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Review

Synthesis of highly ¹³C enriched carotenoids: access to carotenoids enriched with ¹³C at any position and combination of positions*

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Carotenoids and their metabolites are essential factors for the maintenance of important life processes such as photosynthesis. Animals cannot synthesize carotenoids de novo, they must obtain them via their food. In order to make intensive animal husbandry possible and maintain human and animal health synthetic nature identical carotenoids are presently commercially available at the multi-tonnes scale per year. Synthetically accessible ¹³C enriched carotenoids are essential to apply isotope sensitive techniques to obtain information at the atomic level without perturbation about the role of carotenoids in photosynthesis, nutrition, vision, animal development, etc. Simple highly ^{13}C enriched C $_1$, C $_2$ and C $_3$ building blocks are commercially available via 99% 13CO. The synthetic routes for the preparation of the ¹³C enriched building blocks starting from the commercially available systems are discussed first. Then, how these building blocks are used for the synthesis of the various ¹³C enriched carotenoids and apocarotenoids are reviewed next. The synthetic Schemes that resulted in ¹³C enriched β-carotene, spheroidene, β-cryptoxanthin, canthaxanthin, astaxanthin, (3R,3'R)-zeaxanthin and (3R,3'R,6'R)lutein are described. The Schemes that are reviewed can also be used to synthetically access any carotenoid and apocarotenoid in any ¹³C isotopically enriched form up to the unitarily enriched form.

Keywords: $\boldsymbol{\beta}\text{-carotene},$ spheroidene, astaxanthin, canthaxanthin, zeaxanthin

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INTRODUCTION

Carotenoids are a class of coloured unsaturated tetraterpenes. Carotenoid biosynthesis takes place in the chloroplast of plant cells, cyanobacteria, algae and fungi via the methylerythritol phosphate pathway distinct from the mevalonic acid pathway for all other isoprenoids (Goodwin, 1980; Thibodeux & Liu, 2009).

The carotenoids occur in membrane carotenoid complexes in photosynthetic organism both in antennae pigments that absorb light and in the photosynthetic reaction centre that convert the electronic energy into energy rich molecules. In the latter complexes, they protect these systems against light damage (Britton, 2009; Telfer *et al.*, 2009).

The products of organic photosynthesis serve as food and energy rich materials in our society. The fossil fuels that we use are the products of photosynthesis in the past. Without carotenoids oxygenic photosynthesis would be impossible and life on earth as we know it would not be possible.

Animals cannot synthesize carotenoids *de novo*. They are dependent on the plant materials in their food. A very important role of some carotenoids is their enzymatic generation of retinoids in the body. In Scheme 1 it is indicated that β -carotene 1 is enzymatically converted into all-*trans* retinal 2 which is further converted into retinoid 3 and retinoic acid 4 (Britton, 2009).



Scheme 1. The enzymatic conversion of β -carotene 1 in the human (animal) body into retinal 2, which can be converted into retinol (vitamin A) 3 and retinoic acid 4.

Besides this important role as source of the retinoids, carotenoids fulfill many other important roles to maintain health in humans and animals (Carotenoids, 2009). The present day intense animal husbandry and maintenance of human health is impossible without industrially produced nature-identical carotenoids. A number of important synthetic, nature-identical carotenoids are produced by BASF and DSM at a 3000 tonnes scale a year (Paust, 1996).

In order to follow the conversions of nutrients in the body isotopically labeled systems are essential. This type of study was pioneered by Schoenheimer with stable isotope labeled macronutrients (Schoenheimer, 1942). At that time this approach for macronutrients like carotenoids would not have worked due to the low sensitivity of the isotope sensitive techniques. In the mean time the analytical techniques have greatly improved in sensitivity such that site-directed stable isotope retinoids with high isotope incorporation at at least 3 positions

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{(10,19,19,19-2H)-retinyl acetate} have become a favorite (Olsen, 1997).

