Protein Targets of Frankincense: A Reverse Docking Analysis of Terpenoids from *Boswellia* Oleo-Gum Resins

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Abstract: **Background:** Frankincense, the oleo-gum resin of *Boswellia* trees, has been used in traditional medicine since ancient times. Frankincense has been used to treat wounds and skin infections, inflammatory diseases, dementia, and various other conditions. However, in many cases, the biomolecular targets for frankincense components are not well established. **Methods:** In this work, we have carried out a reverse docking study of *Boswellia* diterpenoids and triterpenoids with a library of 16034 potential druggable target proteins. **Results:** *Boswellia* diterpenoids showed selective docking to acetylcholinesterase, several bacterial target proteins, and HIV-1 reverse transcriptase. *Boswellia* triterpenoids targeted the cancer-relevant proteins (poly(ADP-ribose) polymerase-1, tankyrase, and folate receptor β), inflammation-relevant proteins (phospholipase A2, epoxide hydrolase, and fibroblast collagenase), and the diabetes target 11β-hydroxysteroid dehydrogenase. **Conclusions:** The preferential docking of *Boswellia* terpenoids is consistent with the traditional uses and the established biological activities of frankincense.

Keywords: frankincense; *Boswellia*; cembranoids; cneorubenoids; boswellic acids; molecular docking

1. Introduction

The genus *Boswellia* (Burseraceae) is made up of resiniferous trees and shrubs that are distributed across India, the Arabian peninsula, and Africa [1,2]. The genus is known for its aromatic terpenoid oleo-gum resin, frankincense. Frankincense has been a part of human religious ceremonies and ethnomedicine for thousands of years [3]. Important frankincense-producing species include *B. carteri*, which grows in Somaliland and Puntland [1], *B. sacra*, found in Yemen, southern Oman, Somalia, and Somaliland [2], *B. frereana*, which is endemic to Somalia [2], *B. papyrifera*, primarily found in Sudan, Eritrea, and Ethiopia [4], and *B. serrata*, which grows primarily in India [5].

Frankincense oleo-gum resin has been used traditionally to treat wounds [6], to treat inflammatory diseases [7], for oral hygiene [8], as well as for its psychoactive effects [9,10]. The biological activities of frankincense have been attributed to its essential oils [11] and its non-volatile diterpenoids and triterpenoids [6]. Although frankincense has been used for various maladies and conditions, and numerous biological activities have been attributed to frankincense, the particular biological targets are not well established. In this work, we have carried out a reverse molecular docking study of *Boswellia* cembranoid diterpenoids (Figure 1), cneorubenoid diterpenoids (Figure 2), and triterpenoids (Figure 3) against a library of 16,034 potential druggable target proteins.

The cembranoids incensole and incensole acetate were detected in the oleo-gum resin of *B. papyrifera*, while serratol was found in *B. carteri*, *B. sacra*, and *B. serrata* [12]. The boscartins have been
isolated from the oleo-gum resin of *B. carteri* [13]. Incensole oxide has been isolated from *B. carteri* and the X-ray crystal structure determined [13,14]; both incensole oxide and incensole oxide acetate have been detected in small concentrations in the essential oil from the resin of *B. papyrifera* [15]. Isoincensole oxide [16,17] and isoincensolol [17] were isolated from *B. carteri* resin. Verticilla-4(20),7,11-triene and serratol and were isolated from *B. carteri* [18] and *B. serrata* [19], respectively.

![Macrocyclic diterpenoids found in Boswellia species.](image)
Boswellia carteri oleo-gum resin is the source of several prenylated aromadendrane (cneorubenoid) diterpenoids (Figure 2) [20,21]. Numerous ursane, oleanane, lupane, dammarane, and tirucallane triterpenoids have been isolated and characterized from Boswellia species (Figure 3) [22]. Boswellia serrata has yielded α-boswellic acid, β-boswellic acid, 3-acetyl-α-boswellic acid, 3-acetyl-β-boswellic acid, 11-keto-β-boswellic acid, and 3-acetyl-11-keto-β-boswellic acid [23]. Boswellia carteri has yielded the oleanane triterpenoids α-boswellic acid, and 3-acetyl-α-boswellic acid; the ursane triterpenoids β-boswellic acid, 3-acetyl-β-boswellic acid, 11-keto-β-boswellic acid, 3-acetyl-11-keto-β-boswellic acid, 3-acetyl-11α-methoxy-β-boswellic acid, 9,11-dehydro-β-boswellic acid, and 3-acetyl-9,11-dehydro-β-boswellic acid; the lupane triterpenoids lupeolic acid and 3-acetyl lupeolic acid; and the tirucallane triterpenoids α-elemolic acid, β-elemonic acid, 3α-hydroxytirucalla-7,24-dien-21-oic acid, 3α-acetoxytirucalla-7,24-dien-21-oic acid, and 3β-hydroxytirucalla-7,24-dien-21-oic acid [24]. Olibanumols A, B, C, H, I, J’ [25], E, F, G [20], K, L’, M, and N [26] have been isolated from the oleo-gum resin of B. carteri. B. carteri resin has also yielded boscartenes L, M, and N, as well as trametenolic acid B, 3-oxotirucalla-7,9(11),24-trien-21-oic acid, and (20S)-3,7-dioxo-tirucalla-8,24-dien-21-oic acid [27].
Figure 3. Cont.
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Figure 3. Triterpenoids isolated from *Boswellia* species.
2. Materials and Methods

