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Olive phenols efficiently inhibit the oxidation of serum albumin-bound linoleic acid and butyrylcholine esterase

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Keywords
Antioxidant, olive phenols, serum albumin, butyrylcholine esterase, lipid peroxidation, binding site.
Abstract

Background: Olive phenols are widely consumed in the Mediterranean diet and can be detected in human plasma. Here, the capacity of olive phenols and plasma metabolites to inhibit lipid and protein oxidations is investigated in two plasma models.

Methods: The accumulation of lipid oxidation products issued from the oxidation of linoleic acid bound to human serum albumin (HSA) by AAPH-derived peroxyl radicals is evaluated in the presence and absence of phenolic antioxidants. Phenol binding to HSA is addressed by quenching of the Trp214 fluorescence and displacement of probes (quercetin, dansylsarcosine and dansylamide). Next, the esterase activity of HSA-bound butyrylcholine esterase (BChE) is used as a marker of protein oxidative degradation.

Results: Hydroxytyrosol, oleuropein, caffeic and chlorogenic acids inhibit lipid peroxidation as well as HSA-bound BChE as efficiently as the potent flavonol quercetin. Hydroxycinnamic derivatives bind noncompetitively HSA subdomain IIA whereas no clear site could be identified for hydroxytyrosol derivatives..

Conclusion: in both models, olive phenols and their metabolites are much more efficient inhibitors of lipid and protein oxidations compared to vitamins C and E.

General significance: low postprandial concentrations of olive phenols may help to preserve the integrity of functional proteins and delay the appearance of toxic lipid oxidation products.
Introduction

Oxidative modification of proteins and polyunsaturated fatty acids (PUFA) play a key role in the development of cardiovascular and neurodegenerative diseases [1-4]. These deleterious oxidation processes involve reactive oxygen and nitrogen species (RONS) such as superoxide, nitric oxide and peroxynitrite that can form from circulating blood leukocytes in response to various inflammatory stimuli [5-6]. Fortunately, protection of endogenous biomolecules is afforded by antioxidant enzymes and a broad range of nonenzymatic antioxidants such as urate, glutathione, vitamins C and E and dietary phenols [7]. Consistently, epidemiological data suggest that the so-called mediterranean diet, typically rich in antioxidants from vegetables, fruits, cereals and olive oil, shows significant protective health effects [8]. Olive oil and olives are rich in oleic acid, a monounsaturated fatty acid, and a variety of phenolic antioxidants such as tocopherols, caffeic acid as well as the more typical tyrosol, hydroxytyrosol and their elenolic esters [9,10]. Interestingly, more than 50% of tyrosol and hydroxytyrosol were found to be absorbed through the intestine after ingestion of olive oil by humans [11-13] and only partially excreted in the urine under their native forms or as glucuronides of the native and catechol-O-methylated forms [14]. On the other hand, caffeic acid is recovered in plasma unchanged or metabolized by methylation, hydrogenation, glucuronidation and sulfation [15,16]. Regarding possible health effects, ingestion by humans of extra virgin olive oil significantly decreased inflammatory markers in plasma [17]. Furthermore, hypocholesterolemic and antiatherosclerotic effects were observed after oral administration of olive phenols to rats and rabbits [18,19]. Olive phenols also decrease levels of reactive oxygen species in macrophages, melanoma and endothelial cells, thus preventing protein damage and lipid peroxidation [20-22]. Part of these activities may involve their strong capacity to scavenge free radicals and inhibit lipid peroxidation [23,24]. In particular,
plasma low-density lipoproteins were shown to be protected from oxidation after absorption of a meal rich in olive phenols [25].

Human serum albumin (HSA) is the most abundant protein in serum and a potent plasma antioxidant [26,27]. Albumin dimers have been proposed to be markers of oxidative stress following plasma exposure to peroxides [28]. HSA plays an important role in the transport of endogenous and exogenous ligands, especially fatty acids [29], xenobiotics such as drugs [26] and dietary components including polyphenols [30-33]. Recently, we developed a physiologically-relevant model of antioxidant activity in the plasma based on the ability of dietary antioxidants to inhibit the peroxidation of HSA-bound linoleic acid [34,35].

The plasma protein butyrylcholine esterase (BChE) belongs to the ubiquitous polymorphic family of choline esterases [36]. This enzyme, which is typically extracted in association with HSA [37], catalyzes the hydrolysis of a variety of esters including succinylcholine, a muscular relaxant, cocaine and heroin. BChE seems to be implicated in lipoprotein metabolism since a strong choline esterase activity was detected in the serum of hyperlipidaemic patients [38]. HSA-bound BChE was shown to be protected against oxidation by several naturally occurring phenols [39].

The present study is aimed at determining the capacity of olive phenols to inhibit the oxidation of HSA-bound linoleic acid and butyrylcholine esterase as potential lipid and protein targets of RONS in plasma. The antioxidants selected are tyrosol, hydroxytyrosol, oleuropein and caffeic acid as the major olive phenols, dihydrocaffeic acid and ferulic acid as caffeic acid metabolites in the intestine and plasma, respectively. The antioxidant potential of chlorogenic acid is addressed not only because it is the main dietary source of caffeic acid but also in view of its demonstrated plasma availability [40]. The flavonol
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1 quercetin, L-ascorbate, α-tocopherol and n-hexadecylcaffeate, a chemically synthesized
2 amphiphilic derivative of caffeic acid [24], are also investigated for comparison.