The separation properties may even change upon introduction of a number of deuterium isotopes; octadeuteroβ-carotene can be separated from natural abundance β -carotene with HPLC techniques (Dueker *et al.*, 1994). Deuterium is situated at the periphery of the molecule, the deuterium-protium mass ratio is 2/1 and the C-D bond is shorter than the C-H bond length. In the case of multiple ¹³C enrichment the isotope is situated in the carbon skeleton of the molecule and the mass ratio is 13/12. For these reasons ¹³C labels systems were used recently as reference in nutritional studies (van Lieshout et al., 2001; Wang et al., 2008). Besides the difference in mass the 13C isotope has I=1/2 which allows the ¹³C NMR studies. Isotope sensitive non-invasive physical techniques such as NMR and resonance Raman studies allow obtaining information at atomic resolution without any perturbation of the carotene-proteins involved in light harvestation and energy conversion during photosynthesis. In this paper we discuss the site-direct ¹³C incorporation in carotenoids at any position and combination of positions for nutritional and spectroscopic studies.

ACCESS TO HIGHLY ¹³C₂, ¹³C₃ ENRICHED BUILDING BLOCKS

In natural abundance carbon contains 1.1% ¹³C and 98.9% ¹²C. Commercially 99% ¹³CO is obtained by cryogenic distillation of carbon monoxide (Lockhart, 1979). The enrichment factor is 90. As indicated from ¹³CO **5**, ¹³CH₃OH **6**, K¹³CN **8** and ¹³CH₃I **9** are easily prepared (Lockhart, 1979). Simple reactions lead to [1-¹³C]-, [2-¹³C]-and [1,2-¹³C₂]-acetonitrile **10** and [1-¹³C]-, [2-¹³C]- and [1,2-¹³C₂]-acetic acid **7** depending if the synthesis of the two carbon building blocks is prepared with only one of the both highly ¹³C enriched building blocks. Ethyl bromoacetate **16** is prepared *via* a bromination and subsequent esterification.

The ${}^{13}C_2$ building blocks have been used to make Wittig reagents (Creemers & Lugtenburg, 2002). The anion of acetonitrile **10** reacts with methyl iodide **9** in high



Scheme 2. The ${}^{13}C_2$ building blocks that are easily available from 99% ${}^{13}CO$ 5 via known procedures.

They are accessible in any site directed highly enriched form up to the ${}^{13}C_3$ enriched form.



Scheme 3. The preparation of the highly ¹³C enriched building blocks for the synthesis of β -carotene. They are available in any ¹³C enrichment in any possible position.

yield to give propionitrile 12. Diethyl chlorophosphate reacts with the anion of acetonitrile 10 and the anion of propionitrile 12 to give diethyl phosphonoacetonitrile 11 and diethyl phosphonopropionitrile 13, respectively. An Arbuzov reaction of ethyl bromoacetate 16 with triethylphosphite gives ethyl diethyl phosphonoacetate 17. The reaction of ethyl bromoacetate 16 with triphenylphosphine gives triphenyl phosphonium acetate 18. Anion of 18 can easily be converted into the corresponding propionate system 19 by treatment with methyl iodide 9.

Depending on the reagents each ¹³C₃ building block is available in high ¹³C enrichment at any carbon position and any combination of carbon positions. The Wittig reaction of diethyl phosphonopropionitrile **13** with benzaldehyde and subsequent DIBAL-H reduction gives 3-phenyl-2-methyl-3-propenal **14**. To obtain the acetal of prop-2-one-1-al **15**, product **14** is treated first with 1,2-dihydroxymethylene benzene to protect the aldehyde, then followed by oxidative cleavage with potassium permanganate (Jansen, 1996).

In Scheme 3 it is indicated that reagents from Scheme 2 are used to prepare ethyl-3-oxo-butyrate **21** via a Blaise reaction of acetonitrile and the zinc derivative of iodoacetate **20**. Monochlorination of **21** and subsequent acid hydrolysis gives 1-chloroacetone **22** (Creemers & Lugtenburg, 2002). The Wittig reaction of product **22** with diethyl phosphonoacetonitrile **11** gives (E/Z)-4-chloro-3-methylbut-2-enenitrile **23**. The Arbuzov reaction of triethylphosphite with product **23** gives the reagent (Z/E)-4-(diethylphosphono)-3-methylbut-2-enenitrile **24** and similarly, diphenyl methylphosphite gives the corresponding (Z/E)-4-(diphenylphosphono)-3-methylbut-2-enenitrile **25**.

