A versatile process for the syntheses of very long chain alkanes, functionalised derivatives and some branched chain hydrocarbons

Gerald M. Brooke,^{*,*a*} Simon Burnett,^{*a*} Shahid Mohammed,^{*a*} David Proctor^{*a*} and Mark C. Whiting^{*b*}

^a Chemistry Department, Science Laboratories, South Road, Durham DH1 3LE, UK

^b Department of Chemistry, University of Bristol, Cantock's Close, Bristol BS8 1TS, UK



An earlier method for synthesising very long straight-chain alkanes of very specific lengths has been improved to give, in some cases, gram amounts of materials. Eleven compounds have been made: $C_{98}H_{198}$, $C_{122}H_{246}$, $C_{162}H_{326}$, $C_{194}H_{390}$, $C_{198}H_{398}$, $C_{210}H_{422}$, $C_{242}H_{486}$, $C_{246}H_{494}$, $C_{258}H_{518}$, $C_{294}H_{590}$ and $C_{390}H_{782}$. The selfcondensation reaction of one of the intermediate aldehydes enabled two chain-branched hydrocarbons to be obtained: $C_{96}H_{193}$ CHRC $_{94}H_{189}$ where $R = CH_3$ and $CH_3(CH_2)_3$. Three long-chain compounds containing carboxylic acid groups have been prepared: $CH_3(CH_2)_{190}CO_2H$, $HO_2C(CH_2)_{48}CO_2H$ and $HO_2C(CH_2)_{192}CO_2H$.

Considerable interest has been shown in recent years in relating the crystallisation processes and crystal morphology of well defined linear long-chain alkanes to that of commercial polyethylene.1 These materials became available in small quantities from the research work of Wegner² and that of Whiting.³ In the German work,² the C₂₄ diyne, HC=C(CH₂)₂₀-C=CH was oxidatively coupled with copper(II) acetate to give a mixture of oligomers, HC=C(CH₂)₂₀C=C[C=C(CH₂)₂₀- $C \equiv C_n H$ (n = 1-16), which were carefully separated by chromatography. As the chain length increased, the separation of higher homologues became more difficult, although this could be offset by coupling the pure dimer, where n = 1. Catalytic hydrogenation gave linear alkanes, of which $C_{\mathbf{384}}H_{770}$ was the longest obtained. In this procedure, chain lengths were limited to multiples of C_{24} and only hydrocarbons could be synthesised.

The basis of Whiting's procedure for the synthesis of long chain alkanes is shown in Scheme 1. The starting material was 12-bromododecanal ethylene acetal 1, which on deprotection gave the aldehyde 3 while reaction with triphenylphosphine gave the phosphonium bromide 5. The reaction of 3, 5 and potassium carbonate in the presence of 18-crown-6 to generate the ylide 7 *in situ* in tetrahydrofuran (THF) gave the chain doubled C_{24} bromo acetal 8. Repetition of the reaction sequence with 8 gave the second chain doubled product, the C_{48} bromo acetal 15, which in turn was converted into the C_{96} bromo acetal 22 (n = 7), and then into the C_{192} bromo acetal 22 (n = 15); the C_{192} acetal 24 (n = 15) was chain doubled to the C_{384} acetal 24 (n = 31).

Deprotection of acetals 22 and 'capping' the resulting aldehydes 25 using a Wittig reagent 29 of any desired length, followed by replacement of bromine in 30 by hydrogen using lithium triethylborohydride (or the reverse sequence) and finally hydrogenation of the polyalkene 31, gave the alkane 32. The chain lengthening reaction was also carried out using moieties having different chain lengths; *e.g.* the C₁₉₂ aldehyde 26 (n = 15) and the C₄₈ acetal ylide 21 gave the C₂₄₀ acetal 24 (n = 19), and ultimately the hydrocarbon C₂₄₆H₄₉₄ which showed chain-folding properties.¹

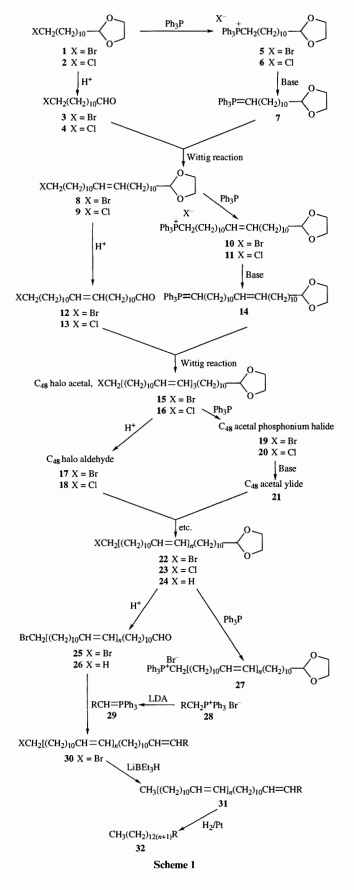
The use of potassium carbonate for generation of the ylides *in* situ was accompanied by a side reaction in which the bromine in some of the product was replaced by carbonate as a nucleophile, producing carbonate esters. Treatment of the reaction product with LiAlH₄ converted carbonate to alcohol which was readily removed by chromatography. In order to maximise the efficiency of the Wittig chain coupling, however, we have

investigated the use of chloro rather than bromo aldehydes in the reaction: primary chlorides are much less reactive towards nucleophiles than the corresponding bromides.

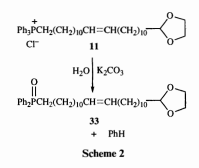
The first perceived major task in the change to 12chlorododecanal ethylene acetal **2** as the basic starting material, however, was to react this primary chloride with triphenylphosphine to form the phosphonium chloride **6**. The C₁₂ chloro acetal **2** and 4 equivalents of triphenylphosphine, when heated and stirred as a melt at 103 °C for 20 days, gave complete removal of the alkyl chloride. Following the work-up, (which included reprotection of some aldehyde which had formed in the procedure), the phosphonium chloride **6** was obtained as a hygroscopic solid (87%) accompanied by the product of hydrolysis of the salt, 12-(diphenylphosphinoyl)dodecanal ethylene acetal, and the β -elimination product, 11-dodecenal ethylene acetal.

The chain doubling reaction involving the C_{12} chloro aldehyde 4 and the C_{12} acetal phosphonium chloride 6 under Whiting's conditions gave very crude C_{24} chloro acetal 9 (86%); accumulated material was carefully distilled under controlled conditions in a short path distillation unit, enabling large quantities of the pure compound 9 to be obtained with ease. In the original work, the assessment of purity was carried out using high temperature analytical vapour phase chromatography, but it was clear that this method would not be applicable to the involatile C48 chloro acetal 16 and higher homologues. An HPLC analysis of the 2,4-dinitrophenylhydrazones (DNPs) of the haloaldehydes derived directly from the acetals has been developed, and represents an important advance in determining the effectiveness of the purification processes undertaken: the DNPs derived from the C_{12} , C_{24} , (and from higher homologues which were prepared later in the work) C₃₆, C₄₈, C₇₂, C₉₆, C₁₄₄ and C192 haloacetals (Cl or Br), were all well resolved from each other. Potential impurities in the C_{24} chloro acetal 9 were the C_{12} chloro aldehyde 4, and the C_{36} chloro acetal 23 (n = 2) formed via deprotection of 9 followed by further reaction with C_{12} acetal ylide 7; both were shown to be absent in the distilled product.

The reaction of the C_{24} chloro acetal 9 with triphenylphosphine required 28 days to give the C_{24} acetal phosphonium chloride 11, which was hygroscopic, and so was stored as a standard solution in anhydrous tetrahydrofuran. The reaction of the ylide 14 from this salt with the C_{24} chloro aldehyde 13 under the original conditions gave only ca. 1% C_{48} chloro acetal 16, and it became clear that the use of potassium carbonate, in our hands, under long reaction times (> 200 h) brought about



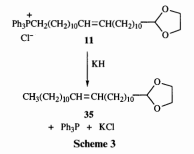
self-condensation reactions of the aldehyde 13, and also hydrolytic cleavage of benzene from the salt 11 to give the unreactive diphenylphosphinoacetal 33 (Scheme 2); water is inevitably formed in any base-catalysed reaction with potassium carbonate. These problems were overcome by preforming the ylide 14, as in a conventional Wittig reaction, using



potassium hydride as the base with 18-crown-6 again as the phase-transfer agent in refluxing tetrahydrofuran. The progress of the reaction was followed by ³¹P NMR spectroscopy until no phosphonium salt remained, and then the chloroaldehyde **13** was added to the deep orange–black solution of the ylide. The yields of C₄₈ chloro acetal **16** were > 80%, and the product was of high purity [free from **13** and C₇₂ chloro acetal **23** (n = 5) by HPLC of DNPs]. However, two minor by-products formed from **11** by loss of Ph₃P were identified in the crude product: (a) the vinyl acetal **34** (from an elimination reaction), a compound

$$CH_2 = CH(CH_2)_9[CH = CH(CH_2)_{10}]_3 - \begin{pmatrix} 0 \\ 0 \end{pmatrix}$$

which would present no problems in subsequent chain-doubling reactions; it would merely appear as a terminal vinyl-containing contaminant of the same overall chain length as the major chain extended product from the Wittig reaction; and (b) the terminal hydrocarbon acetal **35** formed by nucleophilic replacement by hydrogen (Scheme 3) rationalised the presence. These two processes of triphenylphosphine (detected by ³¹P



NMR) accompanying the formation of the ylide. This would cause contamination of the final products with shorter chain hydrocarbons. Removal of the C_{24} acetal 35 from the C_{48} chloro acetal 16 was achieved by a combination of sublimation at 190 °C and 0.05 mmHg, and column chromatography, but the same problem was expected to arise in subsequent chaindoubling reactions where purification using sublimation would not be applicable: only very difficult chromatography for the removal of this non-halogen containing compound could be used. Consequently, we sought to find an alternative powerful base to generate the ylide, and yet which was also nonnucleophilic. Preliminary experiments with lithium diisopropylamide (LDA) fulfilled these requirements admirably. Moreover, the ¹³C NMR spectrum of the C₂₄ chloro acetal 9 from 2 and 6 showed a high *cis* to *trans* ratio of 4:1, similar to the material prepared in the original potassium carbonate-induced Wittig reaction,² so that potential problems of compound solubility with higher homologues were not expected. In general, ylide formation is essentially instantaneous at -10 to -25 °C, as is the subsequent reaction with aldehydes.

