
$$\frac{1}{2}$$

$$\frac{1}{2}$$

$$\frac{1}{2}$$

$$\frac{3}{2}$$

ultimate precursor could then be enticed into a pinacolic coupling to deliver the dodecahedrane framework 7 as shown in scheme 2.



Emanating from this retrosynthetic analysis, our synthetic approach to $\underline{1}$ was divided into three stages of increasing complexity viz., (i) development of a new methodology towards the synthesis of the starting tetraquinane synthon, e.g., $\underline{5}$; (ii) adoption of a suitable cyclopentanone annulation and inversion strategy (viz., conversion of $\underline{4} \rightarrow \underline{2}$) and (iii) deployment of the molecular stitching plan using the two $-CX_2$ functionalities in $\underline{2}$ to close-in the sphere (scheme 2). Progress towards the implementation of this theme is described in this account.

Stage 1

Synthesis of the tetraquinanedione 5 and its further elaboration

At the time of inception of this effort towards dodecahedrane, there were only two reported syntheses of functionalized tetraquinanes available in literature (Fukunaga and Clement 1977; Paquette et al 1978), both of which suffered from severe limitations during scale-up processes and one of them (Fukunaga and Clement 1977) did not deliver the requisite functionality. Therefore, we first turned our attention to a generalised approach to functionalized tetraquinanes. Borrowing pointers from our successful approach to triquinane synthesis (Mehta et al 1979, 1981), we devised a photo-thermal olefin metathetic approach as shown in scheme 3.

For the synthesis of tetraquinanedione $\underline{5}$ as shown in scheme 4, the starting materials were identified as 7-t-butoxynorbornadiene $\underline{8}$ and tetrachlorodimethoxy-cyclopentadiene $\underline{9}$. Diels-Alder reaction between $\underline{8}$ and $\underline{9}$ followed by acetone sensitized $\pi^2 s + \pi^2 s$ closure of the *endo*-adducts resulted in the hexacyclic product $\underline{10}$ in 57% yield (see also Astin and MacKenzi 1975, Byrne *et al* 1974). A three-step sequence consisting of (i) dechlorination using Li-t-BuOH-THF, (ii) hydrolysis

using 3N HCl in dioxane and (iii) PCC oxidation produced the symmetrical (13 C NMR: $\delta 210.0$, 48.8, 42.4, 41.8) caged, bis-homopentaprismane dione $\underline{11}$ in an overall yield of 55% (Mehta and Nair 1983, 1985b). Flash vacuum pyrolysis (Mehta et al 1979, 1981) of this diketone through a quartz tube preheated to 580°C provided the tetraquinanedione $\underline{5}$ in 70% yield (Mehta et al 1983; Paquette et al 1986; Sedelmeier et al 1986). The structure and $C_{2\nu}$ symmetry of this dione could be readily deduced from its 1 H NMR spectrum and 13 C NMR signals at $\delta 214.2$, 133.5, 62.7, 43.2. Following this procedure, the dione $\underline{5}$ could be prepared in multi-gram quantities quite uneventfully.

In order to introduce an equivalent of $-CX_2$ group as per the theme of scheme 1, the tetraquinanedione 5 was further elaborated to the diester 15 (ester group as an equivalent of $-CX_2$ functionality) (G Mehta and K R Reddy, unpublished results). This was achieved through a two-fold carbonyl homologation followed by oxidation as shown in scheme 5. Wittig olefination on 5 with methoxymethylphosphorane gave a 1:2 mixture of 12 and 13. Acid hydrolysis of these enolethers to the thermodynamically more stable dialdehyde 14 and oxidation of the dialdehyde with PDC in DMF followed by esterification of the resulting diacid using diazomethane gave the exo, exo-diester 15. This diester was identical to that obtained by reductive cleavage and isomerization of the Hedaya-Paquette diester 16, scheme 5 (Paquette et al 1978). The diester 15 was considered eminently suitable for further evolution towards the target 1.