2.1. Ligand Preparation

Each ligand structure was prepared using Spartan’16 v. 2.0.7 (Wavefunction, Inc., Irvine, CA, USA). The lowest-energy conformations of the ligands were determined using the Merck Molecular Force Field (MMFF) [28]. In the case of the cembranoid macrocyclic ligands, further conformational analysis was carried out using density functional theory at the M06-2X/6-31G* level [29] with SM8 [30] aqueous solvent model [31].

2.2. Reverse Molecular Docking

A reverse molecular docking study was carried out on each of the Bosellia terpenoids with the sc-PDB database of druggable binding sites [32]. Each compound was examined against the 16034 protein targets contained in the sc-PDB database. Prior to docking, all solvent molecules were removed from the protein structures. Co-crystallized enzyme cofactors were retained as cofactors and co-crystallized substrates or inhibitors were retained as ligands. Molecular docking was carried out using Molegro Virtual Docker v. 6.0.1 (Molegro ApS, Aarhus, Denmark) [33] as previously reported [34]. A python script was written to generate the Molegro input files; the jobs were run as a batch from the mvd.exe command line executable. The script took the co-crystallized ligands in each protein and wrote an input file that defined the search space for that docking as a sphere centered on the ligand’s center of mass. A 15-Å radius sphere was centered on the binding sites of each protein structure in order to permit each ligand to search. Standard protonation states of each protein, based on neutral pH, were used and charges were assigned based on standard templates as part of the Molegro Virtual Docker program. Each protein was used as a rigid model without protein relaxation. Flexible-ligand models were used in the docking optimizations. Different orientations of the ligands were search and ranked based on their “rerank” energy scores. A total of 100 runs for each ligand were carried out.

2.3. Conformational Analysis of Boscartol D

All calculations were carried out using Spartan’16 for Windows (Wavefunction, Inc., Irvine, CA, USA). Conformational profiles were carried out using molecular mechanics with the MMFF force field. Conformations with relative energies <20 kJ/mol were re-evaluated, with geometry optimization, using density functional theory (DFT, M06-2X/6-31G*) with a nonpolar solvent (CHCl$_3$) model.

3. Results and Discussion

3.1. Cembranoid Diterpenoids

The macrocyclic cembranoid diterpenoids examined in this study are shown in Figure 1. The top-binding protein targets for each of the Boswellia cembranoids are summarized in Table 1. Included in Table 1 are the median docking energies for comparison. The top binding proteins for boscartin A were acetylcholinesterase (AChE) enzymes, Torpedo californica (TcAChE) and human (HsAChE). Boscartin B docked preferentially with human N-acetylgalactosaminyltransferase (HsGTA) as well as with the bacterial targets Serratia marcescens chitinase B (SmChiB), Helicobacter pylori peptide deformylase (HpPDF), and Mycobacterium tuberculosis 7,8-diaminopelargonic acid synthase (MtBioA). The proteins with the most exothermic docking for boscartin C were Escherichia coli aspartate transaminase (EcAspTA), murine acetylcholinesterase (MmAChE), and Daboia russelii (Russell’s viper) phospholipase A$_2$ (DrPLA2). Boscartin D showed excellent docking with TcAChE (PDB 2cek, $E_{\text{dock}} = -115.3$ kJ/mol) and EcAspTA. The best protein targets for boscartin E were TcAChE, HpPDF, and MtBioA. Boscartin F showed preferential docking energies with human pyruvate kinase M2 (HsPKM2), TcAChE, and HpPDF. Boscartin G showed excellent docking properties with acetylcholinesterases TcAChE and HsAChE. The proteins with the most exothermic docking
energies with boscartin H were human N-acetylgalactosaminyltransferase (HsGTA), DrPLA2, SmChiB, and MmAChE.