4 Materials and Methods

5 Chemicals

6 Hydroxytyrosol (HT) and n-hexadecylcaffeate were synthesized as previously described
7 [25]. Dansylsarcosine (DNSS), dansylamide (DNSA), caffeic acid, chlorogenic acid,
8 dihydrocaffeic acid, p-coumaric acid, (±) α-tocopherol, quercetin, L-ascorbic acid, 2,2’-
9 azo-bis(2-methylpropionamide dihydrochloride) (AAPH), moxisylyte, linoleic acid
10 (LH), 13(S)-hydroperoxy-(9Z,11E)-octadecadienoic acid, 13(S)-hydroxy-(9Z,11E)-
11 octadecadienoic acid, human serum albumin (fatty acid-free, A-1887 and fraction V, A-
12 1653) were purchased from Sigma-Aldrich (Saint-Quentin Fallavier, France). Ferulic acid,
13 tyrosol and oleuropein were purchased from Extrasynthèse (Genay, France). BCECF
14 (2’,7’-bis(carboxyethyl)-5(6)-carboxyfluorescein) was purchased from Calbiochem (la
15 Jolla, CA, USA). All reagents were of the highest purity available (95-99%) and were used
16 without purification. All solvents used were analytical grade.

17 Inhibition of the peroxidation of HSA-bound linoleic acid

18 HSA (fatty acid free) was dissolved in a 50 mM NaH₂PO₄ - 100 mM NaCl buffer at pH 7.4
19 (Millipore Q-Plus water) previously eluted on Chelex 100 (0.4 mequiv./mL - Bio-Rad) to
20 remove contaminating metal traces. To 3 mL of 0.5 mM HSA was added under stirring 30
21 µL of a freshly prepared 200 mM solution of linoleic acid in MeOH. After 10 min of
22 homogenization, 30 µL of a 5 mM solution of antioxidant in MeOH was added followed 1
23 min later by 42 mg of AAPH. The reaction mixture was stirred in an oven (225 rpm) at 37
24 °C for 10 h. Aliquots (200 µL) were removed every hour for lipid analysis and every 10
min for antioxidant analysis then placed immediately on ice. Protein removal was achieved by acidification with 10 µL of 1 M HCl, addition of 400 µL of a 500 µM solution of L-ascorbic acid in THF/MeCN (2:3), then centrifugation at 10000 g during 3 min. For HPLC analyses, 30 µL of the supernatant were immediately injected. All experiments were run in triplicates.

Analyses of lipid peroxidation products

HPLC analyses were carried out on a Hewlett-Packard 1100 apparatus coupled to a UV-Vis diode array detector. An Alltima C18 reverse phase column (150 mm x 4.6 mm, pre-column of 4.6 x 7.5 mm, particle size 5 µm, Alltech) was used for the chromatographic separations at 35°C. The solvent system was a gradient of A (0.05% aqueous HCO₂H) and B (MeCN) with 5% B at 0 min, 20% B at 4 min, 50% B at 13 min, 60% B at 23 min, 73% B at 38 min and 100% B at 45 min with a flow rate of 1 mL/min. UV detection was set at 234 nm (HPODE and HODE), 280 nm (KODE) or 210 nm (linoleic acid LH). Calibration curves provided the following conversion factors (defined as concentration (µM) = peak area (mAU/min) x Cf): Cf₂₁₀nm (LH) = 1.6, Cf₂₃₄nm (HPODE) = 10, Cf₂₃₄nm (HODE) = 10 and Cf₂₈₀nm (KODE) = 2 [41]. Cf values of HPODE and HODE were obtained with 13(S)-hydroperoxy-(9Z,11E)-octadecadienoic acid and 13(S)-hydroxy-(9Z,11E)-octadecadienoic acid, respectively. Cf of KODE was obtained using a mixture of 4 synthesized isomers.

Antioxidant consumption

The decay of the antioxidants was followed with the same HPLC conditions as above except for the gradient which was 5% B at 0 min, 20% B at 4 min, 50% B at 13 min and 100% B at 15 min. The solvents were a solution of 50 mM KH₂PO₄, 5 mM...
hexadecyltrimethylammonium bromide and 1 mM EDTA at pH 5 for L-ascorbic acid. and
isocratic MeOH for α-tocopherol.

Fluorescence spectroscopy

Tryptophan fluorescence. To 2 mL of 75 µM HSA in a pH 7.4 buffer (50 mM phosphate – 100 mM NaCl) were added via syringe aliquots of 7.5 mM phenol solutions in MeOH. The fluorescence spectra were monitored at 25 °C with a Xenius spectrofluorometer (SAFAS, France) by excitation of the tryptophan residue at 295 nm. The excitation and emission slit widths were set at 10 nm.

Marker displacement. To 2 mL of 75 µM HSA in the above pH 7.4 buffer were added 20 µL of 7.5 mM marker solutions in MeOH. After equilibration, aliquots of 30 mM phenol solutions in MeOH were then added and the fluorescence spectra were monitored at 25 °C by excitation of either HSA-bound quercetin at 450 nm or HSA-bound DNSS or DNSA at 360 nm [30].

Fluorescence intensities were corrected from the inner filter effect of the phenolic ligand according to the following relation [42]:

\[ F_{\text{corr.}} = F_{\text{meas.}} \varepsilon c(l_2-l_1)\ln 10 / (10^{-l_1 c} - 10^{-l_2 c}) \]

Where \( F_{\text{corr.}} \) and \( F_{\text{meas.}} \) are respectively the corrected and measured fluorescence intensities, \( \varepsilon \) is the molar absorption coefficient of the added phenol at the excitation wavelength, \( l_1 \) and \( l_2 \) are the lower and upper distances from the bottom cell where light is transmitted to the photomultiplier and \( c \) is the phenol concentration. In this system, \( l_1 \) and \( l_2 \) were evaluated as 0.45 and 0.85 cm, respectively.