Ethyl 3-methylbutenoate **26** is obtained by the reaction of acetone and the Wittig reagent **18** (Scheme 2), followed by the reduction with DIBAL-H and subsequent treatment with aqueous HBr to obtain 1-bromo-3-methylbut-3-ene **27**. The latter is coupled with ethyl acetoacetate **21**. Subsequently, the alkylated product is hydrolyzed and decarboxylated to obtain 6-methylhept-5-ene-2-one **28**.

¹³C ENRICHED β -CAROTENE

In Scheme 4 it is indicated that starting from β -ionone **29** with the reagents described in Schemes 2 and 3 [12,12',13,13',14,14',15,15',20,20'-¹³C₁₀]-all-*E* β -carotene **1** is prepared (Lugtenburg *et al.*, 1999). β -Ionone **29** gives a chain extension with diethyl phosphonoacetonitrile **11**, subsequent DIBAL-H reduction gives all-*E* and 9-*Z*- β -ionylidene acetaldehyde **30** which can easily be separated by column chromatography.



Scheme 4. The synthesis of $[12,12',13,13',14,14',15,15',20,20'-1^{3}C_{10}]$ -all-*E* β -carotene 1.

The product **all-**E **30** is treated with anion of (Z/E)-4-(diethylphosphono)-3-methylbut-2-enenitrile **24a**. When the reaction is carried out at above -20° C in the absence of excess base, the Wittig reagent occurs only in the extended form without a trace of geometric isomers. The resulting retinonitrile is formed in the all-E form only, careful DIBAL-H reduction gives all-E retinal **2**.

A subsequent McMurry reductive coupling of all-*E* retinal **2** gives all-*E* β -carotene **1**. The reactions in Scheme 4 have been carried out with ${}^{13}C_5$ -phosphonate **24** resulting in [12,12',13,13',14,14'15,15',20,20'-{}^{13}C_{10}]all-*E* β -carotene **1** with ${}^{13}C$ incorporation of 99% at each of the mentioned positions (Lugtenburg *et al.*, 1999).

It is clear that *via* the reactions in Scheme 4 all-*E* β -carotene **1** can be prepared in any possible ¹³C enriched form that is symmetrically incorporated at the position 12 up to 12' and 20 up to 20'. Curiously carrying out the reactions with the corresponding ¹³C₅-diphenyl building block gives an about 1:1 mixture of for vision research important 11-*Z*-retinal **2** and all-*E* retinal **2** which can easily be separated by column chromatography (Wang *et al.*, 2004). The difference in electronegativity between the ethoxy and phenoxy group on the phosphorous is presumably the cause of this result.

The introduction of better electron withdrawing groups *via* Arbuzov reactions with 1-chloroacetone **22** was unsuccessful. Lately, the conversion of (Z/E)-4-(diethylphosphono)-3-methylbut-2-enenitrile **24** into the corresponding dichloro derivative has been reported (Monbaliu *et al.*, 2010). It is to be expected that *via* this phosphonyl chloride many better electron withdrawing groups can be introduced leading to the hope that using these reagents 11-Z-retinal can be obtained in pure 11-*cis*



Scheme 7. Synthetic Scheme to prepare both $[14-1^{3}C]$ -all-*E* spheroidene and $[14'-1^{3}C]$ -all-*E* spheroidene.







Scheme 6. The preparation of the site directed mono-acetal of the ${}^{\rm 13}{\rm C}_{\rm 10}$ -dialdehyde in all possible ${}^{\rm 13}{\rm C}$ isotopomomers.

form only, which will be an important step for vision research.

