Structure–activity relationships of fraxamoside as an unusual xanthine oxidase inhibitor

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ABSTRACT

Fraxamoside, a macrocyclic secoiridoid glucoside featuring a hydroxytyrosol group, was recently identified as a xanthine oxidase inhibitor (XOI) comparable in potency in vitro to the standard antigout drug allopurinol. However, this activity and its considerably higher value than its derivatives oleuropein, oleoside 11-methyl ester, and hydroxytyrosol are not explained by structure–activity relationships (SARs) of known XOIs. To exclude allosteric mechanisms, we first determined the inhibition kinetic of fraxamoside. The resulting competitive mechanism prompted a computational SAR characterization, combining molecular docking and dynamics, which fully explained the behavior of fraxamoside and its derivatives, attributed the higher activity of the former to conformational properties of its macrocycle, and showed a substantial contribution of the glycosidic moiety to binding, in striking contrast with glycoside derivatives of most other XOIs. Overall, fraxamoside emerged as a lead compound for a new class of XOIs potentially characterized by reduced interference with purine metabolism.

Introduction

Xanthine oxidoreductase (XOR) is a 290 kDa dimeric molybdenum-containing enzyme involved in the catabolism of purines in humans. Each subunit contains one molybdopterin cofactor, two distinct [2Fe–2S] centers, and one FAD cofactor and catalyzes the oxidation of hypoxanthine to xanthine and xanthine to uric acid at the molybdopterin center, with concomitant reduction of NAD+ to NADH at the FAD center. XOR may act as dehydrogenase (XDH) and oxidase (XO): the XDH utilizes NAD+ or NADH as final electron acceptor, whereas XO uses exclusively O2 as electron acceptor, generating O2•− and H2O2 during its mechanism of reaction.

Since XO generates significant amounts of ROS, it represents an important source of free radicals, potentially determining oxidative damage to living tissues. Furthermore, XO is deeply involved into the development of hyperuricemia, a risk factor for gout, cardiovascular diseases, hypertension, and metabolic syndrome. The contribution of the glycosidic moiety to binding, in striking contrast with glycoside derivatives of most other XOIs. Overall, fraxamoside emerged as a lead compound for a new class of XOIs potentially characterized by reduced interference with purine metabolism.

Fraxamoside is a macrocyclic secoiridoid glucoside isolated from Fraxinus americana and Olea europaea L. leaves decoction, also containing, among many others, the compounds: oleuropein (Compound 2) and oleoside 11-methyl ester (Compound 3), and hydroxytyrosol (Compound 4) (Figure 1(a)). These four molecules are strictly related to each other, sinceCompound 2 derives from the condensation of Compounds 3 and 4, and Compound 1 is the product of cyclization of Compound 2. Thus, these compounds feature different combinations or topologies of three moieties, illustrated in Figure 1(b) for Compound 1: a hydroxytyrosol group (hty), present in Compounds 1, 2, and 4, both a secoiridoid moiety (sit) and a β-D-glucose unit (βgl), contained in Compounds 1–3. Compounds 1 and 2 share
the same combination and connectivity of groups, except for the extra ethereal bond in 1 between the alcoholic function at C6’ of ηgl and the C2” atom of hty, with consequent formation in this compound of a 14-membered ring.

Interestingly, Compounds 1–3 (especially 1) represent a significant departure from traditional scaffolds active on XO, that, independently on their natural or synthetic origin and on their purine-like or non-purine nature, are almost invariably characterized by largely planar structures.

Methods
Reagents
Xanthine was purchased from Fluka; Xanthine oxidase from bovine milk (0.2 U/mg), DMSO, allopurinol, FeCl2, and XO from Sigma-Aldrich (St. Louis, MO). Ethanol was purchased from Merck (Darmstadt, Germany). Fraxamoside, extracted from Olea europaea L. leaf decoction, was kindly supplied by Prof. M. Iorizzi.

XO activity assay
XO catalyzes uric acid formation through oxidation of xanthine (or hypoxanthine) and therefore its activity can be monitored by following the increase of uric acid formation by the absorbance at 295 nm (Cary50 UV-Vis Spectrophotometer equipped with a Peltier thermoelectric controller, Varian Medical Systems, Inc.)21,22.