The respective molar absorption coefficients were evaluated in the buffer by UV-vis. spectroscopy. They were calculated as the slope of the absorbance vs. concentration plot for at least 5 incremental aliquots (\( n = 2-3 \)). Couples of \( \varepsilon \) are given in M\(^{-1}\)cm\(^{-1}\) at 295 and
360 nm, respectively: caffeic acid (14100, 900), chlorogenic acid (14000, 5750), ferulic
acid (14470, 600), dihydrocaffeic acid (495, NA), quercetin (9400, NA), oleuropein (690,
NA) and hydroxytyrosol (420, NA).

Inhibition of butyrylcholine esterase oxidation [39]

Fluorescence analyses were carried out on a SAFAS flx apparatus containing a ten-cell
holder (optical pathlength: 1 cm). The temperature was controlled by a water-thermostated
bath. To 3.4 mL of a pH 7.4 phosphate buffer (50 mM KH$_2$PO$_4$ - Na$_2$HPO$_4$, ionic strength
set at the human blood value of 174 mM by KCl) were added 400 µL of a 250 µM solution
of HSA (fraction V) and 40 µL of an antioxidant solution in MeOH (except for n-
hexadecylcafeate dissolved in MeOH/DMSO (1/1)) to a final concentration of 0.1-100
µM. The solution was incubated for 10 min at 40 °C under stirring (150 rpm). Then, 100
µL of a freshly prepared 80 mM solution of AAPH in phosphate buffer were added and the
solution was incubated for 120 min at 40 °C under stirring (150 rpm). Then, the incubated
solution was placed in a quartz cell and 15 µL of a 19.2 µM solution of BCECF in water
followed by 30 µL of a freshly prepared 0.1 M solution of moxisylyte in water were added.
After brief manual stirring, the fluorescence measurement was started at 37 °C over 15 min
(excitation wavelengths: 442 nm and 503 nm, emission wavelength: 530 nm). The
percentage of remaining activity $P_{RA}$ was defined as $100x(p/p_{control})$, $p_{control}$ being the slope
of the $I_{BCECF}$ vs. time plot without antioxidant and AAPH, and $p$ the slope of the $I_{BCECF}$ vs.
time plot with the antioxidant and AAPH (slopes evaluated in the linear portion of the
curves between 4 and 12 min). $P_{RA}$ was plotted as a function of logC (C: antioxidant
concentration, 8 values selected around the EC$_{50}$) to extract pEC$_{50}$ = -logEC$_{50}$.

Data analysis
The Prism V3.0 program (GraphPad Software, San Diego, USA) was used in the curve-fitting procedures using a sigmoid equation for the calculation of EC50 values.

**Results**

In both antioxidant tests, a constant flow of hydrophilic peroxyl radicals is produced via the thermal decomposition of the diazo compound AAPH. The capacity of olive phenols and other antioxidants (Fig. 1) to protect HSA-bound linoleic acid (LH) and butyrylcholine esterase (BChE) against oxidation was evaluated.

**Inhibition of the peroxidation of HSA-bound linoleic acid by olive phenols**

The peroxidation of HSA-bound linoleic acid (C18:2) leads to a mixture of hydroperoxyoctadecadienoic (HPODE), hydroxyoctadecadienoic (HODE) and oxooctadecadienoic (KODE) acids. Within each family, a mixture of four (9/13) regio- and (Z/E) diastereoisomers could be evidenced by mass spectrometry and NMR [42]. After 4 hours, additional products resulting from the cleavage of the hydrocarbon chain could be detected (data not shown).

In the absence of antioxidant, a short lag phase of *ca.* 1 hour was observed in the accumulation of HPODE, HODE and KODE (Fig. 2). Then, the rate of formation of the oxidation products reached a maximal value and eventually decreased. The stronger trend toward saturation was achieved for HPODE, which emerged as the main products of the early phase of peroxidation and as likely precursors of HODE and KODE.

Lipid peroxidation was followed by HPLC for 10 hours with different hydroxytyrosol concentrations (5 - 100 µM). The lowest antioxidant concentration showing significant inhibition was 10 µM. In the presence of 50 µM hydroxytyrosol, no consumption of linoleic acid was detectable during the first 2 hours (Fig. 2). Peroxidation...
then set in with a profile similar to that observed in the absence of antioxidants although the rates of HPODE, HODE and KODE formation remained lower than the maximal rates without antioxidant. In addition, the HODE and KODE concentrations at 10 h were found to be decreased by, respectively, 17 % and 28 %.

For naturally occurring phenols having a 1,2-dihydroxybenzene (catechol) moiety, the lag phase in HPODE formation ($T_{lag}$) is 2-3 times as long as the period required for total antioxidant consumption ($T_a$) (Table 1), which shows that some of their oxidation products still retain a capacity to inhibit the peroxidation of linoleic acid in agreement with our previous observations [34,35]. This phenomenon reaches a maximum with quercetin, which is most rapidly consumed (shortest $T_a$) and however emerges as the most efficient inhibitor (longest $T_{lag}$). The small phenols hydroxytyrosol and caffeic acid display roughly the same efficiency as their more hydrophilic conjugates oleuropein and chlorogenic acid showing that the bulkiness of these hydrophilic substituents does not hinder the antioxidant activity. It just seems that the oxidation products of caffeic acid are better inhibitors than those derived from chlorogenic acid. Unlike catechols, tyrosol and $\alpha$-tocopherol come up as modest inhibitors, which moreover are not totally consumed after lipid peroxidation has resumed ($T_{lag} < T_a$). Remarkably, the amphiphilic $n$-hexadecylcaffeate is both less efficient and less rapidly consumed than caffeic acid. Its consumption coincides with the outburst of peroxidation as if its oxidation products (formed during the lag phase) had no residual antioxidant activity. Finally, the hydrophilic L-ascorbate is rapidly consumed and exerts the weakest protection against lipid peroxidation.