The photo-thermal olefin metathetic approach to the tetraquinanes described above was also adopted for the synthesis of oxa-tetraquinane 18, as this was more readily accessible through C_5 and C_6 building blocks as compared to the tetraquinanedione 5. Model studies for bis-cyclopentannulation and ring inversion strategy were, therefore, initially conducted using 18 (Mehta and Nair 1985). As

Scheme 5

shown in scheme 6, tetraquinane $\underline{18}$ was synthesized from benzoquinone and cyclopentadiene via the oxa-bird cage compound $\underline{17}$ in relatively few, high yielding steps.

Stage 2

Bis-cyclopentannulation of tetraquinanes. Synthesis of spheroidal hexaquinanes

As eluded to above, the next important step towards the conquest of dodecahedrane was the appendage of two cyclopentane rings on $\underline{15}$ as in $\underline{4}$. A perusal of the available methods in literature for cyclopentannulation of olefins suggested the Greene methodology (Green and Depress 1979) of dichloroketene addition to olefins followed by regiospecific ring expansion to be superior to others as it would render strategic placement of the carbonyl group. For the exploratory studies, the oxa-tetraquinane 18 was utilised as it was readily obtained in large quantities. When oxa-tetraquinane 18 was treated with an excess of dichloroketene, generated in situ, by using trichloroacetylchloride and a Zn-Cu couple according to procedure of Bak and Brady (1979), a bis-dichloroketene adduct 19 was obtained. While the regiochemical assignment of 19 follows from its ¹H and ¹³C NMR data, the exo-stereochemistry is based on the propensity of folded polyquinanes to react from the open convex face. The bis-adduct 19 was ring expanded using diazomethane in ether and the now redundant chlorine atoms were removed using Zn/NH₄Cl/MeOH milieu (Noyori et al 1974) to give the hexaquinanedione 20 as shown in scheme 7.

Scheme 7

Even though bis-(cyclopentanone)annulation of $\underline{18}$ had been achieved, it could be successfully utilised for the synthesis of dodecahedrane only if the two newly appended cyclopentane rings could be projected within the cavity of the polyquinane framework. This required inversion of stereochemistry at four centres in $\underline{20}$ (marked with asterisks). This transformation was envisaged in a three step sequence as illustrated in scheme 8.

Thus, the first step in the transformation was the introduction of double bonds in conjugation with the carbonyl groups. For this, any of the three general methods known viz., (i) the Reich procedure of α -phenylselenylation followed by dehydroselenation (Reich et al 1975; Reich 1979), (ii) the Saegusa procedure using Pd(II) mediated dehydrosilylation of an enolsilylether (Ito et al 1978) or (iii) the

Mincionne procedure using $PdCl_2$ in refluxing t.BuOH to effect direct dehydrogenation (Mincionne et al 1977) could be utilised. While all three methods proved successful, the Mincionne procedure gave the best results. Reaction of hexaquinanedione $\underline{20}$ with $PdCl_2$ in refluxing t BuOH transformed it into a mixture of bis-enones $\underline{21}$, $\underline{22}$, $\underline{23}$ and the monoenone $\underline{24}$ in approximately equal amounts as illustrated in scheme 9 (Mehta and Nair $\underline{1985a}$).

Scheme
$$\frac{9}{9}$$

$$\frac{PdCl_2}{t.BuOH}$$

$$\frac{20}{75\%}$$

$$\frac{23}{24}$$

The next task was the relocation of double bonds in bis-enones. As it is well known that α,β -unsaturated carbonyl compounds equilibrate with their β,γ -isomers in the presence of base, we decided to subject the bis-enones to this treatment. In order to avoid damage to the ether linkage, a mild base like DBU was chosen for the purpose. The bis-enone 22 under the influence of DBU isomerised to 25. The structure of 25 was arrived at through its 13 C NMR and 500 MHz 1 H NMR data. In the case of 22 the protons adjacent to ether linkage appeared as a doublet (J = 4.3 Hz), whereas in 25 they appeared as a doublet of a doublet ($J_1 = J_2 = 5.3$ Hz). Catalytic hydrogenation of 25 resulted in the projection of the two cyclopentanone units within the cavity, thus giving rise to 26. In this manner, the second stage in our approach to dodecahedrane, viz., bis-cyclopentanone annulation and ring inversion was successfully demonstrated on model oxatetraquinane 18, scheme 10.