The assay was carried out in a 500-μL final volume reaction mixture, containing 50 mM phosphate buffer, pH 7.8, 0.1 mM EDTA, 50 μM xanthine as substrate, 3.2 U/L XO, and different amounts of each compound as potential XO inhibitor. The assay was started by adding 10 μL of 160 U/L XO solution to the reaction mixture, without or with inhibitors, and it was followed in kinetics by reading the absorbance up to 2 min at 20 °C, thus calculating the rate of acid uric formation in the linear range of the reaction. The kinetic parameters of the XO reaction were derived from Lineweaver–Burk plots obtained measuring the rate of uric acid formation at different xanthine concentration (ranging between 0.4 and 10 μM). Inhibition studies were carried out by measuring the kinetic parameters in the presence of different inhibitors. The Ki of fraxamoside and allopurinol was measured by the abscissa intercept of the secondary plots of apparent Km (K’m) versus [I], according to the following equation: \( K'_{m} = \frac{[I]}{K_{m}/K_i + K_{m}} \).

Statistical analysis
The kinetic parameters of XO were determined by Lineweaver–Burk plot using a standard curve-fitting program. At each single concentration of substrate, the activity was evaluated in triplicate. The correlation coefficient \( R^2 \) between the concentration of the samples and the relative effect was always greater than 0.900. Correlation and regression analyses and statistical analysis were made with Student’s t-test, using the software KaleidaGraph (Reading, PA). p values lower than 0.05 were considered to be significant.

Conformational analysis of Compound 1 and parameterization of Compounds 1–3 for amber force field
Starting ligand geometries were built with Gmealcal 2.992,23 in a standard extended geometry for oleuropein and Compound 3, in an arbitrary strainless macrocycle ring conformation for

Figure 1. XO ligand structures, atom, and fragment naming. (a) The structures of fraxamoside (Compound 1), oleuropein (Compound 2), oleoside 11-methyl ester (Compound 3), and hydroxytyrosol (Compound 4) are shown along with the atom naming scheme (reported for 2). (b) Representation of the different fragments discussed in the paper, highlighted for 2 by a dark gray area (hty) with a medium gray enclosure surrounded by a dotted line (cathecol), a light gray area ([ηgl]), and a thick black contour line (air).
fraxamoside, and they underwent energy minimization (EM) at molecular mechanics level first, using Tripos 5.2 force field parametrization\textsuperscript{24}, and then at AM1 semi-empirical level. To explore the conformational space accessible to Compound 1 macrocycle, which is kept rigid in subsequent docking calculations, a molecular dynamics (MD)-based simulated annealing approach (SA) was adopted, by applying cycles of MD with the following general temperature profile: 10 ps of heating from 1 K to $T_{\text{max}}$ 50 ps of simulation at $T_{\text{max}}$ and 10 ps of cooling from $T_{\text{max}}$ to 1 K, with a starting value of 500 K of $T_{\text{max}}$. A final EM provided the new sampled conformer. $T_{\text{max}}$ was varied between 500 and 1500 K according to the following protocol: it was kept constant until three SA cycles produced no new structure within 10 kcal mol$^{-1}$ from the best current conformer, in which case it was increased by 200 K. If good structures were located at this higher temperature, $T_{\text{max}}$ was decreased by 200 K, and the procedure was reiterated. After reaching an upper limit of 1500 K, the SA was terminated if no new good structure was obtained after nine more SA cycles. The default Chemical MD values were used for all other MD parameters involved in the SA procedure.

The fraxamoside structures selected from SA underwent geometry optimization with the ORCA 3.0.2 program\textsuperscript{25} at the DFT level, using B3LYP functional\textsuperscript{26,27}, Def2-TZVP basis set\textsuperscript{28,29}, Def2-TZVPP/J auxiliary basis set, DFT-D3 empirical dispersion correction with Becke–Johnson damping\textsuperscript{30,31}, COSMO water solvent model\textsuperscript{32}, and the RUCOSX approximation to speed-up the calculation of two-electron integrals\textsuperscript{33}. Simultaneous convergence tolerances of $5.0 \times 10^{-6}$ Eh for the energy change between subsequent steps, 3.0 $\times 10^{-4}$ Eh bohr$^{-1}$ for the maximum gradient, 1.0 $\times 10^{-4}$ Eh bohr$^{-1}$ for the RMS gradient, 4.0 $\times 10^{-3}$ bohr for the maximum atomic displacement, and 2.0 $\times 10^{-3}$ bohr for the RMS atomic displacement between subsequent steps, were applied.

The AMI-EM structures of the SA conformation of fraxamoside with the sir cycle in equatorial position on the macrocycle, of oleuropein, and Compound 3, were fully optimized using GAMESS program\textsuperscript{34} at the Hartree–Fock level with STO-3G basis set, followed by a single-point HF energy evaluation at the 6–31G* level to derive the partial atomic charges for the ligands by the RESP procedure\textsuperscript{35}.