Possible synergies between antioxidants were also checked. Combining L-ascorbate (50 µM) and hydroxytyrosol or caffeic acid (1 µM) in concentrations close to the circulating concentrations did not allow to detect the possible regeneration of the phenolic antioxidants by L-ascorbate. Moreover, 25 µM of hydroxytyrosol combined with 25 µM of
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11 caffeic acid gave an intermediate $T_{lag}$ meaning additive antioxidant effects (data not shown).

3 The antioxidants can be classified from the HPODE concentration accumulated in the presence of a fixed antioxidant concentration and after a given peroxidation period (Fig. 3A): quercetin $>$ caffeic acid $=$ oleuropein $=$ chlorogenic acid $=$ hydroxytyrosol $=$ dihydrocaffeic acid $>$ $n$-hexadecylcaffeate $=$ tyrosol $=$ ferulic acid $=$ $\alpha$-tocopherol $>$ L-

9 ascorbate. The same order was established from their capacity to delay KODE production (Fig. 3B).

10 Binding of olive phenols to HSA

11 The binding of olive phenols to HSA was investigated by fluorescence spectroscopy. The fluorescence of HSA single tryptophan (Trp214) was monitored at 340 nm (ex. 295 nm) upon addition of increasing olive phenol concentrations. At a phenol/HSA ratio of 1, chlorogenic, caffeic and ferulic acids promoted a strong quenching of the Trp214 fluorescence as observed for quercetin [30] (Fig. 4). By contrast, oleuropein, hydroxytyrosol and dihydrocaffeic acid did not modify the hydrophobic environment of Trp214, thus indicating no binding to the pocket accommodating the three hydroxycinnamic acids. Because the various phenols absorb light at the excitation wavelength, a correction for the inner filter effect was applied after evaluation of the respective molar absorption coefficients. Trp214 is part of the so-called subdomain IIA which is known to bind flavonoids [30-31] and drugs such as warfarin [26].

22 The binding site location for olive phenols was also investigated by displacement of probes bound to HSA. DNSA and quercetin were selected as markers of subdomain IIA and DNSS of subdomain IIIA [43-44]. In a first series of experiments, olive phenols were added to albumin and quercetin held in equimolar concentrations (75 µM). With a mean
binding constant of $125 \times 10^3 \text{M}^{-1}$ [30-31], 72% of quercetin is bound to HSA. For an olive phenol/albumin ratio of 20, the decrease of the fluorescence intensity of HSA-bound quercetin at 530 nm (excitation at 450 nm) was less than 30% for ferulic, caffeic and chlorogenic acids and around 10% for hydroxytyrosol and oleuropein. Although the three hydroxycinnamic acids are more modest HSA ligands than quercetin [33], their inability to displace quercetin from its binding site suggests that their interaction with HSA does not take place in competition with quercetin and rather points to the formation of ternary complexes and subsequent weak conformational changes of HSA. This may also be the case for HT and its elenolic derivative oleuropein as evidenced by the fluorescence patterns displayed at higher albumin-ligand molar ratios.

When olive phenols were added to DNSS/HSA complexes ($K_b = 1.3 \times 10^5 \text{M}^{-1}$ [44], 73% bound), the fluorescence was more strongly quenched than with the quercetin probe. Chlorogenic acid efficiently displaces DNSS, ranked in the literature as a subdomain IIIA probe, leading to a fluorescence quenching of 50% for a chlorogenic acid/HSA ratio of 3. By contrast, hydroxytyrosol had no effect on the DNSS-HSA fluorescence. Unexpectedly, similar fluorescence quenching patterns were observed for DNSA, which is considered a subdomain IIA probe (data not shown). Thus, it can be proposed that either olive phenols bind without selectivity to both subdomains IIA and IIIA or that DNSS and DNSA are in fact markers for a same subdomain. Low-affinity binding sites for dansyl probes have also been proposed [44] as observed for ANS (1-anilino-8-naphtalenesulfonamide), a structurally related probe which binds both subdomain IIIA ($K_b = 8.7 \times 10^5 \text{M}^{-1}$) and subdomain IIA ($K_b = 7.9 \times 10^4 \text{M}^{-1}$) [45].

**Inhibition of the oxidation of HSA-bound butyrylcholine esterase**
The experiments were conducted in the presence of moxisylyte, an ester substrate of BChE, and the pH-sensitive fluorescent indicator BCECF [39]. The pH drop induced by moxisylyte hydrolysis quenches the fluorescence of BCECF. In the presence of AAPH, the oxidative degradation of BChE strongly attenuates the quenching of BCECF fluorescence. Intermediate situations are observed in the presence of an antioxidant (Fig. 6). Over the typical period of the enzymatic experiments, both moxisylyte and BCECF did not undergo significant oxidative degradation (data not shown). Ferulic acid and the antioxidants having a catechol moiety are the most efficient at inhibiting the oxidative degradation of BChE (EC$_{50} < 10 \mu$M) (Table 2). The differences between them are not statistically significant. Ascorbate and tyrosol are both much less active, a likely consequence of the poor stability of the former and the weak reducing potential of the latter.

**Discussion**

After oral absorption, olive phenols are recovered (mostly as conjugates) into the human blood stream [11-14] where they could develop interesting antioxidant activities (in addition to more specific effects following distribution to other tissues). The blood plasma contains important biomolecules that are sensitive to oxidative stress such as polyunsaturated fatty acids and proteins. Moreover, the major plasma protein serum albumin (0.53 - 0.68 mM in adult blood) is the transporter of free fatty acids and a wide range of xenobiotics [26]. HSA displays three repetitive binding domains named I, II, III, in which long-chain fatty acids are asymmetrically distributed over up to 7 binding sites [29]. HSA-bound fatty acids circulate in blood plasma at a concentration near 1 mM. Among them, PUFA are essential lipids that are sensitive to oxidation processes leading to potentially toxic hydroperoxides and aldehydes via highly oxidizing intermediates such as oxyl and peroxyl radicals [46]. In plasma, such PUFA oxidation products could mediate
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oxidative modifications of HSA (carbonylation, loss of Lys residues) [47,48]. In this work, olive phenols and other common antioxidants are tested for their capacity to inhibit the oxidative degradation of linoleic acid and butyrylcholine esterase in the presence of human serum albumin.