The reaction of the C_{48} chloro acetal **16** with triphenylphosphine at 100–103 °C to form the phosphonium chloride **20** was incomplete after 60 days. Consequently, half of the stock of **16**

Table 1 'Capping' reactions to form bromo polyenes and polyenes

Aldehyde	Ylide	Product
C_{96} bromo aldehyde 25 ($n = 7$)	C_2 ylide 29 (R = CH ₃)	C_{98} bromo polyene 46
C_{120} bromo aldehyde 25 ($n = 9$)	C_2 ylide 29 ($R = CH_3$)	C_{122} bromo polyene 47
C_{96} bromo aldehyde 25 ($n = 7$)	C ₆₆ ylide 45	C_{162} bromo polyene 48
C_{192} bromo aldehyde 25 ($n = 15$)	C_2 ylide 29 (R = CH ₃)	C_{194} bromo polyene 49
C_{192} bromo aldehyde 25 ($n = 15$)	C_{18} ylide 43	C_{210} bromo polyene 50
C_{192} aldehyde 26 ($n = 15$)	C_{50} ylide 44	C_{242} polyene 51
C_{192} bromo aldehyde 25 ($n = 15$)	C_{66} ylide 45	C_{258} bromo polyene 52
C_{384} aldehyde 26 ($n = 31$)	C_6 ylide 29 (R = C_5H_{11})	C ₃₉₀ polyene 53
$CH_{3}(CH_{2})_{4}CH=CH(CH_{2})_{10}CH_{2}X$ 37 X = Br	$CH_{3}[CH=CH(CH_{2})_{10}]_{4}CH_{2}X$ $39 X = Br$	$CH_{3}(CH_{2})_{4}[CH=CH(CH_{2})_{10}]_{5}CH_{2}X$ 41 X = Br
$38 X = P^+ Ph_3 Br^-$	$40 X = P^+ Ph_3 Br^-$	$42 X = P^+ Ph_3 Br^-$
CH ₃ (CH ₂) ₄ CH=CH(CH ₂) ₁₀ CH=PPh ₃	CH ₃ [CH=CH(CH ₂) ₁₀] ₄ CH=PPh ₃	CH ₃ (CH ₂) ₄ [CH=CH(CH ₂) ₁₀] ₅ CH=PP

44

which had been prepared was converted into the C_{48} bromo acetal 15 using tetrabutylammonium bromide in 1-bromopropane,³ for reaction with triphenylphosphine in acetonitrile, which with the addition of 2,2-dimethyl-1,3-dioxolane, formed the phosphonium bromide 19 unaccompanied by any deprotected acetal, in only 3.5 days. Unlike the chlorides 6 and 11, the phosphonium bromides 5, 10, 19 and, subsequently, all the bromides were non-hygroscopic, and so presented no difficulties in handling.

43

The C₉₆ chloro acetal **23** (n = 7) was prepared in 77% yield from the C_{48} chloro aldehyde 18 and the C_{48} acetal ylide 21 generated from bromide 19 and LDA. All of this material was subjected to the trans-halogenation reaction with tetrabutylammonium bromide to give the C_{96} bromo acetal 22 (n = 7) so that future reactions could be carried out using bromo compounds only, as in the original Whiting procedure.³ The reaction did not proceed as expected, due to contamination of the solvent/reagent by water, and faulty technique. The product consisted of the C₉₆ bromo aldehyde 25 (n = 7) (3 parts) and the self-condensation product of this compound 36 (2 parts), readily identified by ¹H NMR spectroscopy. Closer inspection of the spectra of both of these materials showed that ca. 25%contained the equivalent of the addition of one molecule of HBr to one C=C bond in the C₉₆ chain to form CHBr. The mixture was subjected to reprotection of the aldehyde functions with ethylene glycol-toluene-p-sulfonic acid: only the saturated aldehyde 25 (n = 7) reacted. The bromo acetal 22 (n = 7) was separated from the α,β -unsaturated aldehyde 36 by

$$\begin{array}{c} BrCH_{2}[(CH_{2})_{10}CH=CH]_{7}(CH_{2})_{10}\\ CH=C(CHO)(CH_{2})_{9}[CH=CH(CH_{2})_{10}]_{7}CH_{2}Br\\ 36\end{array}$$

chromatography for subsequent reactions. Triphenylphosphine reacted exclusively with the primary alkyl bromide group of 22 (n = 7) to give the C₉₆ phosphonium bromide 27 (n = 7): a model experiment of triphenylphosphine with the secondary bromide 6-bromododecane occurred very slowly, and the characteristic shift of the CH-P⁺Ph₃ was absent from 27 (n = 7), so no branched chain impurities could be introduced subsequently.

The chain doubled compound, the C_{192} bromo acetal 22 (n = 15) was formed in 77% yield from the C_{96} bromo aldehyde 25 (n = 7) and the ylide derived from the C_{96} acetal phosphonium bromide 27 (n = 7). The bromo acetal 22 (n = 15) was deprotected and the resulting bromo aldehyde 25 (n = 15) was 'capped' with the C₆ unit 29 $(R = C_5H_{11})$. Treatment of the C_{198} bromo polyene 30 $(n = 15, R = C_5H_{11})$ with lithium triethylborohydride gave the C_{198} polyene 31 $(n = 15, R = C_5H_{11})$ still containing the secondary bromide functionality: this was removed by dehydrobromination with 1,8-diaza-

bicyclo[5.4.0]undec-7-ene (DBU) and the product hydrogenated to give the known hydrocarbon³ $C_{198}H_{398}$ **32** (n = 15, $R = C_5H_{11}$), in much larger quantities.

45

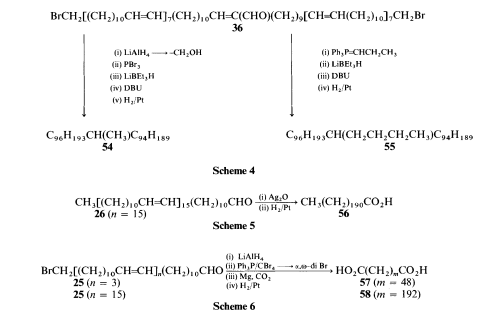
The C₁₉₂ bromo aldehyde **25** (n = 15) underwent Wittig reactions with the C₄₈ acetal ylide **21** and with the ylide from the C₉₆ acetal phosphonium bromide **27** (n = 7) to give the C₂₄₀ bromo acetal **22** (n = 19) and the C₂₈₈ bromo acetal **22** (n = 23), respectively. Both in turn were 'C₆-capped', hydrodebrominated, dehydrobrominated and hydrogenated as before to give the hydrocarbons³ C₂₄₆H₄₉₄ **32** (n = 19), R = C₅H₁₁) and C₂₉₄H₅₉₀ **32** $(n = 23, C_5H_{11})$, respectively.

The work described above was carried out starting with 12chlorododecanal ethylene acetal 2 derived from 1 kg of 12chlorododecanol via a Swern oxidation to the aldehyde 4 which was then converted to the stable 12-chlorododecanal acetal 2;3 on standing at room temperature, the aldehyde 4 began to trimerise within 24 h, so it was always used as quickly as possible in Wittig reactions. New work was started with 2 derived from 3 kg of 12-chlorododecanol. It was first converted to 12-bromododecanal ethylene acetal 1 by the transhalogenation reaction,³ and then chain extension reactions were carried out to give C24, C36, C48, C96 and C192 bromo acetals as before, while reaction of the C_{96} bromo aldehyde 25 (n = 7) with the C₂₄ acetal ylide 14 gave the C₁₂₀ bromo acetal 22 (n = 9). The C₃₈₄ acetal³ 24 (n = 31) was formed from the C_{192} aldehyde 26 (n = 15) and the ylide from the C_{192} acetal phosphonium bromide 27 (n = 15).

Three new primary bromides were prepared for incorporation into 'capping' agent precursors: (a) the C₁₈ bromo compound 37 from the C₁₂ bromo aldehyde 3 and the C₆ ylide 29 (R = C₅H₁₁); with triphenylphosphine it gave the C₁₈ phosphonium bromide 38, *i.e.* 28 [R = CH₃(CH₂)₄CH=CH(CH₂)₁₀]; (b) the C₅₀ bromo compound 39 from the C₄₈ bromo aldehyde 17 and the C₂ ylide 29 (R = CH₃); it was converted into the C₅₀ phosphonium bromide 40, *i.e.* 28 {R = CH₃[CH=CH-(CH₂)₁₀]₄} with triphenylphosphine; and (c) the C₆₆ bromo compound 41 from the C₄₈ bromo aldehyde 17 and the C₁₈ ylide from 38; it gave the C₆₆ phosphonium bromide 42, *i.e.* 28 {R = CH₃(CH₂)₄[CH=CH(CH₂)₁₀]₅} with triphenylphosphine. The three new 'capping' agents 43, 44 and 45, and their use, are given in Table 1.

Bromopolyenes were converted to polyenes with lithium triethylborohydride, and the resulting polyenes hydrogenated. Seven new straight chain hydrocarbons were obtained: C_{98} - H_{198} , $C_{122}H_{246}$, $C_{162}H_{326}$, $C_{194}H_{390}$, $C_{210}H_{422}$, $C_{242}H_{486}$ and $C_{258}H_{518}$ as they were required for the physical study of phase changes;⁴ $C_{390}H_{782}$ (12 mg) had been prepared earlier.³ Two branched chain hydrocarbons $C_{96}H_{193}$ CH(CH₃)-Two branched chain hydrocarbons $C_{96}H_{193}$ CH(CH₃)-

Two branched chain hydrocarbons $C_{96}H_{193}CH(CH_3)$ - $C_{94}H_{189}$ 54 and $C_{96}H_{193}CH(CH_2CH_2CH_2CH_3)C_{94}H_{189}$ 55 were prepared from the branched C_{192} dibromo α,β unsaturated aldehyde 36, as shown in Scheme 4. Although



the unsaturated aldehyde was prepared serendipitously, this condensation reaction should be general for the synthesis of singly-branched long chain alkanes.

Finally, the versatility of the procedures described in this paper is illustrated by the possibility of being able to convert the α -bromine and the ω -acetal groups in the α , ω -bromoacetals 22 into other functionalities. Thus, we have synthesised the very long chain monocarboxylic acid CH₃(CH₂)₁₉₀CO₂H 56 by the route shown in Scheme 5. Two dicarboxylic acids have also been prepared: HO₂C(CH₂)₄₈CO₂H 57 and HO₂C(CH₂)₁₉₂-CO₂H 58 (Scheme 6).

Experimental

NMR spectra were recorded on the following instruments at the frequencies listed: a Varian VXR 400S [1H (399.952 MHz), 13C (100.582 MHz)], and a Bruker AMX 500 [¹H (500.139 MHz)]. Absorption multiplicities have been abbreviated as follows: s (singlet), d (doublet), t (triplet), q (quintet) and m (multiplet). All chemical shifts are given in ppm with respect to TMS, present in CDCl₃ used as solvent. Silica refers to Merck silica gel F60 (230-400 mesh). Analytical HPLC was performed on a Star 5065 instrument fitted with Hypersil 5 SAS 25 cm \times 4.6 mm C₁ reverse phase column. Elemental analyses were performed on an Exeter Analyical Inc CE440 elemental analyser. Melting points were determined on a Gallenkamp melting point apparatus, unless otherwise stated. Purification of C_{12} and C_{24} materials was carried out by short path distillation as described, using a short path distillation unit (KDL-1), manufactured by U.I.C. GmbH Alzenau, Germany. All reaction solvents were dried in the appropriate manner: acetonitrile and dichloromethane were freshly distilled from P2O5; THF was distilled from LiAlH4 and NaH immediately prior to use. All light petroleum solvents were distilled on a rotary evaporator prior to use. The fraction distilling in the range 40-60 °C was used unless otherwise stated. 'Ether' refers to diethyl ether.