**Docking**

Docking studies were performed with AutoDock 4.2\textsuperscript{36}. The crystallographic structures of XO (PDB: 3NVY) and ligands were processed with AutoDock Tools (ADT) package version 1.5.6rc1\textsuperscript{36} to merge nonpolar hydrogens, calculate Gasteiger charges, and select rotatable side-chain bonds. Grids for docking evaluation with a spacing of 0.375 Å and 40-40-60 points, centered in the ligand binding pocket, were generated using the program AutoGrid 4.2 included in Autodock 4.2 distribution. In total, 100 molecular docking runs for each “docking calculation” were performed adopting a Lamarckian Genetic Algorithm (LGA) and the following associated parameters: 100 individuals in a population with a maximum of 15 million energy evaluations and a maximum of 37 000 generations, followed by 300 iterations of Solis and Wets local search. Flexibility was used for all rotatable bonds of docked ligands and selected residue sidechain dihedrals in protein binding site, namely Leu648, Glu802, Leu873, Ser876, Phe1009, Val1011, and Leu1014. For each docking run, the representative binding poses with most favorable binding energy from the most populated clusters were selected for the subsequent MD simulations of ligand–XO complexes.

**Molecular dynamics, energy analysis, and binding free energy evaluation**

The selected complexes for each ligand were completed by addition of all hydrogen atoms and underwent EM and then MD simulations with Amber12 pmemd.cuda module\textsuperscript{37,38}, using f12SB version of AMBER force field\textsuperscript{39} for the protein and gaff parameters\textsuperscript{40} for the ligands. Parameters from Kikuchi et al.\textsuperscript{41} were used for the diothioimolbdenum(VI) ion.

To perform MD simulation in solvent, complexes were confined in TIP3P water periodic boxes exhibiting a minimum distance between solute and box surfaces of 10 Å, using the tleap module of AmberTools13\textsuperscript{42}. Each system was then neutralized by addition of counternions (Na$^+$) and underwent 1000 steps of EM with solute atoms harmonically restrained to their starting positions using a force constant of 10 kcal mol$^{-1}$ Å$^{-1}$. The resulting structures were submitted to 90 ps restrained MD (5 kcal mol$^{-1}$ Å$^{-1}$) at constant volume, gradually heating the system to 300 K, followed by 60 ps restrained MD (5 kcal mol$^{-1}$ Å$^{-1}$) at constant temperature (300 K) and pressure (1 atm) to adjust system density. Production MD simulations were carried out at constant temperature (300 K) and pressure (1 atm) for 10 ns with a time-step of 2 fs. Bonds involving hydrogens were constrained using the SHAKE algorithm\textsuperscript{43}.

Nonbonded energy contribution analysis has been performed with the cpptraj module of AmberTools13\textsuperscript{35} on the fully-solvated MD trajectories, following the minimum image convention and applying a 12 Å cutoff on both van der Waals and electrostatic interactions. For these latter a simple shifting function has been applied, to smoothly reduce the electrostatic interaction to zero at $r_{\text{cut}}$ thus avoiding step-like truncation effects at the cutoff distance:

$$E_{\text{elec}} = k \sum q_i q_j (1 - r_{ij}^2/r_{\text{cut}}^2)^2,$$

where $q_i$, $q_j$ are the charges of atoms $i$ and $j$ and the distance between them, respectively, and $k$ includes the dielectric constant ($=1$ in vacuo), the constant terms and all the unit conversion factors required by the units used for $q$, $r$, and $E$.

Gibbs binding free energies ($\Delta G_h$) have been calculated from MD trajectories with the sietraj program\textsuperscript{44,45}. They are reported as mean values ± SEM.

**Results**

**Redetermination of IC$_{50}$ in pure water for Compounds 1–4 and allopurinol**

IC$_{50}$ values on Compounds 1–4 were previously determined in 5% (v/v) methanol-water solution to allow comparisons with compounds scarcely soluble in pure water\textsuperscript{17}. However, since methanol inhibits XO, it can affect the experimental characterization of XO inhibition by other ligands\textsuperscript{46,47}. Thus, we determined the IC$_{50}$ values for Compounds 1–4 and allopurinol reported in Table 1 in the total absence of methanol (see “Methods”).

| Compound | IC$_{50}$ (µM) | $K_i$ (µM) | Inhibition mode |
|----------|----------------|------------|----------------|
| 1        | 16.1 ± 0.7     | 0.9 ± 0.25 | Competitive    |
| 2        | 335 ± 17       | 53.0$^0$   | Competitive$^0$ |
| 3        | 259 ± 13       | 37.0$^0$   | Competitive$^0$ |
| 4        | 1126 ± 22      | No inhibition$^+$ | -$^0$         |
| Allopurinol | 22.4 ± 0.2   | 1.9 ± 1.0 | Competitive    |

$^0$Data obtained from Flemmig et al.\textsuperscript{12}.