Scheme 10

With the experience of the above model studies, it was now the opportune time to aim at dodecahedrane itself. The endo, endo-diester 33 and the diacetate 34 derived from it appeared to be the most appropriate starting materials. The latter had the advantage of not being susceptible to epimerisation during the subsequent operations. Also, the molecular stitching plan (scheme 1) could be first tested with

 $\underline{35}$ which would only undergo a two-fold cyclisation to furnish an octaquinanedione $\underline{36}$ or $\underline{37}$, scheme 11.

Scheme 11

$$CO_2Me$$
 CO_2Me
 CH_2OAc
 CH_2OC
 CH_2OC

Thus, the diester 15 was routinely elaborated to diacetate and then bis-cyclopentannulated employing the dichloroketene addition methodology outlined earlier in this article, scheme 7. The resulting hexaquinanedione 38 was dehydrogenated to the bis-enones 39 and 40, using the PdCl₂-t·BuOH dehydrogenation procedure. Attempts were now directed to relocate the enone double bonds to the tetrasubstituted ring junction position in order to effect cyclopentane inversion. Towards this end, the bis-enone 39 of axial symmetry was exposed to be a C_{20} -dioxaoctaquinane 41 formed via intramolecular Michael addition of the endo-hydroxymethyl groups to the two enone moieties, scheme 12. The structure of exhibited diagnostic signals at δ 4·15–4·12 (2H, dd, J_1 = 2·4 Hz, J_2 = 9·6 Hz), tionality.

In order to circumvent the deviation leading to 41, DBU was deployed as the non-nucleophilic base for the deconjugation of enone moieties in 40. However, in this case another type of intramolecular Michael addition intervened and the novel heptacyclic ene-dione 42 was obtained. The ¹H and ¹³C NMR data are fully to 43 were thwarted by unanticipated formation of 41 and 42, we were impelled to seek a different strategy to gain access to the key folded hexaquinane 2.

Believing that the inversion of the cyclopentenone rings in 39 and 40 was complicated by the presence of two endo-acetoxymethyl substituents protruding within the spherical cavity (cf. $22 \rightarrow 25$ change), we decided to operate on an exo, bay any steric congestion within the developing sphere during the crucial ring inversion manoeuvre. However, provision was kept to project the exo-substituent into the endo-position at an appropriate stage to execute the molecular stitching

Scheme 12

CO₂Me

33 CO₂Me

43 CH₂OAc

CH₂OAc

CH₂OAc

CH₂OAc

$$\frac{39}{80}$$
 CH₂OAc

 $\frac{39}{40}$ CH₂OAc

plan (stage 3) as per scheme 1. Towards this objective, the exo, exo-diester 15, readily obtainable from the endo, endo-isomer through thermodynamic equilibration with methanolic sodiummethoxide was chosen as the starting tetraquinane derivative. The diester 15 was once again bis-cyclopentannulated via dichlor-oketene addition, diazomethane ring expansion and reductive dechlorination sequence to furnish the hexaquinanedione 44, scheme 13. The dione 44 was