NMR analyses

¹H NMR was used in the identification of all compounds. After each chain doubling or chain extension reaction, the -CH =CH- to $-CH(OCH_2)_2$ (ethylene acetal group) ratio increases. This ratio was used to identify a material's chain length, although accurate measurement became more difficult as the number of carbon–carbon double bonds increased. Due to the difficulty in removing the last traces of light petroleum from some products, measurement of the intensity of the $-CH_2$ signal in the hydrocarbon chain proved less accurate. Integral measurements for the final alkanes also proved to be inaccurate, due to difficulties in shimming the spectrometer at 120 °C so no reliable ratio was achieved between the $-CH_3$ end groups and the internal $-CH_2$ - groups; the ¹H NMR could only be used to demonstrate the absence of carbon-carbon double bonds.

Preparative chromatography and HPLC analysis of 2,4-dinitrophenylhydrazone derivatives

After each chain doubling or chain extension reaction, the resulting α, ω -haloacetals were separated from the slowereluting precursors by preparative chromatography on silica. Nevertheless, it was vital to be able to assess the effectiveness of the separations. ¹H NMR analysis is not sensitive to low levels of contamination by homologous compounds, so the 2,4dinitrophenylhydrazone (DNP) derivative of the product was prepared and analysed by HPLC to determine the purity, since homologues of these were well separated.

The following method was used to prepare all DNP derivatives. The bromoacetal (0.1 g) was stirred with 2,4-dinitrophenylhydrazine (0.2 g) and concentrated sulfuric acid (0.5 ml) in butyl alcohol (10 ml) for 16 h at 45 °C. On cooling to room temperature, the orange DNP derivative precipitated, which was filtered, washed with methanol and dried.

Materials

12-Chlorododecanol (4 kg) was synthesised and supplied by High Force Research Ltd., Unit 1D, Mountjoy Research Centre, Stockton Road, Durham City, DH1 3SW, using the published route.^{3,5} 12-Chlorododecanal ethylene acetal 2 was prepared in 83% overall yield by the Swern oxidation of the 12chloro alcohol in 200 g batches followed by acetalisation and short path distillation at 85 °C and 4×10^{-2} mbar.³ 12-Bromododecanal ethylene acetal 1 was prepared from the chloroacetal 2 by refluxing with tetrabutylammonium bromide in 1-bromopropane which had been dried beforehand by heating under reflux in a Soxhlet unit containing molecular sieves (4 Å).⁴ After 42 h, ¹H NMR showed 9% of unreacted 2, and 15% of aldehyde present. Reprotection of the aldehyde with ethylene glycol/toluene-p-sulfonic acid and a further transhalogenation reaction gave complete conversion of 2, but again, reprotection of aldehyde had to be carried out (the addition of 2,2-dimethyl-1,3-dioxolane would prevent deprotection from occurring). Pure 12-bromododecanal ethylene acetal 1 was obtained by short path distillation at 80 °C and 6 \times 10⁻³ mbar.

C₁₂ Phosphonium salts

11-(Dioxolan-2-yl)undecyl(triphenyl)phosphonium chloride (the C₁₂ acetal phosphonium chloride) 6. The 12-chloro acetal 2 (200 g) and triphenylphosphine (800 g) were heated at 102.5-103.5 °C and the melt stirred for 20 d. Acetonitrile (700 ml) was slowly added to the hot mixture, and when the solution had cooled to ca. 45 °C, light petroleum and acetonitrile were used to transfer the mixture to a separating funnel. The acetonitrile layer was washed with light petroleum (16 \times 100 ml) and the combined petrol extracts were re-extracted with acetonitrile $(4 \times 50 \text{ ml})$. Removal of the solvent in vacuo from the combined acetonitrile extracts gave the crude phosphonium salt (456 g), the ¹H NMR of which showed the presence of deprotected acetal (13%). Reprotection of the aldehyde was carried out by heating a mixture of the crude salt, ethylene glycol (60 ml), toluene-p-sulfonic acid (3.1 g) and dichloromethane (600 ml) at reflux temperature overnight. The reaction mixture was cooled to room temperature, washed with dilute aqueous potassium carbonate $(2 \times 1 \text{ l containing 3 g of the})$ base), followed by water (2 \times 1.5 l); in the latter case, it was necessary to add a trace of potassium carbonate after each wash to break up the emulsion which formed. The organic phase was dried (Na₂SO₄), and the solvent removed in vacuo. Dry ether (1 1) was added to the viscous liquid which was stirred vigorously; the mixture was heated to boiling and left to cool overnight. The upper layer was decanted and replaced by more ether (1 l) and the procedure repeated again. This was carried out four more times. After the final decantation, there remained a white solid which was dried under high vacuum to give the C_{12} acetal triphenylphosphonium chloride 6 (351 g, 87%), a hygroscopic material, identified by comparison of its ¹H NMR spectrum with that of the authentic C_{12} acetal phosphonium bromide. Examination of the ether washings by ¹H NMR spectroscopy showed the presence of 12-diphenylphosphinoyldodecanal ethylene acetal. Evaporation of the petrol extracts, which removed the triphenylphosphine, and careful sublimation of the residue at 35 °C, gave 11-dodecenal ethylene acetal; $\delta_{\rm H}$ 5.80 (m, H_a), 4.88 (d, H_c) and 4.87 (s, H_b) in $H_bH_{c,trans}$ C=CH_{a,trans}.

11-(Dioxolan-2-yl)undecyl(triphenyl)phosphonium bromide³ (the C_{12} acetal phosphonium bromide) 5. Compound 5 was prepared from 12-bromododecanal ethylene acetal 1 triphenylphosphine (1.1 equiv.) and 2,2-dimethyl-1,3-dioxolane (0.1 equiv.) in anhydrous acetonitrile at reflux temperature for 72 h under nitrogen and worked up as before.

Wittig reactions using aldehydes, phosphonium salts and $K_2CO_3/18$ -crown-6 *in situ*

24-Chlorotetracos-12-enal ethylene acetal³ (the C_{24} chloro acetal) 9. This compound was prepared as before, using the C_{12} phosphonium chloride rather than the bromide salt. Purification of the C_{24} chloro acetal was accomplished by four passes through the short path distillation unit, two at 105 and one at 115 °C, 1×10^{-2} mbar, to remove C_{12} components, and once at 140 °C, at 1×10^{-2} mbar, to distil the C_{24} chloro acetal from longer chain compounds, principally the C_{36} chloro acetal. HPLC analysis of the product (70% THF, 30% water), showed that it was exclusively the C_{24} chloro acetal, the ¹H NMR spectrum of which was identical with that of an authentic sample.

23-(Dioxolan-2-yl)tricos-12-enyl(triphenyl)phosphonium

chloride (the C_{24} acetal phosphonium chloride) 11. Compound 11 was prepared from the C_{24} chloro acetal 9 and triphenylphosphine as before over 28 d. Its ¹H NMR spectrum was identical with that of the authentic phosphonium bromide.³ Once again there was NMR evidence of terminal CH₂=CH– and Ph₂P(O)CH₂– by-products being formed. The salt was very hygroscopic and was dissolved in anhydrous tetrahydrofuran as a standard solution for use in Wittig reactions.

36-Chlorohexatriaconta-12,24-dienal ethylene acetal (the C_{36} chloro acetal) 23 (n = 2). The C_{24} chloro aldehyde 13 was

reacted as before with the C₁₂ acetal phosphonium chloride **6** and to give the *product* **23** (n = 2), isolated by chromatography on silica using light petroleum–ether (85/15, v/v); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 2 × (CH=CH)], 4.84 [t, CHO₂(CH₂)₂], 3.9 [m, O(CH₂)₂O] and 3.5 (t, CH₂Cl), in the ratio 4.4: 1.0:4.2:2.0. It formed a 2,4-*dinitrophenylhydrazone*, mp 44–48.5 °C (Found: C, 68.65; H, 9.85; N, 7.3. C₄₂H₇₁ClN₄O₄ requires C, 68.96; H, 9.78; N, 7.66%).

Wittig reactions using KH and phosphonium salts to pregenerate the vlide

48-Chlorooctatetraconta-12,24,36-trienal ethylene acetal (the C_{48} chloro acetal) 16. A solution of the C_{24} phosphonium chloride 11 (17.76 g) in tetrahydrofuran (40 ml), 18-crown-6 (0.805 g) and potassium hydride (1.198 g) was heated and stirred at 55 °C for 48 h. Examination of a small sample of the solution by ${}^{31}P$ NMR indicated 85% ylide (δ 12.6), 13% Ph₂P(O)CH₂-material (δ 27.2), 1% unreacted phosphonium chloride (δ 23.9) and 1% triphenylphosphine (δ -5.1). The C₂₄ chloro aldehyde 13 (8.49 g) was added to the ylide and heating was continued for 2 h. The mixture was washed through a short column of silica using light petroleum-ether (the new work-up procedure) to give crude product which was chromatographed on silica, (loading ca. 10 g of the material onto 1 kg silica) using light petroleum-ether (90:10, v/v) as eluent to give pure 48chlorooctatetraconta-12,24,36-trienal ethylene acetal 16, mp 41-43 °C (Found: C, 78.4; H, 12.8. C₅₀H₉₃ClO₂ requires C, 78.83; H, 12.31%); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 3 × (CH=CH)], 4.84 [t, $CHO_2(CH_2)_2$], 3.9 [m, O(CH₂)₂O] and 3.53 (t, CH₂Cl); signals were in the ratio 6.32:1:4.18:2.04. HPLC of the DNP derivative (70% THF, 30% water) indicated it was free from C₂₄ and C₇₂ materials. Fractions containing a slower eluting component were collected and heated in vacuo with stirring at 180 °C and 0.05 mmHg, volatile material being collected on a water-cooled finger. Examination of this crude sublimate by ¹H NMR showed the presence of only a very small proportion of $-CH_2Cl$ (at ca. δ 3.5), but, significantly, a large CH₃- triplet at *ca.* δ 0.9, in proportion to the alkene and acetal features, which indicated the major presence of tetracos-12-enal ethylene acetal 35.

48-Bromooctatetraconta-12,24,36-trienal ethylene acetal³ (the C_{48} bromo acetal) 15. Compound 15 was prepared by a halogen exchange reaction on the chloro compound 16 using the method described above for 12-bromododecanal ethylene acetal 1.

47-(Dioxolan-2-yl)heptatetraconta-12,24,36-trienyl(triphenyl)phosphonium bromide³ (the C₄₈ acetal phosphonium bromide) 19. Compound 19 was prepared from the C₄₈ bromo acetal 15, triphenylphosphine and 2,2-dimethyl-1,3-dioxolane (5 ml) in acetonitrile at reflux temperature over 112 h and worked up as before. The product 19 (75%) was analysed by ¹H NMR and shown to be identical to an authentic sample.

72-Chlorodoheptaconta-12,24,36,48,60-pentaenal ethylene acetal (the C_{72} chloro acetal) 23 (n = 5). A crude sample was prepared using the C_{48} chloro aldehyde 18 and the C_{24} acetal phosphonium chloride 11, and was converted into the DNP derivative for use as a standard for purity checks by HPLC.