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The unusual structural features of fraxamoside as a XO inhibitor and activity patterns of its related Compounds 2–4 discussed in the introduction prompted us to experimentally determine XO inhibition kinetics and $K_i$ for fraxamoside and, as a reference, allopurinol and to compare the results with available data on Compounds 2 and 4. Kinetic analysis demonstrated that XO activity was competitively inhibited by fraxamoside. In fact, as reported in Figure 2(a), at increasing concentrations of inhibitor the value of $V_{max}$ remained unchanged, whereas $K'_m$ for xanthine progressively increased in the presence of concentrations of fraxamoside of 0.19, 0.83, and 1.90 μM (inset to Figure 2(a)). These data allowed to figure out a $K_i$ for fraxamoside of 0.9 ± 0.25 μM (Table 1), a value comparable to that measured for allopurinol under the same experimental conditions (1.9 ± 1.0 μM; Figure 2(b)). Allopurinol exhibited a competitive inhibition mode, like oleuropein, the most abundant component in Olea europea extract, widely studied for its significant antioxidant activity, as well as for many other nutraceutical properties. The $K_i$ value of fraxamoside is at least 50 times lower than that of oleuropein and comparable to that of allopurinol (Table 1).

Overall, these results show a competitive mechanism for the binding of fraxamoside, thus delimiting the region to be explored with computational techniques to the active site of the enzyme. They also parallel the relative scale of potency of the different ligands observable on IC50 values (Table 1) under the same experimental conditions adopted for $K_i$ determination.

### Molecular docking and molecular dynamics simulations

**General features**

A combined approach of molecular docking and MD simulations was undertaken to investigate the binding modes of Compounds 1–3 into the active site of XO, in order to rationalize the experimental trends observed in XO inhibition.

In all our calculation, we only employed the so-called MoCo domain (~800 residues) of XOR, hereinafter referred to as XO, which is the site of the oxidative hydroxylation of purines. The investigated molecules significantly differ in shape and size from quercetin, the ligand crystallized in complex with the selected XO structure used as target. Thus, the sidechains of some critical residues, both at the entrance of, and inside the binding site, namely Leu648, Glu802, Leu873, Ser876, Phe1009, Val1011, and Leu1014, were kept flexible in all the docking runs. The stability of all the distinct poses within the best 20 complexes by energy from each docking run was assessed by MD.
Figure 4. Representative structure of the ligand binding site of the final model of the XO complex with fraxamoside (Compound 1). The site of the representative frame from the last 5 ns of MD for the XO-fraxamoside complex III model is shown adopting a partially transparent tan ribbon representation for protein backbone, and sticks for ligand, prosthetic groups, and protein sidechains of residues involved in contacts with either ligands or dioxothiomolybdenum(VI) ions. Atoms are colored with the following scheme: O = red, N = blue, H = white, S = yellow, P = orange, Mo = light-blue, cyan = C atoms from residues involved in conserved interactions with all structurally characterized XO ligands, solid tan = C atoms from residues exhibiting specific interactions with fraxamoside and magenta = C atoms in fraxamoside. Ligand-protein H-bonds are depicted with a green spring.

In principle, the fraxamoside macrocycle can assume two major conformations, with the ethereal linkage between the C1 of \( \beta gl \) and the C1 of the sir cycle in either axial or equatorial position. SA calculations selected one main conformer for each macrocycle conformation. These two structures were optimized at DFT(B3LYP)/Def2-TZVPP level in implicit water. Since the equatorial form is by far (19.7 kcal mol\(^{-1}\)) more stable than the axial conformer, only the former was considered in docking calculations.

**XO-fraxamoside (Compound 1) complex**

The analysis of docking results led to the selection of four XO-fraxamoside\(^1\) complex models for the subsequent MD simulation step. The four poses differ in the portion of molecule that docks into the XO active site region: (i) hty, (ii) \( \beta gl \), (iii) both of them, (iv) none of them. MD trajectory analysis showed that complex I was fairly unstable, as shown by a positional RMSD time plot of ligand heavy atoms after protein backbone atom best fit (Figure 3(a)), due to the loss of the starting \( \pi \) stacking between hty and Phe914, and perpendicular \( \pi \)–\( \pi \) interaction with Phe1009.