Scheme 13

regioselectively transformed into a single bis-enone of axial symmetry following the Saegusa procedure (Ito et al 1978). When exposed to acid or base to relocate the enone moieties, the bis-enone $\underline{45}$ once again exhibited recalcitrant behaviour. After many exploratory attempts, we were most gratified to observe that $\underline{45}$ could be converted into bis-ketal $\underline{46}$ under carefully controlled acetalisation in the presence of camphorsulphonic acid. The 10 line ¹³C NMR spectrum fully supported its structure and identified the location of two double bonds at the tetrasubstituted bridgehead position. Catalytic hydrogenation of $\underline{46}$ and deacetalisation under the mildly acidic conditions furnished the folded hexaquinanedione $\underline{47}$, scheme 13. The arrival at the long sought $\underline{47}$ was indicated by its 400 MHz $\overline{}$ H NMR spectral parameters and 8 line $\overline{}$ 3°C NMR spectrum (δ 219·3, 175·8, 57·4, 55·6, 52·3, 49·4, 43·0, 40·6). The structure of this key penultimate precursor of dodecahedrane, by our approach was further secured through single crystal X-ray diffraction studies. With a reasonable supply of $\underline{47}$ at our disposal, we set about giving expression to the stage 3 of our plan, scheme 1.

Stage 3

The molecular stitching plan. End game en route to dodecahedrane

In order to orchestrate the intramolecular ring closure on to a secododecahedrane derivative, it was imperative that the ester groups of 47 be made to fall within the cavity. To achieve this, we proposed to exploit the propensity of the spheroidal systems towards protonation from the convex face. Thus, inversion of the two ester moieties in 47 was envisaged via deprotonation-kinetic protonation employing non-nucleophilic bases, scheme 14. The resulting 48, a surrogate for 2 (scheme 1) was simultaneously expected to result in a cascade of anion-induced ring closures leading to the secododecahedranediol-dione 49.

Scheme 14

O OMe
$$base$$

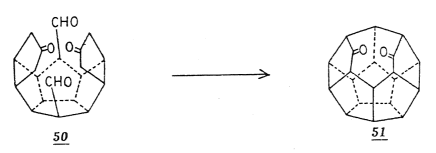
O OMe $base$

O

49

Thus, 47 was exposed to a variety of bases (LDA, Li-HMDS, KH, t-BuOK-t-BuOH etc.) but the outcome was very disappointing. In a majority of these reactions, considerable loss of precious material occurred and no product could be firmly characterised. Consequently, we realised that the choice of diester functionality for effecting the four-fold anionic ring closure needed a change to a more electrophilic group. Presently we are in the process of preparing the hexacyclic diketodialdehyde 50 and hope that it will be possible to entice it into cyclisation to deliver the secondodecahedranedione 51, scheme 15.

Scheme 15



In summary, we have conceived a new, short approach to the $C_{20}H_{20}$ hydrocarbon sphere, dodecahedrane. To attain this formidable objective, a new synthesis of functionalised tetraquinanes based on photo-thermal olefin metathesis reaction has been developed. The Greene methodology has been successfully applied for the *bis*-cyclopentannulation of several tetraquinanes to furnish functionalised hexaquinanes. It has been possible to invert and project the newly appended cyclopentane rings into the spheroidal cavity. Thus, the advanced C_{20} -hexaquinane precursor of dodecahedrane <u>47</u> has been realised. Attempts to convert <u>47</u> to a secododecahedrane derivative *en route* to the target molecule <u>1</u> have so far not succeeded. However, firm ground work has been laid for making repeated assaults on the final objective, which has been appropriately referred to, as the 'Mount Everest of alicyclic chemistry' (Grubmuller 1979).

Acknowledgements

This research has been supported by the Council of Scientific and Industrial Research, Government of India, and the University Grants Commission through a special assistance programme in organic chemistry. KRR and MSN thank CSIR for the award of research fellowships. Finally, we appreciate the timely help of Dr T N Guru Row and his colleagues, NCL, Pune for X-ray crystal structure determination of 47.