| Ligand       | PDB  | $E_\text{dock}$ (kJ/mol) | Target Protein                                                                 |
|--------------|------|--------------------------|-------------------------------------------------------------------------------|
| Boscartin A  | 1e66 | −119.7                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|              | 2cek | −123.7                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|              | 4bdt | −113.5                   | human acetylcholinesterase (HsAChE)                                           |
|              |      |                          | Median docking energy                                                         |
| Boscartin B  | 1h0g | −107.8                   | Serratia marcescens chitinase B (SmChiB)                                       |
|              | 2ew5 | −104.6                   | Helicobacter pylori peptide deformylase (HpPDF)                                |
|              | 3fu  | −107.5                   | Mycobacterium tuberculosis 7,8-diaminopelargonic acid synthase (MtBioA)         |
|              | 3vlo | −110.4                   | human fucosylgalactoside α-N-acetylgalactosaminyltransferase (HsGTA)            |
|              |      |                          | Median docking energy                                                         |
| Boscartin C  | 1ahg | −112.5                   | Escherichia coli aspartate aminotransferase (EcAspTA)                         |
|              | 1fv0 | −108.2                   | Daboia russelli (Russell’s viper) phospholipase A2 (DrPLA2)                    |
|              | 1q83 | −110.4                   | murine acetylcholinesterase (MmAChE)                                          |
|              | 4g1n | −106.5                   | human pyruvate kinase isozyme M2 (HsPKM2)                                     |
|              |      |                          | Median docking energy                                                         |
| Boscartin D  | 1ahg | −114.4                   | Escherichia coli aspartate aminotransferase (EcAspTA)                         |
|              | 1xzd | −106.3                   | Thermotoga maritima GTP-binding protein TrmE (TrmTrmE)                         |
|              | 2cek | −115.3                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|              |      |                          | Median docking energy                                                         |
| Boscartin E  | 1ahg | −110.4                   | Escherichia coli aspartate aminotransferase (EcAspTA)                         |
|              | 1e66 | −111.7                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|              | 2ew5 | −107.5                   | Helicobacter pylori peptide deformylase (HpPDF)                                |
|              |      |                          | Median docking energy                                                         |
| Boscartin F  | 2ew5 | −109.5                   | Helicobacter pylori peptide deformylase (HpPDF)                                |
|              | 3ifm | −107.3                   | Helicobacter pylori peptide deformylase (HpPDF)                                |
|              | 4g1n | −108.5                   | human pyruvate kinase isozyme M2 (HsPKM2)                                     |
|              |      |                          | Median docking energy                                                         |
| Boscartin G  | 1e66 | −126.4                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|              | 2cek | −116.8                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|              | 4bdt | −118.7                   | human acetylcholinesterase (HsAChE)                                           |
|              |      |                          | Median docking energy                                                         |
| Boscartin H  | 1fv0 | −106.9                   | Daboia russelli (Russell’s viper) phospholipase A2 (DrPLA2)                    |
|              | 1w1t | −106.0                   | Serratia marcescens chitinase B (SmChiB)                                       |
|              | 2gyw | −105.9                   | murine acetylcholinesterase (MmAChE)                                          |
|              | 3vlo | −109.3                   | human fucosylgalactoside α-N-acetylgalactosaminyltransferase (HsGTA)            |
|              |      |                          | Median docking energy                                                         |
| Incensole    | 2cek | −111.9                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|              | 3ug  | −109.7                   | human aldo-keto reductase 1C3 (HsAKR1C3)                                      |
|              | 1h0g | −102.4                   | Serratia marcescens chitinase B (SmChiB)                                       |
|              |      |                          | Median docking energy                                                         |
| Incensole acetate | 1ahg | −106.9                   | Escherichia coli aspartate aminotransferase (EcAspTA)                         |
|               | 3jun | −103.7                   | Burkholderia cepacia phenazine biosynthesis protein A/B (BcPhzA/B)            |
|               |      |                          | Median docking energy                                                         |
| Incensole oxide | 1ahg | −108.6                   | Escherichia coli aspartate aminotransferase (EcAspTA)                         |
|               | 2cek | −103.1                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|               | 3up  | −103.5                   | Burkholderia cepacia phenazine biosynthesis protein A/B (BcPhzA/B)            |
|               | 3mee | −106.7                   | HIV-1 reverse transcriptase (HIV-1 RT)                                         |
|               |      |                          | Median docking energy                                                         |
| Incensole oxide acetate | 1ahg | −119.0                   | Escherichia coli aspartate aminotransferase (EcAspTA)                         |
|               | 1q83 | −109.6                   | murine acetylcholinesterase (MmAChE)                                          |
|               | 2cek | −110.4                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|               | 3ifm | −114.3                   | Torpedo californica acetylcholinesterase (TcAChE)                             |
|               | 3mee | −113.5                   | HIV-1 reverse transcriptase (HIV-1 RT)                                         |
|               |      |                          | Median docking energy                                                         |
| Isoincensole oxide | 3up  | −103.3                   | Burkholderia cepacia phenazine biosynthesis protein A/B (BcPhzA/B)            |
|               |      |                          | Median docking energy                                                         |
Incensole docked well with TcAChE, human aldo-keto reductase 1C3 (HsAKR1C3), and SmChiB. Incensole acetate preferentially targeted bacterial proteins EcAspTA, HpPDF, and HIV-1-RT. Isoincensole oxide docked well to BcPhzA/B and TcAChE. The protein targets that showed the best docking energies with isoicensole oxide were DrPLA2, Staphylococcus aureus multidrug binding protein QacR (SaQacR), and human dehydroepiandrosterone sulfotransferase (HsSULT2A1).