During MD simulation of complex II, fraxamoside\(^1\) binding mode underwent an initial transition, associated to a sudden increase to about 3 Å of the RMSD value shown in Figure 3(a), determined by a ligand rotation with loss of the initial H-bonds between sir carboxylic O11 and Lys771 and between \( \beta gl \) O6' and Thr1010, whereas the H-bonds of \( \beta gl \) O2'/3' with Glu802 and hty O5'/6' with Glu879 persist during the whole simulation.

Complex III was quite stable during MD (Figure 3(a)), and preserved all its starting interactions: the hty pendant is sandwiched between Phe914 and Phe1009, forming \( \pi \) stacking with the former and a \( \pi \)–\( \pi \) perpendicular interaction with the latter, while O6' engages a H-bond with the oxygen atom of the dioxothiomolybdenum(VI) ion. \( \beta gl \) O3'/4' are H-bonded to Glu802, and the sir ethylidene group forms hydrophobic interactions with Leu648 (Figure 4), all other protein–ligand interactions being shown in Figure 5.

Instead complex IV resulted unstable, drifting toward dissociation (Figure 3(a)) during the simulation. According to calculated \( \Delta G_0 \) values, complex III is more stable than II by 1.5 kcal mol\(^{-1}\) (9.33 ± 0.02 kcal mol\(^{-1}\) versus 7.76 ± 0.02 kcal mol\(^{-1}\), respectively), thus for XO-fraxamoside only model III will be discussed hereafter.

The relative contributions of the groups in which fraxamoside can be ideally fragmented to the ligand nonbonded interaction energy with XO and the solvent, calculated with the shifting function for the electrostatic part described in "Methods", show that hty is responsible for 29.0% (29.0% with XO, 0% with solvent) of the total fraxamoside–environment interaction, while sir only contributes 15.8% (10.7% XO, 5.1% solvent) and the \( \beta gl \) is the most strongly interacting group in this binding mode, accounting for 55.2% (50.8% XO, 4.4% solvent) of the total.

**XO-oleuropein (Compound 2) complex**

The classification adopted for the poses of fraxamoside can be also applied to the selected docking models for XO-oleuropein (Compound 2) complex. However, this latter exhibits additional poses corresponding to minor variants of the II and III orientations, arising from the availability in oleuropein of the primary (O6') hydroxyl group of \( \beta gl \). In fact, complex II has three variants, with O6' alternatively pointing toward either the dioxothiomolybdenum(VI) ion (II-1), or Glu802 (II-2) or Thr1010 (III-3). However, only complex II-3 was stable throughout the MD simulation (Figure 3(b)), stabilized by the H-bonds between \( \beta gl \) O2'/3' and Glu802, \( \beta gl \) O6' and Thr1010, hty O5' and Glu879, and hty O6' and Thr1010 carbonyl backbone (Figure 6(b)).

Complex III, corresponding to the most stable XO-fraxamoside pose, has two variants, characterized by different orientations of \( \beta gl \) O6', facing either hty C2' (III-1), thus mimicking fraxamoside ring closure, or Lys771 sidechain (III-2). Additionally, to ensure the most complete comparison with the favorite binding mode of fraxamoside, we also built and simulated by MD a XO-oleuropein complex model (III") directly derived from the III model of the XO-fraxamoside complex by ring opening followed by local EM of the resulting structure of oleuropein. All III complexes deeply rearranged during MD simulations, losing the initial stacking interaction between hty and Phe914. The final position of oleuropein in all III complexes was shifted toward the rim of the cleft: in III-1 \( \beta gl \) retained the initial H-bond interactions with Glu802, but hty formed intramolecular H-bonds with \( \beta gl \) rather than the protein, in III-2 \( \beta gl \) drifted toward the external rim replacing its initial H-bonds to Glu802 with H-bonds to Lys771 and Asp872, this latter being outside the cleft, and complex III" rearranged toward a structure essentially overlapping that observed in II-3 MD simulation.

Complex IV, featuring neither \( \beta gl \), nor hty inside the pocket, was fairly unstable during the simulation period, the ligand drifting toward the external rim of the catalytic pocket as shown by ligand RMSD plots in Figure 3(b). Complex I, with hty inside and \( \beta gl \) at the pocket entrance, remained stable after an initial rearrangement (Figure 6(a)). Oleuropein formed six H-bonds with the protein: two double interaction between hty
O5’/O6’ and Arg880, \( \beta_{gl} \) O2’/3’ and Glu879, and two single H-bonds, between \( hty \) O6’ and, alternatively, one of the O or the S atom of the dioxothiomolybdenum(VI), \( \beta_{gl} \) carbonylic O11, and Ser876. Moreover, oleuropein is involved in \( \pi-\pi \) stacking and perpendicular \( \pi-\pi \) interactions of its \( hty \) group with Phe914 and Phe1009, respectively, and in the van der Waals interactions shown in Figure 5. Since calculated \( \Delta G_0 \) predicts this latter more stable than \( II-3 \) by 0.7 kcal mol\(^{-1}\) (\(-8.80 \pm 0.02 \) kcal mol\(^{-1}\) versus \(-8.06 \pm 0.02 \) kcal mol\(^{-1}\), respectively) only model \( I \) will be discussed hereafter for XO-oleuropein.