References

Astin K B and Mackenzi K 1975 J. Chem. Soc., Perkin I 1004
Bak D A and Brady W T 1979 J. Org. Chem. 44 107
Baldwin J E and Beckwith P L M 1983 J. Chem. Soc., Chem. Commun. 279
Baldwin J E, Beckwith P L M and Wallis J D 1984 J. Chem. Soc. Perkin II 53

Baum M W, Guenzi A, Johnson C A and Mislow K 1982 Tetrahedron Lett. 31

Byrne L T, Rye A R and Wege D 1974 Aust. J. Chem. 27 1961

Carcellar E, Garcia M L, Moyano A, Pericas M and Serratosa F 1986 Tetrahedron 42 1831

Deslongchamps P and Soucy P 1981 Tetrahedron 37 4385

Dixon D A, Deerfield D and Graham G D 1981 Chem. Phys. Lett. 78 161

Eaton P E 1979 Tetrahedron 35 2189

Eaton P E, Bunnele W H and Engel P 1984 Can. J. Chem. 62 2612

Eaton PE, Muller RH, Carlson GR, Cullison DA, Cooper GF, Chou T-C and Krebs E-P 1977 J. Am.

Ermer O 1977 Angew Chem., Int. Ed. Engl. 16 411

Fessner W-D, Murthy B A R C, Worth J, Hunkler D, Fritz H, Prinzbach H, Roth W D, Schleyer P v R, McEwen A B and Maier W F 1987 Angew. Chem., Int. Ed. Engl. 26 452, and references therein Fukunaga T and Clement R A 1977 J. Org. Chem. 42 270

Greene A E and Depress J P 1979 J. Am. Chem. Soc. 101 4003

Grubmuller P 1979 Ph D dissertation, Freidrich-Alexander Universitat, Erlangen-Nurnberg, West

Ito Y, Hirao T and Saegusa T 1978 J. Org. Chem. 43 1011

Jacobson I T 1967 Acta. Chem. Scand. 21 2235

McKervey M A, Vibuljian P, Ferguson G and Siew P Y 1981 J. Chem. Soc., Chem. Commun 912 Mehta G 1978 J. Sci. Ind. Res. (India) 37 256

Mehta G and Nair M S 1983 J. Chem. Soc., Chem. Commun. 439

Mehta G and Nair M S 1985a J. Chem. Soc., Chem. Commun. 629

Mehta G and Nair M S 1985b J. Am. Chem. Soc. 107 7519

Mehta G, Reddy A V and Srikrishna A 1979 Tetrahedron Lett. 4863

Mehta G, Srikrishna A, Reddy A V and Nair M S 1981 Tetrahedron 37 4543

Mincionne E, Ortaggi G and Sirna A 1977 Synthesis 773

Monego T A 1982 Ph D dissertation, Michigan State University

Noyori R, Baba Y and Hayakawa Y 1974 J. Am. Chem. Soc. 96 3336

Paquette L A 1979 Top Curr. Chem. 79 43

Paquette L A 1984 Strategies and tactics in organic synthesis (New York: Academic Press) p. 175 Paquette L A and Miyahara Y 1987 J. Org. Chem. 52 1265

Paquette L A, Nakamura K and Engel P 1986 Chem. Ber. 119 3782

Paquette L A, Ternansky R J, Balogh D W and Kentgen G 1983 J. Am. Chem. Soc. 105 5446 Paquette L A, Wyvratt M J, Berk H C and Moerck R E 1978 J. Am. Chem. Soc. 100 5845 Reich H J 1979 Acc. Chem. Res. 1222

Reich H J, Renga J M and Reich I L 1975 J. Am. Chem. Soc. 97 5434

Repic O 1976 Ph D dissertation, Harvard University

Roberts W P and Shoham G 1981 Tetrahedron Lett. 4895

Schleyer P v R 1957 J. Am. Chem. Soc. 79 3292

Schulman J M and Disch R L 1978 J. Am. Chem. Soc. 100 5677

Schulman J M, Venanzi T and Disch R L 1975 J. Am. Chem. Soc. 97 5335

Sedelmeier G, Fessner W-D, Pinkos R, Grund C, Murthy B A R C, Hunkler D, Rihs G, Fritz H, Kruger C and Prinzbach H 1986 Chem. Ber. 119 3442

Ternansky R J, Balogh D W and Paquette L A 1982 J. Am. Chem. Soc. 104 4503

Woodward R B, Fukunaga T and Kelley R C 1964 J. Am. Chem. Soc. 86 3162