Wittig reactions using lithium diisopropylamide (LDA) and phosphonium salts to pregenerate the ylide

24-Bromotetracos-12-enal ethylene acetal³ (the C_{24} bromoacetal) 8. The C_{12} acetal triphenylphosphonium bromide 5 (302 g, 0.530 mol, 1.1 equiv.) and THF (700 ml) were cooled to -10 °C whilst stirring under nitrogen. Lithium diisopropylamide (LDA) (1.5 mol l⁻¹ in THF; 337 ml, 0.506 mol, 1.05 equiv.) was added, giving a deep orange-red colour, characteristic of the ylide 7, and the mixture was stirred for 0.5 h at -10 °C. The temperature was held at -10 °C whilst a solution of the C_{12} bromo aldehyde 3 (127 g, 0.482 mol, 1 equiv.) in THF (10 ml) was added with stirring, causing the colour to fade, leaving a very pale cream-yellow mixture. This was allowed to reach room temperature over 16 h. Water

J. Chem. Soc., Perkin Trans. 1, 1996 1639

(20 ml) was added to the reaction mixture and the THF removed under vacuum leaving a slurry which was dissolved in acetonitrile (500 ml) and extracted with light petroleum $(7 \times 250 \text{ ml})$. The combined petrol layers were washed with water $(3 \times 250 \text{ ml})$, the aqueous layer being extracted with further light petroleum in each case. The petrol layers were combined, dried over sodium sulfate and the solvent was removed under vacuum, leaving an orange oil. This was passed through a short pad of silica (500 g) using light petroleumdiethyl ether (90:10, v/v) to remove any remaining phosphorus compounds and/or lithium salts. The resulting yellow oil (215 g) was purified by short path distillation to remove C_{12} impurities (3 distillations at 100 °C, 4 × 10⁻³ mbar, conditions under which the C_{12} chain lengths distilled, leaving C_{24} and C_{36} material) and C_{36} impurities (1 distillation at 155 °C, 4 \times 10⁻³ mbar, conditions under which C₂₄ compounds distilled, leaving C_{36} material in the residue). The resulting colourless oil 8 (138 g, 60%) formed a white crystalline solid at room temperature, the ¹H NMR of which was identical with that of an authentic sample. Analysis of the DNP derivative by HPLC (70% THF, 30% water) showed less than 0.2% of C_{12} material to be present and no C_{36} impurity.

23-(Dioxolan-2-yl)tricos-12-enyl(triphenyl)phosphonium bromide³ (the C_{24} acetal phosphonium bromide) 10. The C_{24} bromo acetal 8, triphenylphosphine, 2,2-dimethyl-1,3-dioxolane and acetonitrile were reacted over 66 h and worked-up as before to give the product 10, the ¹H NMR of which was identical with that of an authentic sample.

36-Bromohexatriaconta-12,24-dienal ethylene acetal (the C_{36} bromo acetal) 22 (n = 2). The ylide 7 from the C_{12} triphenylphosphonium bromide 5 and LDA was treated with the C_{24} bromo aldehyde 12 as before, to give the *product* 22 (n = 2), isolated by chromatography on silica using light petroleum-diethyl ether (90:10, v/v); $\delta_{\rm H}(\rm CDCl_3)$ 5.35 [m, $2 \times (\rm CH=\rm CH)$], 4.85 [t, $\rm CHO_2(\rm CH_2)_2$], 3.9 [m, $\rm O(\rm CH_2)_2O$] and 3.4 (t, $\rm CH_2Br$), in the ratio 4.19:1.02:4:2.00. The DNP derivative was prepared and used as a standard for HPLC analysis.

48-Bromooctatetraconta-12,24,36-trienal ethylene acetal³ (the C₄₈ bromo acetal) **15.** The ylide **14** from the C₂₄ triphenylphosphonium bromide **10** and LDA was treated with the C₂₄ bromo aldehyde **12** as before to give the product **15** (58%), isolated by chromatography on silica using light petroleum-diethyl ether (90:10, v/v). The DNP derivative was prepared and analysed by HPLC which showed it was free from C₂₄ and C₇₂ materials.

72-Bromodopheptaconta-12,24,36,48,60-pentaenal ethylene acetal (the C₇₂ bromo acetal) **22** (n = 5). The ylide **14** from the C₂₄ triphenylphosphonium bromide **10** and LDA was reacted with the C₄₈ bromo aldehyde **17** as before to give the product, separated by chromatography on silica using light petroleum– diethyl ether (90:10, v/v); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 5 × (CH=CH)], 4.85 [t, CHO₂(CH₂)₂], 3.9 [m, O(CH₂)₂O] and 3.4 (t, CH₂Br), in the ratio 9.05:1.05:4:1.91. The DNP derivative was prepared and used as a standard for HPLC analysis.

96-Chlorohexanonaconta-12,24,36,48,60,72,84-heptaenal ethylene acetal (the C₉₆ chloro acetal) 23 (n = 7). The ylide 21 from the C₄₈ triphenylphosphonium bromide 19 and LDA was treated with the C₄₈ chloro aldehyde 18. The work-up procedure was modified by filtering the products from the Wittig reaction through a short column of silica using light petroleum-diethyl ether (90:10, v/v), followed by loading *ca*. 10 g of the C₉₆ chloro acetal onto 1 kg silica, and chromatography using light petroleum-ether (90:10, v/v) as eluent, to give 96-*chlorohexanonaconta*-12,24,36,48,60,72,84*heptaenal ethylene acetal* 23 (n = 7) (76%), mp 41–43 °C (Found: C, 82.6; H, 13.1. C₉₈H₁₈₁ClO₂ requires C, 82.49; H, 12.78%); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 7 × (CH=CH)], 4.84 [t, CHO₂(CH₂)₂], 3.9 [m, O(CH₂)₂O] and 3.53 (t, CH₂Cl), in the ratio 14.29:1:4.03:1.95. HPLC analysis of the DNP derivative indicated the presence of *ca.* 0.7% C_{48} contamination and no C_{144} impurities.

96-Bromohexanonaconta-12,24,36,48,60,72,84-heptaenal ethylene acetal³ (the C₉₆ bromo acetal) 22 (n = 7). Via Wittig reaction. This compound was prepared as in the previous experiment only using the C₄₈ bromo aldehyde 17 instead of 18, and was identified by ¹H NMR. HPLC analysis of the DNP derivative showed no C₄₈ or C₁₄₄ materials present.

Via halogen exchange. The C₉₆ chloro acetal 23 (n = 7) (67 g) was reacted with tetrabutylammonium bromide in 1bromopropane as described above for the reaction with the C12 chloro acetal 2. After 120 h, 10% -CH₂Cl remained, and deprotected acetal (i.e. aldehyde) was present. Further treatment with tetrabutylammonium bromide in 1-bromopropane gave a product which mainly consisted of two components: the C_{96} bromo aldehyde 25 (n = 7) [*i.e.* deprotected C_{96} bromo acetal 22 (n = 7)] and 2-(94-bromotetranonaconta-10,22,-34,46,58,70,82-heptaenyl)-98-bromooctanonaconta-2,14,26,38,-50,62,74,86-octaenal (the C_{192} dibromo- α,β -unsaturated aldehyde) 36, readily identified from its ¹H NMR spectrum. The mixture was treated with ethylene glycol and toluene-p-sulfonic acid monohydrate in refluxing bromoethane: only the C₉₆ bromo aldehyde underwent conversion to acetal. The two components were separated by chromatography on silica (1 kg) with a loading of ca. 4 g of mixture, using light petroleumether (90:10, v/v) as eluent. The ¹H NMR of the slower moving component was identical with that of an authentic sample of the C_{96} bromo acetal 22 (n = 7), but with an additional bromomethylene signal; $\delta_{\rm H}({\rm CDCl_3})$ 5.36 [m, $7 \times (CH=CH)$], 4.84 [t, $CHO_2(CH_2)_2$], 4.02 (tt, CHBr), 3.9 $[m, O(CH_2)_2O]$ and 3.4 (t, CH_2Br) were in the ratio 14.6:1.06:4.16:2.0, after making the allowance for the loss of one -CH=CH- for each -CHBr-. The secondary bromide functionality was identified by comparison of a model bromide made by reacting the C24 bromo acetal 8 with hydrobromic acid (48%) in a sealed tube at 105 °C.

The faster moving component, 2-(94-bromotetranonaconta-10,22,34,46,58,70,82-heptaenyl)-98-bromooctanonaconta-2,14,-26,38,50,62,74,86-octaenal (the branched C_{192} dibromo- α , β unsaturated aldehyde) **36a** had the characteristic features of -CH₂CH=C(CHO)- group; $\delta_{\rm H}$ (CDCl₃) 9.37 (CHO), 6.45 [t, -CH₂CH=C(CHO)], 5.36 [m, 14 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 14.8:2.0, after correcting for the secondary -CHBr- (at δ 4.02).

95-(Dioxolan-2-yl)pentacontanona-12,24,36,48,60,72,84-heptaenyl(triphenyl)phosphonium bromide³ (the C₉₆ acetal phosphonuium bromide) 27 (n = 7). This compound, completely free from secondary bromide (see below), was prepared by the previously described method using a C₉₆ bromo acetal 22 (n = 7): triphenylphosphine: 2,2-dimethyl-1,3-dioxolane ratio of 1:4:0.1, respectively. The excess triphenylphosphine was separated from the product by chromatography on silica using dichloromethane-methanol (98:2, v/v) as eluent, and the salt was eluted with dichloromethane-methanol (95:5, v/v); $\delta_{\rm H}$ (CDCl₃) 7.75 (m, 3 × C₆H₅), 5.35 [m, 7 × (CH=CH)], 4.85 [t, CHO₂(CH₂)₂], 3.9 [m, O(CH₂)₂O] and 3.7 (m, CH₂P⁺Ph₃) in the ratio 15.2:14.7:0.95:4:1.75.

The C₉₆ bromo acetal **22** (n = 7) described above, which contained secondary bromide functionality, reacted with triphenylphosphine exactly as in the previous experiment, exclusively at the primary alkyl bromide site even though excess triphenylphosphine was used. The ¹H NMR was identical with that of **27** (n = 7), apart from the additional signal at δ 4.02 (tt, CHBr). There were no signals in the region δ 4.2–4.6 [expected for –CH(P⁺Ph₃)–, which were found in a sample obtained by the reaction of 6-bromododecane with triphenylphosphine in acetonitrile and isolated by chromatography on silica using dichloromethane–methanol (80:20, v/v) as eluent].

120-Bromocoshecta-12,24,36,48,60,72,84,96,108-nonaenal ethylene acetal (the C_{120} bromo acetal) 22 (n = 9). Treatment of

the C₉₆ bromo aldehyde **25** (n = 7) with the ylide **14** from the C₂₄ phosphonium bromide **10** gave the C₁₂₀ bromo acetal **22** (n = 9), mp 41–42 °C (Found: C, 81.2; H, 12.7. C₁₂₂H₂₂₅BrO₂ requires C, 81.23; H, 12.57%); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 9 × (CH=CH)], 4.84 [t, CHO₂(CH₂)₂], 3.9 [m, O(CH₂)₂O] and 3.53 (t, CH₃Br) in ratio 18.8:1.0:4.1:2.0.