Linoleic acid, the most abundant PUFA in vivo, was selected as the first potential target of ROS in plasma. The linoleic acid/HSA molar ratio was set at 4:1 so as to mimic metabolic conditions reached during strenuous exercise and in diabetes (normal physiological ratio 2:1) [26]. The protection by antioxidants of HSA-bound linoleic acid against hydrophilic peroxyl radicals generated in the aqueous phase must be governed by the two following factors:

- the intrinsic reactivity of the antioxidant at reducing peroxyl radicals. H-atom or electron transfer from the antioxidant to the peroxyl radicals is largely governed by the stability of the radical derived from the antioxidant. For phenolic antioxidants, it is well known that semiquinone radicals derived from catechol (1,2-dihydroxybenzene) nuclei are stabilized by a combination of electronic and intramolecular hydrogen bonding effects. Consequently, antioxidants having a catechol group typically come up as more efficient than analogs having a single phenolic OH group.

- the affinity of the antioxidant for HSA (measurement by the binding constant $K_b$) and its availability to the peroxyl radicals. Dietary flavonoids such as quercetin that are partially absorbed from the gastro-intestinal tract are detected in plasma as HSA-bound conjugates [49]. Up to now, binding constants to HSA were not available for the true circulating polyphenol conjugates. For commercially available polyphenols (aglycones, glycosides) and chemically synthesized models of conjugates (flavonoid sulfates), $K_b$ values have been evaluated in the range $10^4$ – $10^5$ M$^{-1}$ [30-33]. This moderate to low affinity is sufficient to ensure that quercetin and the hydroxycinnamic acids investigated in this work are
predominantly bound to HSA given the high HSA/ligand molar ratio used, which is expected to be even higher in the plasma conditions. Because data dealing with the affinity of hydroxytyrosol derivatives towards HSA are lacking, an insight was brought into the location and affinity of these small ligands. Ferulic acid, caffeic acid and its quinic ester chlorogenic acid all quenched Trp214 fluorescence as efficiently as quercetin suggesting binding in subdomain IIA or in proximity of the interdomain IIA/IIIA. By contrast, dihydrocaffeic acid, HT and oleuropein showed no quenching and thus do not display a high-affinity binding site in these regions. Displacement studies were next conducted to refine the location of olive phenols. The three hydroxycinnamic acids are slightly weaker HSA ligands than quercetin ($K_b$ (ferulic acid/HSA) = $2.2 \times 10^4$ M$^{-1}$, $K_b$ (chlorogenic acid/HSA) = $4.4 \times 10^4$ M$^{-1}$) [33]. However, if they bind to HSA in competition with quercetin, their affinity for HSA would be sufficient to totally quench the fluorescence of HSA-bound quercetin given the high ligand/quercetin molar ratios used. This is by far not the case and rather points to noncompetitive binding. Unexpected results were obtained in the displacement of specific probes for subdomains IIA (DNSA) and IIIA (DNSS) in that similar fluorescence patterns were observed. Again, the three hydroxycinnamic acids deeply affected the binding of the dansyl ligands to HSA whereas the small-sized hydroxytyrosol showed no effect. Multiple binding sites may be suggested for hydroxycinnamic acids whereas no clear site could be identified for hydroxytyrosol and oleuropein.

A detailed investigation has shown that quercetin bound to HSA ($K_b \approx 125\times10^3$ M$^{-1}$) is oxidized as rapidly as free quercetin under the flow of peroxyl radicals produced by the thermal decomposition of AAPH as if the HSA matrix exerted no screening between the peroxyl radicals generated in the aqueous phase and quercetin in its binding cavity within subdomain IIA [35]. This observation suggests long-range electron transfer processes
operating between the periphery of the protein (the primary sites of radical attack) and the quercetin binding site. The lag phase observed in the peroxidation of HSA-bound linoleic acid has been interpreted as the period needed for AAPH-derived peroxyl radicals to oxidize some peripheral HSA residues thereby triggering a minimal conformational change that leads to LH molecules being more exposed to the protein-borne radicals. After quercetin, the most efficient peroxidation inhibitors are caffeic acid, chlorogenic acid and the typical olive antioxidants hydroxytyrosol and oleuropein. Among these four hydrophilic antioxidants, only the two first are known to be (weak) HSA ligands. Hence, hydrophilic antioxidants with a catechol group clearly emerge as the best peroxidation inhibitors as if their activity was governed by their intrinsic H-donating properties more than by their binding to HSA. Moreover, the presence of a large hydrophilic sub-unit that could be expected to alter the binding to HSA (caffeic acid vs chlorogenic acid, hydroxytyrosol vs oleuropein) does not significantly affect the antioxidant efficiency.