U-13C ENRICHED β-CAROTENE

 $[U^{-13}C]$ -All-*E* retinal **2** can be converted easily into $[U^{-13}C]$ -β- carotene **1** *via* the McMurry coupling shown in Scheme 4. The reactions to obtain $[U^{-13}C]$ -all-*E* retinal **2** are shown in Scheme 5 (Creemers & Lugtenburg, 2002). $[U^{-13}C]$ -6-Methylhept-5-ene-2-one $[U^{-13}C]$ -28 prepared *via* Scheme 3 is coupled with ${}^{13}C_2$ -diethyl phosphonoace-tonitrile **11** giving $[U^{-13}C]$ -3,7-dimethylocta-2,6-dieneni-trile $[U^{-13}C]$ -31 which upon treatment with concentrated sulphuric acid gives the α-cyclocitronitrile **31a** and subsequent DIBAL-H treatment gives $[U^{-13}C]$ -α-cyclocitral $[U^{-13}C]$ -32. The α-form is fully converted into the conjugated $[U^{-13}C]$ -β-cyclocitral $[U^{-13}C]$ -33.

Reaction of $[U^{-13}C]^{-33}$ with $(Z/E)^{-4}$ -(diethylphosphono)-3-methylbut-2-enenitrile 24 above $-20^{\circ}C$ and subsequent DIBAL-H reduction gives pure $[U^{-13}C]$ -all- \mathcal{E} β -ionylidene acetaldehyde $[U^{-13}C]$ -all- \mathcal{E} 30. Repeating this sequence results in pure $[U^{-13}C]$ -all- \mathcal{E} retinal $[U^{-13}C]$ -all- \mathcal{E} 2. $[U^{-13}C]$ -All- \mathcal{E} retinal 2 is expected to give $[U^{-13}C]$ -all- \mathcal{E} β -carotene 1 without any problem. It is clear that besides $[U^{-13}C]$ -all- \mathcal{E} β -carotene 1 all

 β -carotenes that have symmetrical ¹³C isotope enrichment are accessible *via* the reactions described in Scheme 5 and the building blocks that are required in ¹³C enriched forms can be prepared according to the literature (Creemers & Lugtenburg, 2002).

The reactions in Scheme 5 are not applicable for carotenoids with different end groups or for asymmetric introduction of ¹³C in symmetric carotenoids. A general approach based on 2,7-dimethylocta-2,4,6-triene-1,8-dial is used in the commercial synthesis of β -carotene **1** *via* Wittig chemistry (Creemers & Lugtenburg, 2002).



Scheme 8. Preparation of the end group of canthaxanthin in any possible ¹³C isotopologues.

In Scheme 6 it is indicated that the acetal of prop-2en-1-al **15** (Scheme 2) can be easily converted into the mono site protected aldehyde of 2,7-dimethylocta-2,4,6triene-1,8-dial **36**. Product **15** is chain extended twice with diethyl phosphonoacetonitrile **11** and DIBAL-H reduction into the acetal conjugated aldehyde **35**. A final chain extension with diethyl phosphonopropionitrile **13** and subsequent reduction gives the required monoacetal of 2,7-dimethylocta-2,4,6-triene-1,8-dial **36**.

In literature the synthesis of building blocks **37** and **39** have been described (Jansen & Lugtenburg, 2009).



 $[4,4'-{}^{13}C_2]-, [12,12'-{}^{13}C_2]-, [13,13'-{}^{13}C_2]-, [14,14'-{}^{13}C_2]-, [15,15'-{}^{13}C_2]-, [20,20'-{}^{13}C_2]-Astaxanthines$





Scheme 10. Reactions to prepare (3,3'-RR,SS,SR)-zeaxanthin 63 in ¹³C enriched form at any carbon position.

The Wittig coupling of the labeled aldehyde **36** with product **37** in refluxing 1,2-epoxybutane and subsequent deprotection gives the conjugated aldehyde **38**.

Repeating this coupling with building block **39**, results in $[14'_{-13}C]$ -spheroidene **40** where as first coupling of **36** with **39**, subsequent deprotection and final coupling with building block **37** gives $[14_{-13}C]$ -spheroidene **40**. These reactions have actually been carried out with the all-*E* $[^{13}C_{10}]$ -7-cyano-2-methylocta-2,4,6-trienal (Jansen, 1996). The reactions in Scheme 7 lead to any carotenoids ^{13}C enriched in any possible way in the central part of the carbon skeleton.