144-Chlorotetratetracontahecta-12,24,36,48,60,72,84,96,108,-120,132-undecaenal ethylene acetal (the C₁₄₄ chloro acetal) 23 (n = 11). A sample of this compound was prepared as in the previous synthesis using the C₉₆ chloro aldehyde obtained from the C₉₆ chloro acetal 23 (n = 7) and the C₄₈ phosphonium bromide 19; $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 11 × (CH=CH)], 4.84 [t, CHO₂(CH₂)₂], 3.9 [m, O(CH₂)₂O] and 3.53 (t, CH₂Cl), in the ratio 22.8:1.0:4.0:2.0. It was converted into the DNP derivative for use as a standard for purity checks by HPLC.

192-Bromodononacontahecta-12,24,36,48,60,72,84,96,108,-120,132,144,156,168,180-pentadecaenal ethylene acetal³ (the C_{192} bromo acetal) 22 (n = 15). The ylide prepared from C_{96} acetal triphenylphosphonium bromide 27 (n = 7) and LDA was treated with the C_{96} bromo aldehyde 25 (n = 7), workedup as before and the product separated by chromatography on silica using light petroleum–ether (96:4, v/v) as eluent to give the pure product (80%), identified by ¹H NMR spectroscopy. Analysis of the DNP derivative showed the absence of both C_{96} homologues and any material at longer retention times.

The C₁₉₂ bromo acetal **22** (n = 15) containing secondary alkyl bromide was obtained in similar yield from secondary alkyl bromide-containing moieties, the ¹H NMR spectrum showing this functionality at δ 4.02 (tt, CHBr).

191-(Dioxolan-2-yl)hennonacontahecta-12,24,36,48,60,72,-84,96,108,120,132,144,156,168,180-pentadecaenyl(triphenyl)phosphonium bromide³ (the C₁₉₂ acetal phosphonium bromide) 27 (n = 15). This compound was prepared as for the C₉₆ acetal phosphonium bromide 27 (n = 7); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 15 × (CH=CH)], 3.9 [m, O(CH₂)₂O] and 3.82 (m, CH₂P⁺-Ph₃). The integral ratio for the alkene protons and the combined aliphatic protons was 29.6:6.

240-Bromotetracontadicta-12,24,36,48,60,72,84,96,108,120,-132,144,156,168,180,192,204,216,228-nonadecaenal ethylene acetal (the C₂₄₀ bromo acetal) 22 (n = 19). Reaction of the C₁₉₂ bromo aldehyde 25 (n = 15) (containing secondary bromide) and the ylide 21 from C₄₈ phosphonium bromide 19 and LDA gave the C₂₄₀ bromo acetal 22 (n = 19) which was purified by chromatography on silica (1 kg) with a loading of *ca*. 4 g using light petroleum–ether (95:5, v/v) as eluent (81% yield), the ¹H NMR spectrum of which was exactly as expected, but with an additional bromomethylene signal; $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 19 × (CH=CH)], 4.84 [t, CHO₂(CH₂)₂], 4.02 (tt, CHBr), 3.9 [m, O(CH₂)₂O] and 3.4 (t, CH₂Br), in the ratio 38.5:1.04:4.21:2 after making the allowance for the loss of one -CH=CH– for each–CHBr–.

288-Bromooctaoctacontadicta-12,24,36,48,60,72,84,96,108,-120,132,144,156,168,180,192,204,216,228,240,252,264,276-tricosaenal ethylene acetal³ (the C₂₈₈ bromo acetal) 22 (n = 23). Treatment of the C₁₉₂ bromo aldehyde 25 (n = 15) with the ylide from C₉₆ phosphonium bromide 27 (n = 7) and LDA (both moieties containing secondary bromine) gave the C₂₈₈ bromo acetal 22 (n = 23) which was purified by chromatography on silica (1 kg) with a loading of *ca*. 4 g using light petroleum–ether (97.5:2.5, v/v) as eluent (57% yield), the ¹H NMR spectrum of which was exactly as expected, but with an additional bromomethylene signal; $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 23 × (CH=CH)], 4.84 [t, CHO₂(CH₂)₂], 4.02 (tt, CHBr), 3.9 [m, O(CH₂)₂O] and 3.4 (t, CH₂Br), in the ratio 45.9:0.91:3.86:2.0, after making the allowance for the loss of one –CH=CH– for each –CHBr–.

384-Bromotetraoctacontatricta-12,24,36,48,60,72,84,96,108,-120,132,144,156,168,180,192,204,216,228,240,252,264,276,288,-300,312,324,336,348,360,372-hentriacontaenal ethylene acetal (the C₃₈₄ bromo acetal) 22 (n = 31). Treatment of the C₁₉₂ bromo aldehyde 25 (n = 15) with the ylide from C₁₉₂ phosphonium bromide 27 (n = 15) and LDA gave the C₃₈₄ bromo acetal 22 (n = 31) which was partially purified by chromatography on silica (see later); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 31 × (CH=CH)], 4.84 [t, CHO₂(CH₂)₂], 3.9 [m, O(CH₂)₂O] and 3.4 (t, CH₂Br), in the ratio 63.0:0.95:3.89:2.0.

Syntheses of long chain primary bromides and their phosphonium salts for 'capping' reactions

(i) Octadec-12-enyl bromide 37. The ylide from hexyl(triphenyl)phosphonium bromide and LDA was treated with 12bromododecanal 3 and then filtered through a short column of silica using light petroleum. The solvent was evaporated to give a viscous liquid 37 (Found: C, 65.6; H, 11.0. $C_{18}H_{35}Br$ requires C, 65.24; H, 10.64%); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 1 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 2.0: 2.10.

(ii) Octadec-12-enyl(triphenyl)phosphonium bromide 38. The bromide 37 was treated with triphenylphosphine (1.2 equiv.) in acetonitrile at reflux temperature for 5 d. The solvent was removed *in vacuo* at 40 °C to give a highly viscous oil from which the excess triphenylphosphine was removed by stirring with light petroleum and decanting the supernatant liquid. Removal of traces of solvent gave a viscous liquid 38 which was used without further treatment in subsequent Wittig reactions; $\delta_{\rm H}(\rm CDCl_3)$ 5.36 [m, 1 × (CH=CH)] and 3.65 (t, CH₂P⁺Ph₃), in the ratio 2.0:2.06.

(iii) Pentaconta-12,24,36,48-tetraenyl bromide 39. The ylide from ethyl(triphenyl)phosphonium bromide and LDA was treated with the C₄₈ bromo aldehyde 25 (n = 3) and the crude product was isolated as in (i), above. Chromatography on silica using light petroleum gave the pure C₅₀ tetraenyl bromide 39 (82%) as a glass (Found: C, 77.45; H, 12.2. C₅₀H₉₃Br requires C, 77.57; H, 12.10%); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 4 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 7.96:2.0.

(iv) Pentaconta-12,24,36,48-tetraenyltriphenylphosphonium bromide 40. The C₅₀ tetraenyl bromide 39 was treated as in (ii), and the crude product was chromatographed on silica using dichloromethane-methanol (98:2, v/v) to give the pure salt 40 as a glass (Found: C, 78.45; H, 10.9. C₆₈H₁₀₈BrP requires C, 78.80; H, 10.50%); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 4 × (CH=CH)] and 3.78 (m, CH₂P⁺Ph₃), in the ratio 8.18:2.0.

(v) Hexahexaconta-12,24,36,48,60-pentaenyl bromide 41. The ylide 43 from octadec-12-enyl(triphenyl)phosphonium bromide 38 and LDA was treated with the C₄₈ bromo aldehyde 17, and the crude product, isolated as in (i), was chromatographed on silica using light petroleum to give the pure C₆₆ pentaenyl bromide 41 (82%), mp 27–28 °C (Found: C, 80.0; H, 12.9. C₆₆H₁₂₃Br requires C, 79.54; H, 12.44%); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 5 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 10.4:2.0.

(vi) Hexahexaconta-12,24,36,48,60-pentaenyl(triphenyl)phosphonium bromide 42. The C₆₆ pentaenyl bromide 41 was treated as in (ii), and the crude product was chromatographed on silica using dichloromethane-methanol (95:5, v/v) to give the pure salt 42 as a glass (Found: C, 80.2; H, 11.0. $C_{84}H_{138}BrP$ requires C, 79.19; H, 11.12%); $\delta_{H}(CDCl_3)$ 5.36 [m, 5 × (CH=CH)] and 3.78 (m, -CH₂P⁺Ph₃), in the ratio 5.6:1.

Capping reactions

(i) $C_{96} + C_2 \longrightarrow C_{98}$. A mixture of ethyl(triphenyl)phosphonium bromide (4.41 g, 11.9 mmol) and anhydrous THF (43.7 ml) was cooled to -20 °C under argon and treated with LDA (1.5 mol 1⁻¹ in THF, 6.3 ml, 9.4 mmol). A proportion of this solution (12.5 ml, 2.38 mmol) was added under argon to a suspension of the C_{96} bromo aldehyde **25** (n = 7) (3.2 g, 1.16 mmol) in anhydrous THF (5 ml) at -20 °C followed by further ylide (3.5 ml, 0.66 mmol). The mixture was allowed to warm to room temperature overnight and washed through a short silica column with light petroleum and the solvent evaporated. Chromatography on silica using light petroleum gave *octa*- nonaconta-12,24,36,48,60,72,84,96-octaenyl bromide (the C_{98} bromo polyene) **46** (\equiv **30**, n = 7, $R = CH_3$), mp 37–38 °C (Found: C, 81.7; H, 12.8. $C_{98}H_{181}Br$ requires C, 81.77; H, 12.67%); $\delta_{\rm H}(\rm{CDCl}_3)$ 5.35 [m, 8 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 16.6:2.

(ii) $C_{120} + C_2 \longrightarrow C_{122}$. The C_{120} bromo aldehyde 25 (n = 9) was reacted with the ylide from ethyl (triphenyl)phosphonium bromide as in (i) to give *docosahecta*-12,24,36,48,60,72,-84,96,108,120-*decaenyl bromide* 47 (\equiv 30, n = 9, R = CH₃) (64%), mp 37–38 °C (from light petroleum) (Found: C, 82.6; H, 12.6. $C_{122}H_{225}Br$ requires C, 82.69; H, 12.79%); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 10 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 20.6:2.0.

(iii) $C_{96} + C_{66} \longrightarrow C_{162}$. The C_{96} bromo aldehyde 25 (n = 7) was added to the ylide 45 from the C_{66} triphenylphosphonium bromide 42, as in (i), to give *dohexacontahecta*-12,24,36,-48,60,72,84,96,108,120,132,144,156-*triadecaenyl bromide* 48 $\{\equiv 30, n = 7, R = C_5H_{11}[CH=CH(CH_2)_{10}]_5\}$ (65%), mp 30-31 °C (Found: C, 83.5; H, 13.0. $C_{162}H_{289}$ Br requires C, 83.62; H, 12.95%); $\delta_{H}(CDCl_3)$ 5.35 [m, 13 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 27.2:2.0.