After the lag phase, the peroxidation rate quickly increases to reach 80 to 90% of that of the uninhibited peroxidation ($R_p^{\text{max}} \approx 1.08 \, \mu \text{M min}^{-1}$, Table 1). This value is much lower than the initiation rate, i.e. the flow of AAPH-derived peroxyl radicals in the aqueous phase (based on the literature [35], $R_i \approx 14 \, \mu \text{M min}^{-1}$ in our conditions). Hence, even when the peroxidation rate is maximal, linoleic acid is largely protected from the attacking peroxyl radicals by the HSA matrix. From the quasi-linear decay of the antioxidant concentration, the rate of antioxidant consumption ($R_a$) can also be estimated (Table 1). Potent antioxidants are rapidly consumed with $R_a$ values in the range 0.8-1.3 $\mu \text{M min}^{-1}$ (period required for complete consumption of the antioxidant $T_a$ in the range 50-60 min). By contrast, $\alpha$-tocopherol and tyrosol, which are poor peroxidation inhibitors, are both more stable ($R_a < 0.4 \, \mu \text{M min}^{-1}$, $T_a > 180$ min). Once more, the rates of antioxidant consumption are much lower than the initiation rate as a consequence of the competition
between the antioxidant and other H-donating species during the lag phase. These species include not only some exposed reducing residues of HSA but also some oxidation products of the antioxidants in which the catechol nuclei are regenerated as a consequence of radical recombination (dimerization, disproportionation) or water addition on quinone or quinonemethide intermediates [24]. It is also noteworthy that the distribution of caffeic acid oxidation products (mainly dimers with biaryl or biaryl ether linkages [24]) is unaffected by the HSA matrix (data not shown). Those observations could be more consistent with an inhibition mechanism involving the unbound fraction in the aqueous phase. Remarkably, α-tocopherol, normally a potent inhibitor of lipid peroxidation, comes up as a poor antioxidant in this model. Its good solubility in the medium (despite its hydrophobic character) may be taken as a clue of its binding to HSA, which could substantially mask its activity. It can also be added that α-tocopherol is especially potent at inhibiting lipid peroxidation in systems where propagation can take place (chain-breaking inhibition by scavenging of lipid peroxyl radicals). However, the encounter between lipid peroxyl radicals (LOO•) and linoleic acid molecules (LH) is an unlikely event in this model where each LH molecule is tightly sequestered within its binding pocket. Finally, L-ascorbate is unique in that it is rapidly consumed while providing only a marginal protection of linoleic acid. These observations are consistent with the high instability of L-ascorbate in the medium reflecting its fast autoxidation rather than its efficient scavenging of peroxyl radicals.

A simple kinetic model can be developed to obtain a semi-quantitative picture of the sequence of oxidative events taking place within the HSA-linoleic acid complex in the absence or presence of an antioxidant. In our system, a large concentration of diazo initiator AAPH ensures a constant flow of hydrophilic peroxyl radicals ROO• (rate R.) that
abstract H atoms from the antioxidant (AH), lipid (LH), or oxidizable protein residues (PH) [Eqs. (1)-(3)]:

$\text{ROO}^\bullet + \text{AH} \xrightarrow{k_{AH}} \text{ROOH} + \text{A}^\bullet$ \hspace{1cm} (1)

$\text{ROO}^\bullet + \text{LH} \xrightarrow{k_{LH}, O_2} \text{ROOH} + \text{LOO}^\bullet$ \hspace{1cm} (2)

$\text{ROO}^\bullet + \text{PH} \xrightarrow{k_{PH}} \text{ROOH} + \text{P}^\bullet$ \hspace{1cm} (3)

The steady-state hypothesis for \( \text{ROO}^\bullet \) is expressed as [Eq. (4)]:

\[ R_i = [\text{ROO}^\bullet](k_{AH}[\text{AH}] + k_{LH}[\text{LH}] + k_{PH}[\text{PH}]) \] \hspace{1cm} (4)

From these relationships, the chemical rates for the consumption of AH and LH are readily deduced [Eqs. (5)-(6)]:

\[ R_{AH} = \frac{d}{dt} [\text{AH}] = k_{AH}[\text{AH}][\text{ROO}^\bullet] = \frac{R_i}{1 + r_{LH}[\text{LH}]/[\text{AH}] + C_{PH}/[\text{AH}]} \] \hspace{1cm} (5)

With \( r_{LH} = k_{LH}/k_{AH}, C_{PH} = k_{PH}[\text{PH}]/k_{AH} \)

\[ R_{LH} = \frac{d}{dt} [\text{LH}] = k_{LH}[\text{LH}][\text{ROO}^\bullet] = R_{AH} r_{LH} \frac{[\text{LH}]}{[\text{AH}]} \] \hspace{1cm} (6)

In fact, an antioxidant of stoichiometry \( n \) can be modelled as \( n \) identical AH sub-units that all transfer a single H-atom with rate constant \( k_{AH} \). We may also write: \([\text{AH}] = nC_a, R_{AH} =\)
\[ nR_a, C_a \text{ being the true antioxidant concentration (typically, } 50 \mu M) \text{ and } R_a \text{ the true rate of } \]

antioxidant consumption (estimated by HPLC titrations). Hence, we have:

\[ R_a = \frac{R_i}{n + r_{LH}[LH] / C_a + C_{PH} / C_a} \quad (7) \]

In fact, the antioxidant is consumed during the lag phase, i.e. when LH consumption is negligible. Hence, Eq. (7) can be simplified as follows:

\[ R_a = \frac{R_i}{n + C_{PH} / C_a} \quad (8) \]

After the lag phase, LH consumption is maximized as the consequence of the consumption of the antioxidant and some specific HSA residues (responsible for the lag phase observed in uninhibited peroxidation). Hence, only the bulk of oxidizable HSA residues are left to compete with linoleic acid for the attacking peroxyl radicals:

\[ R_{LH}^{\text{max}} = \frac{R_i}{1 + C_{PH}^{\text{bulk}} / r_{LH}[LH]} = \frac{R_i}{1 + AE_{PH}^{\text{bulk}} [PH^{\text{bulk}}]/[LH]} \quad (9) \]

With \( AE_{PH}^{\text{bulk}} = k_{PH}^{\text{bulk}} / k_{LH} \)

If only the oxidation of the antioxidant and specific HSA residues are considered during the lag phase, we may write:
Taking \( R_i = 14 \, \mu M \, \text{min}^{-1}, \) \( T^0_{\text{lag}} = 60 \, \text{min} \) (no antioxidant), we obtain: \([PH^{sp}] = 0.84 \, \text{mM}, \) i.e. 1.7 residues per HSA molecule. For potent antioxidants (50 \( \mu M \)), \( T_{\text{lag}} \approx 150 \, \text{min} \). Thus, we have: \( n \approx 25 \), which is unrealistic. Consequently, we must assume that PH residues from the bulk already contribute. Taking a more reasonable \( n \) value of 4 [24], we obtain from Eq. (10): \([PH] = 1.9 \, \text{mM}, \) i.e. almost 4 residues per HSA molecule.