In Scheme 8 the acid catalyzed conversion of 3,7-dimethylocta-2,6-dienenitrile **31** into the cycloderivative **31a** has been discussed. Epoxidation of the double bond with *meta*-chloroperbenzoic acid (*m*-CPBA) gives the epoxide derivative **41**.

Treatment of product **41** with a base gives (4R.5)-4hydroxy- β -cyclocitronitrile **42** (Lugtenburg *et al.*, 1999; Jansen, 2000). Pyridinium chlorochromate oxidation and subsequent ketalisation with ethylene glycol gives compound **43** which upon DIBAL-H reduction gives the 4,4'-protected β -cyclocitral **44**. Wittig-Horner reaction with the ¹³C₅-phosphonato derivative results in the β -ionylidene acetoester **45**, which upon treatment with DIBAL-H and subsequent deprotection gives 4-oxo- β ionylidene ethanol **46**.

In Scheme 9 it is shown that upon treatment with two equivalents LDA, the alcoholate anion of 46 is treated with oxaziridine to give (3*R*.*S*)-3-hydroxy-4-oxo- β -ionylidene ethanol 47. Treatment with triphenyl

phosphine and HBr results in the Wittig salt **48** (Becker *et al.*, 1981). The Wittig coupling of the phosphonium salt **48** with the ${}^{13}C_{10}$ -dialdehyde **49** in 1,2-epoxybutane gives (3,3'-RR,SR,SS)-astaxanthin **50**. The $[4,4'-{}^{13}C_2]$ - and the other isotopologues of astaxanthin indicated in Scheme 9 have been prepared.

In Scheme 10 it is shown that (4R.5)-4hydroxy- β -cyclocitronitrile 42 is converted by acid catalyzed dehydration into the cyclohexadienenitrile 51 (van Wijk *et al.*, 2003). Oxidation with *m*-CPBA acid results epoxidation of 3,4-double bond in product 52. DIBAL-H reduction results in (3R.5)-3-hydroxy- β -cyclocitral 53, chain extension and subsequent DIBAL-H reduction gives the 3-hydroxy- β -ionylidene acetaldehyde 54 followed by protection with TBDMSCI and reduction with NaBH₄ and subsequent reaction with Ph₃P.HBr gives the Wittig salt 56. The lat-

ter could be converted into zeaxanthin **63**. In literature (3R, 3'R)-all-E β -carotene-3,3'-diol enriched with ¹³C at 12,12',13 and 13' has been reported (Khachik *et al.*, 1995). Product **59** can also be converted into (3R*S*)-3-hydroxy all-E retinal **55** which upon McMurry coupling gives the corresponding (3,3',-RR, *SR*, *SS*)-zeaxanthin **63** (Lugtenburg *et al.*, 1999).

In Scheme 11 it is indicated that α -cyclocitral **32** is prepared in any ¹³C isotopologue *via* Scheme 5 that can be converted into α -ionone **57** by treating it with the imine of butylamine and acetone. If a ¹³C enriched isotopologue is required α -cyclocitral **32** could be coupled with the diethyl phosphonoacetonitrile **11** first and then the obtained nitrile could be converted into α -ionone **57**.



Scheme 11. Synthesis of (3*R*)-3-hydroxy- β -ionylidene ethyl triphenyl phosphonium bromide (3*R*)-56 from α -cyclocitral 32.



Scheme 12. Preparation of (3R, 3'R)-zeaxanthin 63 and β -cryptoxanthin 66.

The conversions of α -ionone **57** to **58** has been described in literature (Khachik & Chang, 2011). Protection of the keto function in **58** and subsequent oxidation with *t*-butyl hydroperoxide in aqueous hypochlorite gives the 3-oxo derivative **58**, which with K-Selectride and deprotection of the keto function results in (3*RS*)-3-hydroxy- α -ionone **59** which upon treatment with KOH gives the corresponding (3*RS*)-3-hydroxy- β -ionone **60**.



Scheme 13. Preparation of (3R,3'R,6'R)-lutein **72** starting from α -ionone.