(iv) $C_{192} + C_2 \longrightarrow C_{194}$. The C_{192} bromoaldehyde 25 (n = 15) was treated with the ylide from ethyl(triphenyl)phosphonium bromide as in (i), to give *tetranonacontahecta*-12,24,36,48,60,-72,84,96,108,120,132,144,156,168,180,192-*hexadecaenyl* bromide 49 (\equiv 30, n = 15, $R = CH_3$) (64%), mp 37–38 °C (from light petroleum) (Found: C, 84.1; H, 13.0. $C_{194}H_{357}$ Br requires C, 84.12; H, 12.99%); δ_{H} (CDCl₃) 5.35 [m, 16 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 33.6:2.0.

(v) $C_{192} + C_6 \longrightarrow C_{198}$. The C_{192} bromo aldehyde 25 (n = 15), containing the CHBr feature, was treated with the ylide from hexyl(triphenyl)phosphonium bromide as in (i), to give *octanonacontahecta*-12,24,36,48,60,72,84,96,108,120,132,144,-156,168,180,192-*hexadecaenyl bromide*³ 30 (n = 15, $R = C_5H_{11}$) (75%). The ¹H NMR spectrum was identical with that of an authentic sample, but with an additional bromomethylene signal; $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 16 × (CH=CH)], 4.02 (tt, CHBr) and 3.4 (t, CH₂Br). The ratio of alkene to bromomethyl protons was 16.5:1, after making the allowance for the loss of one –CH=CH– for each–CHBr–.

(vi) $C_{192} + C_{18} \longrightarrow C_{210}$. The C_{192} bromo aldehyde 25 (n = 15) was added to the ylide 43 from the C_{18} triphenylphosphonium bromide 38, as in (i), to give *decadicta*-12,24,-36,48,60,72,84,96,108,120,132,144,156,168,180,192,204-*hepta-decaenyl bromide* 50 { \equiv 30, n = 15, $R = C_5H_{11}$ [CH=CH-(CH₂)₁₀]} (60%), mp 31–32 °C (Found: C, 84.05; H, 13.0. $C_{210}H_{384}$ Br requires C, 84.29; H, 13.03%); δ_{H} (CDCl₃) 5.35 [m, 17 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 35.0:2.0.

(vii) $C_{192} + C_{50} \longrightarrow C_{242}$. The C_{192} aldehyde 26 (n = 15) was added to the ylide 44 from the C_{50} phosphonium bromide 40, as in (i), to give *dotetracontadicta*-2,14,26,38,50,62,74,86,98,110,-122,134,146,158,170,182,194,206,218,230-*cosaene* 51 { \equiv 31, n = 15, R = CH₃[CH=CH(CH₂)₁₀]₄} (60%), mp 32–33 °C (Found: C, 86.9; H, 13.5. $C_{242}H_{446}$ requires C, 86.71; H, 13.24%); δ_{H} (CDCl₃) 5.35 [m, 17 × CH₂CH=CHCH₂] and 2.0 [m, CH₂CH=CHCH₂], in the ratio 2.0:3.8.

(viii) $C_{240} + C_6 \longrightarrow C_{246}$. The C_{240} bromo aldehyde 25 (n = 19), containing the -CHBr- feature, was treated with the ylide from hexyl(triphenyl)phosphonium bromide as in (i) to give *hexatetracontadicta*-12,24,36,48,60,72,84,96,108,120,132,144,-156,168,180,192,204,216,228,240-*cosaenyl bromide*³ **30** (n = 19, $R = C_5H_{11}$). The ¹H NMR spectrum was identical with that of an authentic sample, but with an additional bromomethylene signal; $\delta_H(CDCl_3)$ 5.36 [m, 20 × (CH=CH)], 4.02 (tt, CHBr) and 3.4 (t, CH₂Br). The alkene and bromomethyl integrals were in the ratio 20.8:1, after making the allowance for the loss of one - CH=CH- for each -CHBr-.

(ix) $C_{192} + C_{66} \longrightarrow C_{258}$. The C_{192} bromo aldehyde 25 (n = 15) was added to the ylide 45 from C_{66} triphenylphosphonium bromide 42, as in (i) to give *octapentacontadicta*-12,24,-

36,48,60,72,84,96,108,120,132,144,156,168,180,192,204,216,-228,240,252-*hencosaenyl bromide* **52** { \equiv **30**, *n* = 15, R = C₅H₁₁[CH=CH(CH₂)₁₀]₅} (36%), mp 36–37 °C (Found: C, 84.3; H, 13.2. C₂₅₈H₄₇₅Br requires C, 84.72; H, 13.08%); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 21 × (CH=CH)] and 3.4 (t, CH₂Br), in the ratio 44.3:2.0.

(x) $C_{288} + C_6 \longrightarrow C_{294}$. The C_{288} bromo aldehyde 25 (n = 23), containing the -CHBr- feature, was treated with the ylide from hexyl(triphenyl)phosphonium bromide as in (i) to give *tetranonacontadicta*-12,24,36,48,60,72,84,96,108,120,132,144,-156,168,180,192,204,216,228,240,252,264,276,288-*tetracosa*-

*enyl bromide*³ **30** (n = 23, $\mathbf{R} = C_5 H_{11}$). The ¹H NMR spectrum was identical with that of an authentic sample, but with an additional bromomethylene signal; $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 24 × (CH=CH)], 4.02 (tt, CHBr) and 3.4 (t, CH₂Br). The alkene and bromomethyl integrals were in the ratio 24.7:1, after making the allowance for the loss of one –CH=CH– for each–CHBr–.

(xi) $C_{384} + C_6 \longrightarrow C_{390}$. The C_{384} aldehyde ³ **26** (n = 31) (see below) was treated with the ylide from hexyl(triphenyl)phosphonium bromide as in (i) to give *nonacontatricta*-6,18,-30,42,54,66,78,90,102,114,126,138,150,162,174,186,198,210,-222,234,246,258,270,282,294,306,318,330,342,354,366,378-*do*-

triacontaene³ **53** (\equiv **31**, n = 31, $R = C_5H_{11}$)³, mp 37–38 °C (Found: C, 86.5; H, 13.5. $C_{390}H_{718}$ requires C, 86.62; H, 13.38%); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 32 × (CH=CH)], 1.95 (m, 64 × CH₂CH=), 1.3 [m, 260 × (CH₂), *i.e.* 520 protons] and 0.88 (2 × CH₃), in the ratio 64.0:128.5:520:5.9.

Replacement of bromine by hydrogen

(i) C_{98} bromo polyene $\longrightarrow C_{98}$ polyene. The C_{98} bromo polyene 46 (2.41 g, 1.67 mmol) and 'Superhydride' (lithium triethylborohydride; 1.0 mol l⁻¹ solution in THF; 10 ml, 0.1 mol), were stirred together at room temperature under argon for 4 h. The product was dissolved in light petroleum and washed with dilute sulfuric acid (0.5 mol 1⁻¹; 200 ml) and water (200 ml). In each case the aqueous layers were extracted with light petroleum, and the organic layers combined and dried over sodium sulfate. The solvent was removed under vacuum, leaving a colourless oil, which was chromatographed on silica using light petroleum, to give octanonaconta-2,14,26,38,50,-62,74,86-octaene (the C_{98} polyene) **59** (\equiv **31**, n = 7, $R = CH_3$), (2.14 g, 94%), mp 37.5-38 °C (Found: C, 86.6; H, 13.6. $C_{98}H_{182}$ requires C, 86.52; H, 13.48%; $\delta_{H}(CDCl_{3})$ 5.35 $[m, 8 \times (CH=CH)], 1.95 (m, 15 \times CH_2CH=), 1.3 [m, (CH_2)_9$ and 7 × $(CH_2)_8$, *i.e.* 130 protons], with intensities in the ratio 16.0:29.1:126; and 1.6 (dd, =CHCH₃) and 0.87 (t, CH₂CH₃).

(ii) C_{122} bromo polyene $\longrightarrow C_{122}$ polyene. The C_{122} bromo polyene 47 was treated as in (i) to give *docosahecta*-2,14,26,38,50,62,74,86,98,110-*decaene* 60 (\equiv 31, n = 9, $R = CH_3$) (98%), mp 31–32 °C (Found: C, 86.5; H, 13.6. $C_{122}H_{226}$ requires C, 86.54; H, 13.45%); $\delta_{\rm H}(\rm CDCl_3)$ 5.35 [m, 10 × (CH=CH)], 1.95 (m, 19 × CH₂CH=), 1.3 [m, CH₂ and 10 × (CH₂)₈, *i.e.* 162 protons], with intensities in the ratio 20.0:37.9:162.0; and 1.60 and 1.64 (both d, =CHCH₃) and 0.87 [t, CH₂CH₃].

(iii) C_{162} bromo polyene $\longrightarrow C_{162}$ polyene. The C_{162} bromo polyene **48** was treated as in (i) to give *dohexacontahecta*-6,18,30,42,54,66,78,90,102,114,126,138,150-*triadecaene* **61** { \equiv **31**, n = 7, $R = C_5H_{11}$ [CH=CH(CH₂)₁₀]₅} (84%), mp 36-37 °C (Found: C, 86.5; H, 13.5. $C_{162}H_{300}$ requires C, 86.55; H, 13.45%); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 13 × (CH=CH)], 2.0 (m, 26 × CH₂CH=) and 1.3 [m, 108 × (CH₂), *i.e.* 216 protons], in the ratio 26.0:53.9:215.

(iv) C_{192} bromo acetal $\longrightarrow C_{192}$ acetal. The C_{192} bromo acetal **22** (n = 15) was treated as in (i) to give dononacontahecta-12,24,36,48,60,72,84,96,108,120,132,144,156,168,180-pentadecaenal acetal³ **24** (n = 15); $\delta_{H}(CDCl_3)$ 5.36 [m, 15 × (CH= CH)] and 4.84 [t, CHO₂(CH₂)₂], in the ratio 31.2:4.

(v) C_{194} bromo polyene $\longrightarrow C_{194}$ polyene. The C_{194} bromo

polyene **49** was treated as in (i) to give *tetranonacontahecta*-2,14,26,38,50,62,74,86,98,110,122,134,146,158,170,182-*hexadecaene* **62** (\equiv **31**, n = 15, $R = CH_3$), mp 35–36 °C (Found: C, 86.35; H, 13.5. C₁₉₄H₃₅₈ requires C, 86.59; H, 13.41%); $\delta_{\rm H}(\rm CDCl_3)$ 5.35 [m, 16 × (CH=CH)], 2.0 [m, 31 × CH₂CH=] and 1.3 [m, 129 × (CH₂), *i.e.* 258 protons], in the ratio 32.0:61.4:272.