For potent antioxidants during the lag phase, we also have: \( R_a \approx 1 \, \mu M \, \text{min}^{-1} \).

From Eq. (8), we thus deduce: \( C_{PH} = k_{PH}[PH]/k_{AH} \approx 10C_a = 0.5 \, \text{mM} \). Hence, \( k_{AH}/k_{PH} \approx 4 \).

Finally, we have: \( R_{p_{\text{max}}} \approx 1 \, \mu M \, \text{min}^{-1} \). Taking \([LH] = 1.8 \, \text{mM} \) at \( R_{p_{\text{max}}} \), we deduce from Eq. (9): \( AE_{PH} \approx k_{PH}/k_{LH} \approx 12 \).

Although very crude, these calculations provide a deeper insight in the mechanisms at work. In summary:

- Uninhibited peroxidation starts after a minimal oxidation of HSA at 1-2 critical residues during the lag phase.

- In the presence of potent antioxidants, the lag phase is much longer and likely involves additional HSA residues. However, the oxidation of the antioxidant is faster than that taking place on the protein.

- After the lag phase, peroxidation of linoleic acid is significant but remains slow because of its competition with oxidation at several HSA residues (\( ca. \) 4), which as a whole is one order of magnitude faster. Potential targets of radical attack on HSA are Trp-214, Cys-34, Tyr (18 residues), His (16 residues) and Met (6 residues) [27,50,51].
Finally, olive phenols are able to inhibit the oxidation of butyrylcholine esterase co-extracted with HSA (Table 2) [39]. EC$_{50}$ values of all phenolic antioxidants except tyrosol lie in the range 2.5 - 6.9 µM. This poor discrimination between the antioxidants despite their structural diversity and variable affinity for HSA once more suggests that a strong H-donating capacity is the main factor for a potent inhibition, which in most cases is likely to operate via direct scavenging of the AAPH-derived peroxyl radicals by the unbound antioxidant fraction in the aqueous phase.

## Conclusion

The biological significance of the antioxidant activity of dietary phenols has been strongly questioned based on their relatively low bioavailability and extensive conjugation after intestinal absorption (a process that may lower the reducing activity of phenolic nuclei). Anyway, the inhibition of lipid and protein oxidation remains an important challenge in terms of preserving important nutrients and avoiding the formation of potentially toxic hydroperoxides and carbonyl compounds. A deeper insight in the chemical mechanisms sustaining the antioxidant activity of dietary phenols in biologically-relevant systems involving lipids and proteins brought some unexpected results. Catechol-bearing dietary phenols, whether strongly or weakly bound to serum albumin (e.g., quercetin and hydroxytyrosol), are much more efficient at inhibiting lipid and protein peroxidation than recognized antioxidants such as vitamins C and E. Finally, although the postprandial plasma concentration of dietary phenols is much lower than the circulating concentrations of α-tocopherol and ascorbate, dietary phenols could help protect HSA itself against oxidative modifications, thereby maintaining its function as a transport protein and a plasma antioxidant.
Acknowledgments

M. Roche is grateful to the General Council of the PACA region and to INRA for financial support and AFIDOL for their interest in this study.

List of Abbreviations

HSA, human serum albumin; BChE, butyrylcholine esterase; PUFA, polyunsaturated fatty acid; HODE, hydroxyoctadecadienoic acid; HPODE, hydroperoxyoctadecadienoic acid; KODE, oxooctadecadienoic acid; AAPH, 2,2’-azo-bis(2-methylpropionamidine dihydrochloride).

References

Olive phenols as antioxidants


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Figure legends

Fig. 1. Chemical structures of the selected antioxidants.

Fig. 2. Oxidation of HSA-bound linoleic acid at 37 °C. Open symbols: inhibition by hydroxytyrosol (HT), close symbols: control without antioxidant. A: Time dependence of total lipids (▲,△) and linoleic acid (■,□). B: HPODE accumulation (■,□) and HT consumption (○). C: HODE accumulation. D: KODE accumulation. Initial concentrations: 0.5 mM HSA, 2 mM LH, 50 mM AAPH, 50 μM HT.

Fig. 3. Inhibition of the oxidation of HSA-bound linoleic acid by the selected antioxidants. A: concentrations of HPODE accumulated after 3 h. B: concentrations of KODE accumulated after 4 h (antioxidant concentration = 50 μM).

Fig. 4. Quenching of HSA Trp fluorescence by the selected phenols. Dihydrocaffeic acid (open circle), hydroxytyrosol (close square), oleuropein (close diamond), caffeic acid (cross), ferulic acid (close circle), quercetin (open square), chlorogenic acid (close triangle) and MeOH (dashed line). HSA concentration = 75 μM. n = 2 (mean dev.<10%)

Fig. 5. Influence of selected phenols on the fluorescence of HSA-bound quercetin (A) and dansylsarcosine (B). Hydroxytyrosol (close square), oleuropein (close diamond), caffeic acid (cross), ferulic acid (close circle), chlorogenic acid (close triangle) and MeOH (dashed line). HSA and marker concentrations = 75 μM. n = 2 (mean dev.<10%)

Fig. 6. Inhibition of the oxidative degradation of HSA-bound BChE by hydroxytyrosol (HT). A: Time dependence of the BCECF fluorescence intensity at 530 nm in the presence...
of different HT concentrations (C) at 40°C. B: Plot of the percentage of remaining BChE activity as a function of logC.
Olive phenols as antioxidants

Fig. 1
Fig. 2
Fig. 3
Fig. 4
Olive phenols as antioxidants

**Fig. 5A**

**Fig. 5 B**
Olive phenols as antioxidants

**Fig. 6**
Table 1. Inhibition of the peroxidation of HSA-bound linoleic acid (initial concentrations: 0.5 mM HSA, 2 mM LH, 50 mM AAPH, 50 µM antioxidant).