The analysis of the relative contributions of the different fragments of oleuropein to the ligand nonbonded interaction energy with the environment provided the following contributions for complex \( k \): 37.8% (35.3% with XO, 2.5% with solvent) for \( hty \), 18.5% (16.9% with XO, 1.6% with solvent) for \( \beta_{gl} \), 43.7% (37.9% with XO, 5.8% with solvent) for \( \beta_{gl} \).
our models of XO complexes with Compounds 1–3. The site of the representative frame from the last 5 ns of MD for the XO-Compound 3 complex II-2 is shown with the same representation and colors used in Figure 4, except for C atoms of Compound 3, in dark blue.

**Figure 7.** Representative structure of the ligand binding site of the final model of the XO complex with oleoside 11-methyl ester (Compound 3). The site of the representative frame from the last 5 ns of MD for the XO-Compound 3 complex II-2 is shown with the same representation and colors used in Figure 4, except for C atoms of Compound 3, in dark blue.

**XO-oleoside 11-methyl ester (Compound 3) complex**
The selected XO-oleoside 11-methyl ester (Compound 3) complexes from docking runs exhibited the same three orientations of the glycosidic ring inside the pocket found in XO-oleuropein, with the primary (6') hydroxyl group of βgl alternatively pointing toward the dioxothiomolybdenum(VI) ion (II-1), Glu802 (II-2), or Thr1010 (II-3).

Complex II-1 rearranged during MD, losing the initial H-bond with dioxothiomolybdenum(VI) ion and forming a pattern of H-bonds with βgl similar to that observed for complex II-2 (shown in Figure 5), whereas the sir carboxylate group is now involved in an intramolecular H-bond.

Complex II-2 was quite stable along MD trajectory (Figure 3(c)): βgl formed a network of H-bonds involving O6’ with Glu802, and O4’ with Arg 880/Thr1010 and the sir carboxylate group (O7 atoms) was H-bonded/salt bridged to Lys771 (Figure 7). The network of intermolecular H-bonds/salt bridge and the other protein–ligand interaction identified in the MD trajectories of this complex are shown in Figure 5.

Complex II-3, even if stably resident within the site throughout the whole MD simulation, featured a reduced number of stabilizing H-bond interactions, since it lose the initial H-bonds/salt bridge between sir carboxylate of Compound 3 and Lys771.

**Discussion**

**Comparison with other known XO ligands**
Experimentally solved complex structures of XO involve ligands (quercetin, PDB: 3NVY; Febuxostat, PDB: 1N5X; trihydroxy-FYX-051, PDB: 3AM9; oxypurinol, PDB: 4BDJ; Y-700, PDB: 1VDV; FYX-051, PDB 1V97) exhibiting widely different binding modes, suggested mechanisms for XO inhibition and potency. In spite of this, a comparison of their short-range contacts with XO shows a conserved subset of interactions shared by all the solved ligands (marked with an asterisk (*) in the first column of Figure 5). When our models of XO complexes with Compounds 1–3 are added to this comparison, it can be easily observed that the subset is substantially preserved (stick representation with cyan C atoms in Figures 4, 6, and 7), with the only exception of Compound 3, which does not form direct interactions with the dioxothiomolybdenum(VI) ion. Another noticeable result is that the three ligands also form most of the contacts included in the wider subsets containing interactions detected in all but either oxypurinol or quercetin PDB ligands (marked with a degree (°) and a caret (^) symbol, respectively, in the first column of Figure 5). In particular, fraxamoside forms all the contacts from these two subsets. While number and exact nature of each interaction in the subset may vary between different ligands, the involved protein residues appear to play a fundamental role in the stabilization of the experimental XO–ligand complexes. In this view, this analysis provides a validation of the proposed models for the complexes of XO with Compounds 1–3 as competent structures for XO inhibition. Furthermore, the three ligands may also give rise to additional interactions, not observed, or corresponding to very loose contacts, in experimental complexes (residues in parenthesis in the first column of Figure 5, stick representation with tan C atoms in Figures 4, 6, and 7). Drug design/lead optimization based on a combination of exploitation of these new potential interaction sites with the possible strengthening of those interactions which in Compounds 1–3 are weaker than in strong XO ligands could represent an effective strategy to improve ligand potency, while retaining their unique structure–activity relationships.