(vi) C_{198} bromo polyene $\longrightarrow C_{198}$ polyene. The C_{198} bromo polyene 30 (n = 15, R = C₅H₁₁) containing the CHBr feature was treated as in (i) to give octanonacontahecta-6,18,30,42,54,66,78,90,102,114,126,138,150,162,174,186-hexadecaene ³ 31 (n = 15, R = C₅H₁₁). The ¹H NMR spectrum was identical with that of an authentic sample, but with an additional bromomethylene signal; $\delta_{\rm H}({\rm CDCl}_3)$ 5.35 [m, (CH=CH)], 4.02 (tt, CHBr), 2.0 (m, 31 × CH₂CH=) and 1.3 [m, (CH₂)]. The residual CHBr was removed by heating the polyene (1.53 g) with 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) (2.32 g) under argon at 150 °C for 4 h. The mixture was cooled, shaken with sulfuric acid (4 mol l^{-1}) and ether-light petroleum, and the organic solvents removed in vacuo at 20 °C to avoid the formation of an emulsion. The solid which separated was filtered and washed with water, dissolved in ether-light petroleum, dried (Na2SO4) and the solvent evaporated. The product was chromatographed on silica using light petroleum-ether (98.5:1.5, v/v) as eluent, to give the halogen-free polyene **31** (n = 15, $R = C_5 H_{11}$).

(vii) C_{210} bromo polyene $\longrightarrow C_{210}$ polyene. The C_{210} bromo polyene 50 was treated as in (i) to give *decadicta*-6,18,30,42,54,66,78,90,102,114,126,138,150,162,174,186,198*heptadecaene* $63 [<math>\equiv$ 31, n = 15, $R = C_5H_{11}CH=CH(CH_2)_{10}$] (87%), mp 31–32 °C (Found: C, 86.5; H, 13.0. $C_{210}H_{388}$ requires C, 86.57; H, 13.42%); $\delta_{H}(CDCl_3)$ 5.35 [m, $17 \times (CH=CH)$], 1.95 (m, 34 × $CH_2CH=$) and 1.3 [m, $140 \times (CH_2)$, *i.e.* 280 protons], in the ratio 34.0:68.8:299.

(viii) C_{246} bromo acetal $\longrightarrow C_{246}$ acetal. The C_{246} bromo polyene 30 (n = 19, $R = C_5H_{11}$) containing the -CHBr- feature was reacted as in (i) and dehydrobrominated as in (vi) to give isomers of hexatetracontadicta-6,18,30,42,54,66,78,90,102,-114,126,138,150,162,174,186,198,210,222,234-cosaene³ 31 (n = 19, $R = C_5H_{11}$); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 20 × (CH=CH)], 1.95 (m, 40 × CH₂CH=), 1.3 [m, 164 × (CH₂), *i.e.* 328 protons] and 0.92 (2 × CH₃), in the ratio 40.0:81:337:6.1.

(ix) \overline{C}_{258} bromo polyene $\longrightarrow C_{258}$ polyene. The C_{258} bromo polyene 52 {= 30, n = 15, R = C_5H_{11} [CH=CH(CH₂)₁₀]₅} was reacted as in (i) to give *octapentadicta*-6,18,30,42,54, 66,78,90,102,114,126,138,150,162,174,186,198,210,222,234,-246-*hencosaene* 64 {= 31, n = 15, R = C_5H_{11} [CH=CH-(CH₂)₁₀]₅} (43%), mp 38–39 °C (Found: C, 86.4; H, 13.5. $C_{258}H_{476}$ requires C, 86.59; H, 13.40%); δ_{H} (CDCl₃) 5.35 [m, 21 × (CH=CH)], 1.95 (m, 42 × CH₂CH=) and 1.3 [m, 172 × (CH₂), *i.e.* 344 protons], in the ratio 42.0:82.1:349.

(x) C_{294} bromo polyene $\longrightarrow C_{294}$ polyene. The C_{294} bromo polyene 30 (n = 23, $R = C_5H_{11}$) containing the -CHBrfeature was reacted as in (i) and dehydrobrominated as in (vi) to give isomers of tetranonacontadicta-6,18,30,-42,54,66,78,90,102,114,126,138,150,162,174,186,198,210,222,-234,246,258,270,282-tetracosaene³ 31 (n = 23, $R = C_5H_{11}$); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 24 × (CH=CH)], 1.95 (m, 48 × CH₂-CH=), 1.3 [m, 196 × (CH₂), *i.e.* 392 protons] and 0.92 (2 × CH₃), in the ratio 48.0:92.6:394.

(xi) C_{384} bromo acetal $\longrightarrow C_{384}$ acetal. The C_{384} bromo acetal 22 (n = 31) was reacted as in (i) and purified by chromatography on silica using light petroleum–ether (96:4, v/v) to give the C_{384} acetal ³ 24 (n = 31); $\delta_{\rm H}$ (CDCl₃) 5.36 [m, 31 × (CH=CH)], 4.82 [t, CHO₂(CH₂)₂] and 3.9 [m, O(CH₂)₂O], in the ratio 63.2:0.9:4.

Hydrogenation of straight chain polyenes

(i) $C_{98}H_{198}$ formation. The C_{98} polyene 59 (2.01 g), 5% Pt on activated carbon (1 g) and ethyl palmitate (10 ml) were stirred

together and heated at 130 °C in an atmosphere of hydrogen for 14 h. Light petroleum (bp 100-120 °C) (10 ml) was added to the mixture at 110 °C which was then allowed to cool slowly to room temperature. Solid material was filtered and washed with light petroleum, placed in a cellulose extraction thimble and extracted for 16 h with boiling light petroleum (bp 100-120 °C) (150 ml) in an apparatus which allowed the hot vapours to heat the thimble through which was passing condensed solvent vapours. After cooling the extract slowly to room temperature, the white solid was filtered, washed with light petroleum, and finally heated at 110 °C and 0.01 mmHg for 48 h to remove traces of dibutyl and dioctyl phthalate to give octanonacontane (1.84 g, 90%), mp 113-114 °C (Found: C, 85.4; H, 14.5. C₉₈H₁₉₈ requires C, 85.50; H, 14.49%); the ¹H NMR in 1,1,2,2tetrachloro[²H₂]ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(ii) $C_{122}H_{246}$ formation. The C_{122} polyene 60 was hydrogenated as in (i) to give *docosahectane* (1.62 g, 97%), mp 116–117 °C (Found: C, 85.6; H, 14.7. $C_{122}H_{246}$ requires C, 85.53; H, 14.47%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(iii) $C_{162}H_{326}$ formation. The C_{162} polyene 61 was hydrogenated as in (i) to give *dohexacontahectane* (2.74 g, 97%), mp 121–121.5° (Found: C, 85.4; H, 14.5. $C_{162}H_{326}$ requires C, 85.55; H, 14.44%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]-ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(iv) $C_{194}H_{390}$ formation. The C_{194} polyene 62 was hydrogenated as in (i) to give *tetranonacontahectane* (1.41 g, 90%), mp 122–123 °C (Found: C, 85.6; H, 14.2. $C_{194}H_{390}$ requires C, 85.56; H, 14.43%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(v) $C_{198}H_{398}$ formation. The C_{198} polyene 31 (n = 15, $R = C_5H_{11}$) was hydrogenated as in (i) to give octanonacontahectane³ (1.46 g); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(vi) $C_{210}H_{422}$ formation. The C_{210} polyene 63 was hydrogenated as in (i) to give *decadictane* (1.23 g, 93%), mp 125–126 °C (Found: C, 85.6; H, 14.4. $C_{210}H_{422}$ requires C, 85.58; H, 14.42%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]-ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(vii) $C_{242}H_{486}$ formation. The C_{242} polyene 51 was hydrogenated as in (i) to give *dotetracontadictane* (1.41 g, 91%), mp 120–121 °C (Found: C, 85.7; H, 14.6. $C_{242}H_{486}$ requires C, 85.58; H, 14.42%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]-ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(viii) $C_{246}H_{494}$ formation. The C_{246} polyene 31 (n = 19, $R = C_5H_{11}$) was hydrogenated as in (i) to give hexatetracontadictane (3.38 g);³ the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(ix) $C_{258}H_{518}$ formation. The C_{258} polyene 64 was hydrogenated as in (i) to give *octapentacontadictane* (0.3 g, 61%), mp 125–126 °C (Found: C, 85.65; H, 14.4. $C_{258}H_{518}$ requires C, 85.58; H, 14.41%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(x) $C_{294}H_{590}$ formation. The C_{294} polyene 31 (n = 23, $R = C_5H_{11}$) was hydrogenated as in (i) to give tetranonacontadictane³ (2.57 g); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]-ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

(xi) $C_{390}H_{782}$ formation. The C_{390} polyene 53 was hydrogenated as in (i) to give nonacontatrictane ³ (0.83 g, 96%), mp 125–126 °C (Found: C, 85.8; H, 14.5. $C_{390}H_{782}$ requires C, 85.60; H, 14.40%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]-ethane at 120 °C showed the absence of alkenic absorptions at δ 5.36.

Syntheses of branched chain alkanes

(a) $C_{96}H_{193}CH(CH_3)C_{94}H_{189}$. (i) The branched C_{192} dibromo- α,β -unsaturated aldehyde containing –CHBr– **36** (4.97 g) in anhydrous THF (20 ml) was added to a stirred suspension of lithium aluminium hydride (1.75 g) in anhydrous THF (35 ml) at –40 °C over 5 min. The excess hydride was destroyed with ethyl acetate, the mixture acidified (4 mol l⁻¹ H₂SO₄), extracted with light petroleum–ether, the extracts dried (Na₂SO₄) and solvent evaporated. Chromatography of the crude product on silica using light petroleum–ether (90:10, v/v) gave the branched C_{192} dibromo unsaturated alcohol, the ¹H NMR of which showed 16% conversion of –CH₂Br to –CH₃; δ_{H} (CDCl₃) 5.36 (m, 29 × CH=C), 4.02 (s, CH₂OH) (masking the CHBr) and 3.4 (t, CH₂Br) in the ratio 29.0:2.16:3.9.

(ii) The branched C_{192} dibromo unsaturated alcohol from (i) (3.06 g) in dry light petroleum (6 ml) was treated at 15 °C with PBr₃ (0.168 g) in dry light petroleum (2.5 ml). After 1 h, the mixture was added to water, extracted with ether, the extracts dried (Na₂SO₄) and solvent evaporated. The crude product was purified by chromatography on silica using light petroleum–ether (98:2, v/v) to give the branched C_{192} *dibromo unsaturated bromide*; $\delta_{\rm H}$ (CDCl₃) 5.62 [t, CH=C(CH₂-Br)], 5.26 [m, 14 × (CH=CH)], 4.0 [s, CH=C(CH₂Br), overlapping the CHBr at δ 4.02] and 3.4 (t, CH₂Br); also $\delta_{\rm H}$ (CDCl₃) 5.15 and 4.9 (both s, CHBrC=CH₂) and 4.5 (t, CH₂CHBrC=CH₂), the result of *ca.* 4% S_N2' displacement of OH.

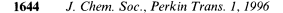
(iii) The branched C_{192} dibromo unsaturated bromide from (ii) was reacted with 'Superhydride' (lithium triethylborohydride) as described before. The ¹H NMR of the C_{192} - C_1 branched product showed the -CHBr- at δ 4.02 and also evidence of S_N2' displacement of Br; $\delta_H(CDCl_3)$ 5.12 [t, CH₂CH=C(CH₃)] and 4.7 [s, CH₂C(=CH₂)CH₂], in the ratio 90:10; and 1.56 and 1.67 [both s, due to geometrical isomers of CH=C(CH₃)]. The bromine from the -CHBr- was removed with DBU as described before.