<table>
<thead>
<tr>
<th>Antioxidant</th>
<th>$T_{lag}$ (min)$^a$</th>
<th>$T_{a}$ (min)$^b$</th>
<th>$R_p^{max}$ (µM/min)$^c$</th>
<th>$R_a$ (µM/min)$^d$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quercetin</td>
<td>163 (± 3)</td>
<td>50</td>
<td>0.92 (± 0.06)</td>
<td>1.11 (± 0.02)</td>
</tr>
<tr>
<td>Caffeic acid</td>
<td>153 (± 11)</td>
<td>60</td>
<td>0.95 (± 0.06)</td>
<td>0.81 (± 0.03)</td>
</tr>
<tr>
<td>Chlorogenic acid</td>
<td>141 (± 4)</td>
<td>60</td>
<td>0.91 (± 0.14)</td>
<td>0.89 (± 0.02)</td>
</tr>
<tr>
<td>Oleuropein</td>
<td>142 (± 9)</td>
<td>60</td>
<td>0.83 (± 0.01)</td>
<td>0.80 (± 0.01)</td>
</tr>
<tr>
<td>Hydroxytyrosol</td>
<td>141 (± 2)</td>
<td>50</td>
<td>0.92 (± 0.02)</td>
<td>1.33 (± 0.11)</td>
</tr>
<tr>
<td>Dihydrocaffeic acid</td>
<td>131 (± 1)</td>
<td>60</td>
<td>0.86 (0.10)</td>
<td>0.91 (0.01)</td>
</tr>
<tr>
<td>n-Hexadecylcaffeate</td>
<td>118 (± 10)</td>
<td>120</td>
<td>1.01 (± 0.05)</td>
<td>0.41 (± 0.01)</td>
</tr>
<tr>
<td>Ferulic acid</td>
<td>110 (± 8)</td>
<td>80</td>
<td>1.19 (± 0.08)</td>
<td>0.61 (± 0.04)</td>
</tr>
<tr>
<td>Tyrosol</td>
<td>86 (± 1)</td>
<td>240</td>
<td>0.78 (± 0.08)</td>
<td>0.26 (± 0.01)</td>
</tr>
<tr>
<td>α-Tocopherol</td>
<td>88 (± 2)</td>
<td>180</td>
<td>0.92 (± 0.07)</td>
<td>0.39 (± 0.02)</td>
</tr>
<tr>
<td>L-Ascorbate</td>
<td>68 (± 7)</td>
<td>40</td>
<td>1.12 (± 0.01)</td>
<td>1.03 (± 0.03)</td>
</tr>
<tr>
<td>Control</td>
<td>56 (± 2)</td>
<td>-</td>
<td>1.08 (± 0.09)</td>
<td>-</td>
</tr>
</tbody>
</table>

$^a$ Lag phase of the [HPODE] vs. time curve; $n = 2 - 4$. $^b$ Period required for complete consumption of the antioxidant (estimated error from duplicates = ± 10 min). $^c$ Maximal peroxidation rate, calculated from $t = 3$ to $5$ h (from $t = 1$ to $3$ h for the control) of the [HPODE] vs. time curve. Values are means (SD), $n = 2 - 4$. $^d$ Rate of antioxidant consumption.
Table 2. Inhibition of the oxidation of HSA-bound BChE (initial concentrations: 25 µM HSA, 2 mM AAPH, 72 nM BCECF, 750 µM moxisylyte).

<table>
<thead>
<tr>
<th>Compound</th>
<th>pEC$_{50}$</th>
<th>EC$_{50}$ (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quercetin</td>
<td>5.61 (± 0.12)</td>
<td>2.5 [1.4 - 4.3]</td>
</tr>
<tr>
<td>Dihydrocaffeic acid</td>
<td>5.48 (± 0.09)</td>
<td>3.3 [2.1 - 5.1]</td>
</tr>
<tr>
<td>Chlorogenic acid</td>
<td>5.46 (± 0.13)</td>
<td>3.5 [1.9 - 6.6]</td>
</tr>
<tr>
<td>Caffeic acid</td>
<td>5.32 (± 0.10)</td>
<td>4.8 [3.0 - 7.5]</td>
</tr>
<tr>
<td>Oleuropein</td>
<td>5.31 (± 0.15)</td>
<td>4.9 [2.4 - 10.0]</td>
</tr>
<tr>
<td>n-Hexadecylcaffeate</td>
<td>5.31 (± 0.27)</td>
<td>4.9 [1.3 - 19.1]</td>
</tr>
<tr>
<td>Ferulic acid</td>
<td>5.27 (± 0.10)</td>
<td>5.4 [3.5 - 8.5]</td>
</tr>
<tr>
<td>Hydroxytyrosol</td>
<td>5.16 (± 0.11)</td>
<td>6.9 [4.1 – 11.7]</td>
</tr>
<tr>
<td>L-Ascorbate</td>
<td>4.46 (± 0.10)</td>
<td>35 [21.5 – 56.9]</td>
</tr>
<tr>
<td>Tyrosol</td>
<td>&lt; 3.84</td>
<td>&gt; 140 [nd]</td>
</tr>
</tbody>
</table>

a pEC$_{50}$ = -logEC$_{50}$; values are means (SD) n = 4; b confidence interval of EC$_{50}$ is valid at 95 %. 