**Structure–activity relationships among the investigated inhibitors**
The complex models also answer the more specific questions arisen in the “Introduction” about the inhibitory properties of Compounds 1–3 on XO. First, they justify the XO inhibition activity exhibited by the most potent compound of the series, fraxamoside, in spite of its size, bulkiness, and the overall structural properties rather unusual for a XO ligand. Second, they explain the non-additivity observed in the XO inhibitory activity within the compound 1–4 series (Table 1), and, in particular, the origin of the higher activity exhibited by fraxamoside versus oleuropein. Third, they explain the stabilizing effect of the sugar unit on the complexes, differently from what observed for other natural products containing phenols groups, whose glycosides are usually less active than the corresponding aglycones.

As for the first point, fraxamoside can accommodate both the βgl and the hty catechol rings deeply inside the XO binding pocket, by engaging a network of stabilizing interactions with both dioxothiomolybdenum(VI) ion, and those XO residues identified in the crystal structures of other XO-inhibitor complexes as critical for ligand binding, such as Glu802, Phe914, and Phe1009, thus reaching a binding mode that is both tight and effective in precluding the access of XO substrates to the active site.

As for the second point, this docking mode appears exclusive to fraxamoside, since it critically depends on the conformational properties of the fraxamoside macrocycle. In fact, oleuropein, in spite of its considerably larger flexibility, appears unable to simultaneously dock in a stable fashion its βgl and hty catechol rings deep inside XO active site. Even those starting poses of oleuropein exhibiting a ligand conformation and relative orientation to XO site very similar to those occurring in XO-fraxamoside III model rapidly lose the interaction of either of the two rings with residues critical for binding to XO during the MD simulations, thus resulting in different binding modes between fraxamoside and oleuropein.
In principle, it would be expected that the higher flexibility of oleuropein on the one hand may allow a better optimization of ligand–protein interactions, in turn leading to complexes enthalpically favored over those accessible to the conformationally restricted fraxamoside, but on the other hand would determine a higher entropic cost due to the higher loss of ligand conformational freedom upon binding. However, even by employing a method to estimate the binding free energy that totally neglects this latter entropic contribution (see “Methods”), the XO-fraxamoside complex is still favored by about 0.5 kcal mol\(^{-1}\) over XO-oleuropein. In this view, oleuropein, rather than “led toward a more favorable binding mode by its flexibility”, appears to be “prevented from assuming exactly the same optimal conformation observed in best XO-fraxamoside models by the sterical bump between βgl C6′ and O6′ and hty C2′ atoms”, which are instead covalently bonded in fraxamoside. The combination of different predicted binding modes and energy contributions between these two ligands suggests that the optimal conformation for binding lies in the quite narrow region of conformational subspace only accessible to the cyclic molecule and not to its linear counterpart, rather than either in their common subspace, which would provide similar binding modes with an exclusively entropic advantage for the cyclic compound over its open form, or in the much wider subspace only accessible to the linear molecule, which would result in different binding modes, but associated to a stronger predicted ΔG\(_{\text{b}}\) (at least when neglecting conformational entropy changes) for the linear compound. The unbalance between energetic gain and “entropic cost” deriving from oleuropein flexibility is also confirmed by an extension of the comparison to Compound 3, a “fragment” of oleuropein devoid of hty, from which oleuropein emerges as the worst inhibitor in the series. Compound 3, although lacking the aromatic moiety, critical for π–π stacking with key aromatic residues in the catalytic site, is able to compensate this gap with, on one hand, a lower reduction of conformational freedom upon binding, and on the other hand, the occurrence of an ionic interaction with Lys771, which helps to stably anchor the molecule in the active site.