Treatment of (3RJ)-3-hydroxy- β -ionone **60** with vinyl acetate in the presence of lipase gives (3J)-hydroxy- β -ionone **60a** and (3R)-acetoxy- β -ionone **61**. The latter has been converted into (3R,3'R)-zeaxanthin **63** *via* coupling of **(3R)**-**56** and dialdehyde **49** in Scheme 12.

The Schemes discussed so far allow access to any C_{40} carotenoid labeled with ¹³C at any position and combination of positions. The only carotenoid that is not accessible in any ¹³C enriched form is (3R,3'R,6'R)-lutein. Recently the synthesis of (3R,3'R,6'R)-lutein in natural form has been published (Khachik & Chang, 2009). Wittig-Horner reaction of α -ionone 57 gives α -ionylidene acetonitrile 67, which upon treatment with t-butyl hydroperoxide and aqueous hypochlorite solution gives the 3-oxo- α -ionylidene acetonitrile 68. Treatment of 68 with K-Selectride and subsequent reaction with vinyl acetate in the presence of lipase and final DIBAL-H reduction gives the (3R,6R)-3-hydroxy-aionylidene acetaldehyde 69. These reactions have also been used for the conversions in Scheme 11.

Coupling of product (**3***R*,**6***R*)-**69** with the protected ¹³C₁₀-phosphonium salt gives compound **71**. The latter is converted in a final Wittig coupling with (**3***R*)-**56** (Scheme 11) into (3*R*,3'*R*,6'*R*)lutein **72**. For the introduction of ¹³C at any carbon position in lutein, α -ionylidene acetonitrile **67** should preferably be made via α -cyclocitral **32** and coupling with

(Z/E)-4-(diethylphosphono)-3-methylbut-2-enenitrile 24 (Scheme 5).

The phosphonium salt **70** is easily available *via* the aldehyde **36**. *Via* the Schemes in this paper all important C_{40} -carotenoids are now accessible in any ¹³C labeled form at any position and combinations of positions. The Schemes for the introduction of ¹³C discussed so far have a few drawbacks. First, the six membered rings contain two di-

astereotopic methyl groups on carbon 1 and carbon 1': when the methyl groups have one ¹³C and one ¹²C isotope, the quaternary carbon atom 1 will be optically active, giving diastereomeric mixtures that cannot be separated. Second, the introduction of substitution in the six membered rings leads to long linear synthetic Schemes resulting in low yields of these carotenoids with ¹³C enrichment in the six membered rings.

For the solution of both problems we have found that 4-methyl-3-carbethoxypent-3-ene-2-one **73** has been converted in one pot with allyl triphenylphosphonium bromide **74** into the cyclic ester **75** (Büchi & Wuest, 1971). We carried out a Knoevenagel condensation of 1,1-dimethoxyacetone **76** with 1-cyanoacetone **77** to give 3-cyano-5,5dimethoxy-4-methylpent-3-ene-2-one **78** (Dawadi & Lugtenburg, 2007). Reaction of the anion of acetone at position 4 of **78** should give the stable anion **79** analogous to the reaction between **73** and **74**. The anion **79** is expected to form



Scheme 14. Synthesis of well defined six membered systems for the easy access to ¹³C labeled end groups of carotenoids.

the phosphate ester 80 upon treatment with diethyl phosphonyl chloride. Subsequent reaction with base should give 3-oxo-β-cyclocitronitrile 81 (Dawadi & Lugtenburg, 2010). Application of the reduction and enzymatic conversion in Scheme 14 gives the building blocks for the end groups of zeaxanthin and lutein with one of the diasteromeric methyl groups in the form of a protected aldehyde, which will allow a site-directed ¹³C introduction. After deprotection and reduction of the aldehyde end groups will be available with well defined chirality at the carbon position 1.

CONCLUSION

In this paper all known13C enriched C40 carotenoids that have been published are reviewed. The Schemes described in this paper allow access to any biologically and commercially important carotenoid in well defined ¹³C enriched form up to the U-13C enriched system. Combined with the reactions reviewed in the 13C enriched retinal papers all apo-carotenoids are now also accessible in any ¹³C enriched form (Dawadi & Lugtenburg, 2010).

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