(iv) The C₁₉₁-CH₃ branched polyene from (iii) (1.72 g) was hydrogenated as described before to give 95-*methylhennonacontahectane* **54** (1.57 g), mp 116 °C (DSC) (Found: C, 86.5; H, 13.7. C₁₉₂H₃₈₆ requires C, 86.53; H, 13.47%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of any alkenic absorptions.

(b) $C_{96}H_{193}CH(CH_2CH_2CH_2CH_3)C_{94}H_{189}$. (i) The branched C_{192} dibromo α,β -unsaturated aldehyde containing –CHBr– **36** (2.66 g), was treated under argon at room temperature with the ylide from propyl(triphenyl)phosphonium bromide and LDA in anhydrous THF (0.035 mol l⁻¹) until the orange–yellow colour persisted. The mixture was washed through a short column of silica with light petroleum–ether (90:10, v/v) and the solvent evaporated. The crude product (2.88 g) was chromatographed on silica to give the 95-(*but*-1-*enyl*) C_{191} *dibromo polyene*; $\delta_{H}(CDCl_3)$ 5.58, 5.72 and 5.93 (associated with CH₃CH₂CH=CHC=CH) and 0.98 and 1.01 (both t, geometrical isomers of CH₃CH₂CH=CH).

(ii) The 95-(but-1-enyl) C_{191} dibromo polyene from (i) was reacted with 'Superhydride' (lithium triethylborohydride) as described before. The ¹H NMR of the 95-(but-1-enyl) C_{191} polyene showed the -CHBr- group at δ 4.02 (CDCl₃); careful removal of all traces of light petroleum by co-distillation with chloroform revealed three clean triplets, at 0.89 (2 × CH₃ at the end of chain), and 0.98 and 1.01 (geometrical isomers of CH₃CH₂CH=CH) in the ratio 2:1. The bromine from the -CHBr- group was removed with DBU as described before.

(iii) The 95-(but-1-enyl) C_{191} polyene from (ii) was hydrogenated as described before to give 95-butylhennonacontahectane 55 (1.84 g), mp 116 °C (DSC) (Found: C, 85.5; H, 14.8. $C_{195}H_{392}$ requires C, 85.56; H, 14.44%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of any alkenic absorptions.



Syntheses of long chain carboxylic acids

(a) CH₃(CH₂)₁₉₀CO₂H. (i) The C₁₉₂ acetal 24 (n = 15) (0.84 g) was deprotected on silica (73 g) impregnated with toluene-*p*-sulfonic acid monohydrate (1.83 g) and water (5.4 ml), with light petroleum–dichloromethane (3:1, v/v) as eluent. The resulting aldehyde 26 (n = 15) was dissolved in a mixture of THF (6 ml) and ethanol (3 ml) and AgNO₃ (1.27 g), and sodium hydroxide (0.35 g) in water (2.5 ml) was added. After stirring for 2 h at room temperature, the mixture was acidified with sulfuric acid (4 mol l⁻¹), extracted with light petroleum–ether, the extracts dried (Na₂SO₄) and the solvent evaporated. The crude product (0.96 g) was purified by chromatography on silica using light petroleum–ether (85:15, v/v) as eluent to give *dononacontahecta*-12,24,36,48,60,72,84,96,108,120,132,144,156,168,180-*penta*-

decaenoic acid, mp 44.5–47 °C (Found: C, 85.8, H, 13.3. $C_{192}H_{354}O_2$ requires C, 85.57; H, 13.24%); $\delta_H(CDCl_3)$ 5.35 [m, 15 × (CH=CH)], 2.36 (t, CH₂CH₂CO₂H), 1.95 (m, 30 × CH₂CH=) and 1.65 (q, CH₂CH₂CO₂H), in the ratio 30.2:2.0:62:1.94.

(ii) The C₁₉₂ carboxylic acid from (i) was hydrogenated as described before, but extracted with octane, to give *dononacontahectanoic acid* **56** (0.60 g), 127.5–130.5 °C (Found: C, 84.6; H, 14.4. C₁₉₂H₃₈₄O₂ requires C, 84.62; H, 14.20%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of any alkenic absorptions.

(b) HO₂C(CH₂)₁₉₂CO₂H. (i) The C₁₉₂ bromo aldehyde 25 (n = 15) (3.29 g) in bromoethane (5 ml) was added to lithium aluminium hydride (0.26 g) in anhydrous ether (10 ml) at 0 to -5 °C. After 30 min, ether (30 ml) was added and the mixture was added to light petroleum–ether over sulfuric acid (4 mol1⁻¹). The organic layer was separated, dried (Na₂SO₄), the solvent evaporated and the crude product chromatographed on silica using light petroleum–ether (90:10, v/v) as eluent to give 192bromodononacontahecta-12,24,36,48,60,72,84,96,108,120,132,-144,156,168,180-pentadecaenol, mp 44–45 °C (Found: C, 84.0; H, 13.0. C₁₉₂H₃₅₅BrO requires C, 83.56; H, 12.96%); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 15 × (CH=CH)], 1.95 (m, 30 × CH₂-CH=), 3.65 (t, CH₂OH) and 3.4 (t, CH₂Br), in the ratio 30:60,7:1.73:1.70.

(ii) A mixture of the C₁₉₂ bromo alcohol from (i) (2.03 g), carbon tetrabromide (0.327 g) suspended in anhydrous dichloromethane (30 ml) at -5 °C was treated with triphenylphosphine (0.318 g) in dry dichloromethane (5 ml) and the temperature allowed to rise to room temperature. After 3 h the mixture was washed through a short silica column using ether, the solvents were evaporated and the crude product (2.22 g) chromatographed on silica using light petroleum-ether (90:10, v/v) to give 1,192-*dibromodononacontahecta*-12,24,-36,48,60,72,84,96,108,120,132,144,156,168,180-*pentadecaene*, mp 37.5–39 °C (Found: C, 82.1; H, 12.7. C₁₉₂H₃₅₄Br₂ requires C, 81.70; H, 12.64%); $\delta_{\rm H}({\rm CDCl}_3)$ 5.35 [m,

15 × (CH=CH)] and 3.4 (t, 2 × CH₂Br), in the ratio 30:3.87. (iii) The C₁₉₂ dibromo compound from (ii) (2.02 g), magnesium powder (1.00 g) and ether (12 ml) freshly distilled from LiAlH₄, were heated together under reflux with anhydrous 1,2-dibromoethane being added (two drops at a

anhydrous 1,2-dibromoethane being added (two drops at a time) to the mixture every 30 min over 2 h. Carbon dioxide, dried by passage through conc. sulfuric acid then through molecular sieves, (4 Å) was bubbled through the mixture for 20 min, which was then acidified (4 mol 1^{-1} H₂SO₄), extracted with ether, the extracts dried and the solvent evaporated. The crude residue was chromatographed on silica using light petroleum–ether (90:10, v/v) as eluent. The slowest moving component was *tetranonacontahecta*-13,25,37,49,61,73,85,97,109,121,133, 145,157,169,181-*pentadecaen*-1,194-*dioic acid*, mp 43–44.5 °C (Found: C, 84.3; H, 12.8. C₁₉₄H₃₅₆O₄ requires C, 84.64; H, 13.03%); $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 15 × (CH=CH)], 2.36 (t, 2 × CH₂CH₂CO₂H), 1.95 (m, 30 × CH₂CH=), 1.65 (q, 2 × CH₂CH₂CO₂H) and 1.3 [128 × (CH₂), *i.e.* 256 protons], in the ratio 30.0:3.92:59:3.75:252.

(iv) The C₁₉₄ dicarboxylic acid from (iii) was hydrogenated as described before, but extracted for 7 days to give *tetranonacontahectane*-1,194-*dioic acid* **58** (0.59 g), mp 131– 133 °C (Koffler block) (Found: C, 83.6; H, 14.0. C₁₉₄H₃₈₆O₄ requires C, 83.72; H, 13.98%); the ¹H NMR in 1,1,2,2tetrachloro[²H₂]ethane at 120 °C showed the absence of any alkenic absorptions; $\delta_{\rm H}$ 2.4 (t, 2 × CH₂CH₂CO₂H), 1.7 (q, 2 × CH₂CH₂CO₂H) and 1.4 (m, 188 × CH₂, *i.e.* 376 protons), in the ratio 4.0:4.17:371.

(c) $HO_2(CH_2)_{48}CO_2H$. (i) The C₄₈ bromo aldehyde 17 was converted, by the method described before, into 48bromooctatetraconta-12,24,36-trienol, mp 45.5–48 °C (Found: C, 75.9; H, 12.1. C₄₈H₉₁BrO requires C, 75.45; H, 12.00%).

(ii) The C₄₈ bromo alcohol from (i) was reacted with PBr₃ as described before, to give the C₄₈ dibromo compound, which in turn reacted with magnesium–carbon dioxide to give *pentaconta*-13,25,37-*trien*-1,50-*dioic acid*; $\delta_{\rm H}$ (CDCl₃) 5.35 [m, 3 × (CH=CH)], 2.36 (t, 2 × CH₂CH₂CO₂H), 1.95 (m, 6 × CH₂CH=), 1.65 (q, 2 × CH₂CH₂CO₂) and 1.3 [32 × (CH₂), *i.e.* 64 protons], in the ratio 6.0:3.95:12.0:3.97:68.7.

(iii) The C₅₀ dicarboxylic acid was hydrogenated as described before, to give *pentacontane*-1,50-*dioic acid* **57** (0.46 g), mp 127.5–130 °C (Koffler block, but two peak maxima on DSC) (Found: C, 79.0; H, 13.1. C₅₀H₉₈O₄ requires C, 78.68; H, 12.94%); the ¹H NMR in 1,1,2,2-tetrachloro[²H₂]ethane at 120 °C showed the absence of any alkenic absorptions; $\delta_{\rm H}$ 2.4

(t, $2 \times CH_2CH_2CO_2H$), 1.7 (q, $2 \times CH_2CH_2CO_2H$) and 1.4 (m, $44 \times CH_2$, *i.e.* 88 protons), in the ratio 4.0:3.87:91.

Acknowledgements

The Authors thank Professors W. J. Feast and D. Saunders for their great encouragement throughout this work, Dr A. Kenwright, Julia Say (Durham branch of the Interdisciplinary Research Centre in Polymer Science and Technology) and Ian McKeag for NMR work and the EPSRC for funding (grants GR/G 59615 and GR/G 66560).

References

- 1 G. Ungar, J. Stejny, A. Keller, I. Bidd and M. C. Whiting, *Science*, 1985, **229**, 386.
- 2 K. S. Lee and G. Wegner, *Makromol. Chem.*, *Rapid Commun.*, 1985, 6, 203.
- 3 I. Bidd, D. W. Holdup and M. C. Whiting, J. Chem. Soc., Perkin Trans. 1, 1987, 2455.
- 4 G. Ungar, personal communication.
- 5 I. Bidd, D. J. Kelly, P. M. Ottley, O. I. Paynter, D. J. Simmons and M. C. Whiting, J. Chem. Soc., Perkin Trans. 1, 1983, 1369.

Paper 5/08126B Received 13th December 1995 Accepted 24th January 1996

[©] Copyright 1996 by the Royal Society of Chemistry