As for the third point, the analysis of all the XO-ligand models obtained in the present paper suggests an active stabilizing contribution of βgl to binding, in sharp contrast with glycoside derivatives of other polyphenols active on XO, usually detrimental to the activity of the corresponding aglycone\(^{30-31}\). For instance, a comparison between the XO inhibitory effect of quercetin and its 3-O-glucoside derivative\(^{37}\) (IC\(_{50}\) = 1.9 and 54.6 μM, respectively) clearly demonstrates a detrimental effect of glycosilation for its inhibitory activity. A comparative analysis between the X-ray structure of quercetin-XO and a model of quercetin 3-O-glucoside-XO complex suggests that this effect may derive from the limited flexibility of the flavonoid system, which produces a change in the exact binding mode of the flavonoid group to host the glucose unit in the XO site (data not shown), thus impairing the network of favorable interactions of the aglycone without a sufficient compensation from the new interactions involving the glycosidic unit. Instead, in our models of all the XO–ligands complexes, βgl substantially contributes to optimal XO binding, thus compensating the relatively small contribution of hty to the overall binding observed in fraxamoside and oleuropein complexes. The relative weight of the overall contribution of hty (29.0%) versus βgl (55.2%) in the best predicted XO binding mode for fraxamoside (model III) is consistent with both experimental data (poor activity of Compound 4 and comparable activity of Compounds 2 and 3\(^{17}\), and other computational results obtained in our MD simulations, such as the aforementioned observed instability of the XO-fraxamoside binding mode I, and labilization of the interaction between hty and XO active site in all the models of XO-oleuropein belonging to the III family. Even in complex I of oleuropein, which exhibits the largest contribution of hty to the total interaction energy of the ligand with its environment (37.8%), the largest term is still provided by βgl (43.7%). While in fraxamoside and oleuropein, the sugar group contributes by both forming direct interaction with the middle-external region of XO active site and improving and protecting the interactions formed in the inner part of the XO active site by hty and βgl, Compound 3, which lacks this latter group, is mainly stabilized by direct interactions of βgl with the middle-internal region of XO active site, where it also forms van der Waals contacts with the important aromatic residues Phe914 and Phe1009. Thus, at present, a large stabilizing contribution from a sugar unit represents, although with different mechanism between Compounds 1/2 and 3, a peculiarity of the ligands characterized in the present study, which could be exploited to design new derivatives of these molecules endowed with improved ligand pharmacokinetic or tolerability properties.

The relative contribution of hty to the interaction of the ligands with XO, although limited in comparison with hty and βgl (10.7% and 16.9% for fraxamoside and oleuropein, respectively) is important to ensure the proper orientation of hty or βgl and provides one of the hydrophobic anchoring groups of the three complexes. After cyclization in fraxamoside, it assumes a conformation capable of simultaneously optimizing hty and βgl interactions with XO.

Conclusions

Fraxamoside (Compound 1) represents a promising lead compound for a new family of XO inhibitors, characterized by peculiar structural features, and particularly, by the exploitation of different binding groups instead of the combination of aromatic/conjugated rings typical of other XO inhibitors. In particular, this study underlines the importance of the conformational properties of the fraxamoside macromolecule. In fact, this ring, quite rigid despite its size, both allows an optimal orientation inside the XO active site of all the “subligands” in which fraxamoside can be ideally split, and, thanks to its rigidity, minimizes the entropic cost associated to conformational freezing upon binding.

Furthermore, the comparison of the modeled static and dynamic properties between the XO complexes of Compounds 1–3 has allowed a rationalization of their relative scale of activity. The availability of a small but “well-targeted” natural library of related compounds in the Olea europaea L. leaf decoction represented a precious contribution to characterize the SAR of the most active compound in the series.

Another relevant implication of the SARs derived for Compounds 1–4 is their strong non-additivity, which suggests a great care in the systematic application of simple additive models in the derivation of SARs or in molecular design.

Overall, the last two observations pointed out the importance of the combination of richness and “smooth variability” of most natural sources, exhibiting a substantially higher complexity and diversity of scaffolds than synthetic libraries featuring several orders of magnitude more compounds, but usually based on additive criteria in their design.

However, the quality of a virtual screening approach depends more critically on computational protocols for binding mode screening, optimization, and analysis when applied to a relatively small and widely structurally variable natural product library than to a more smoothly variable synthetic (especially combinatorial) library, since the latter usually benefits of both a fairly larger
statistical base, and of cancelations of errors on energy/scoring terms often occurring within a similar series of compounds. Thus, more reliable validations of the proposed binding modes for the former are required, also allowing for extensive rearrangements of the models deriving from docking, possibly at a cost in terms of required computational times, largely compensated by the lower number of compounds to be screened. Our docking + MD-based protocol proved effective in discriminating between stable and unstable binding modes and allowed from moderate to substantial rearrangements of the starting models and a careful comparison of potentially similar binding modes between different ligands.

The ensemble of structure–activity relationships derived from the overall computational and biochemical data is expected to significantly contribute to the identification and design of highly selective ligands toward XO, with small or no affinity for other targets involved in the metabolism of purines and their derivatives. In this view, a relevant feature is represented by the possibility of obtaining fraxamoside and its analogs, as well as building blocks, from a relatively inexpensive source, the leaves of *Olea europaea* L., i.e. a by-product of the olive cultivation.

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