Every cembranoid ligand showed excellent docking properties to acetylcholinesterases (Table 2). Acetylcholinesterase. MmAChE = human acetylcholinesterase.

The lowest-energy docked pose of boscartin G with TcAChE (PDB 1e66, Figure 4A) shows the ligand to adopt the lowest-energy conformation as calculated by density functional theory at the

Table 1. Cont.

| Ligand        | PDB a | E_{dock} | Target Protein                                                                 |
|---------------|-------|----------|---------------------------------------------------------------------------------|
| Isoincensole  | 1fv0  | −108.2   | Daboia russelli (Russell’s viper) phospholipase A₂ (DrPLA2)                      |
|               | 1jus  | −107.9   | *Staphylococcus aureus* multidrug binding protein (SaQacR)                      |
|               | 2gp4  | −102.2   | human dehydroepiandrosterone sulfotransferase (HsSULT2A1)                       |
|               |       | −74.0    | Median docking energy                                                           |
| Serratol      | 1e66  | −104.1   | *Torpedo californica* acetylcholinesterase (TcAChE)                            |
|               | 2cek  | −106.1   | *Torpedo californica* acetylcholinesterase (TcAChE)                            |
|               | 2ow5  | −103.2   | *Helicobacter pylori* peptide deformylase (HpPDF)                               |
|               | 4bdt  | −103.0   | human acetylcholinesterase (HsAChE)                                            |
|               |       | −78.1    | Median docking energy                                                           |
| Verticillatriene | 1w4l  | −104.3   | *Torpedo californica* acetylcholinesterase (TcAChE)                            |
|               | 3ifm  | −100.8   | *Torpedo californica* acetylcholinesterase (TcAChE)                            |
|               | 3izc  | −103.2   | *Torpedo californica* acetylcholinesterase (TcAChE)                            |
|               |       | −63.6    | Median docking energy                                                           |

PDB: Protein Data Bank code.

Table 2. MolDock molecular docking energies (kJ/mol) of *Boswellia* cembranoids with acetylcholinesterase protein targets. a

| Ligand       | TcAChE 1e66 | TcAChE 1h22 | TcAChE 1w4l | TcAChE 2cek | TcAChE 3ifm | TcAChE 3izc | TcAChE 1q83 | TcAChE 2gyw | TcAChE 4bdt |
|--------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|
| Boscarkin A  | −119.7      | −98.4       | −99.5       | −123.7      | −98.2       | −100.6      | −95.9       | −101.4      | −113.5      |
| Boscarkin B  | −95.1       | −98.8       | −92.5       | −106.3      | −99.2       | −94.5       | −94.3       | −85.8       | −106.3      |
| Boscarkin C  | −87.7       | −90.8       | −94.7       | −96.9       | −104.1      | −94.2       | −110.4      | −94.0       | −86.6       |
| Boscarkin D  | −99.7       | −102.8      | −93.1       | −115.3      | −103.7      | −93.9       | −92.5       | −93.6       | −90.2       |
| Boscarkin E  | −111.7      | −96.5       | −94.5       | −108.7      | −95.5       | −92.9       | −86.0       | −94.8       | −90.6       |
| Boscarkin F  | −93.2       | −97.0       | −98.5       | −104.8      | −103.7      | −95.8       | −101.1      | −82.4       | −77.3       |
| Boscarkin G  | −126.4      | −99.4       | −94.5       | −116.8      | −101.5      | −95.6       | −111.1      | −94.1       | −118.7      |
| Boscarkin H  | −100.1      | −106.3      | −105.6      | −103.0      | −104.2      | −98.8       | −105.9      | −97.9       | −95.8       |
| Incensole    | −89.7       | −92.1       | −89.5       | −111.9      | −94.1       | −89.6       | −90.3       | −87.7       | −53.4       |
| Incensole acetate | −88.6      | −97.2       | −102.8      | −102.5      | −102.3      | −101.7      | −87.5       | −85.0       | −42.3       |
| Incensole oxide | −89.6      | −93.2       | −92.7       | −103.1      | −96.9       | −89.3       | −101.9      | −95.5       | −93.5       |
| Incensole oxide acetate | −109.8     | −98.7       | −96.8       | −110.4      | −114.3      | −95.8       | −109.6      | −97.7       | −102.6      |
| Isoicensole oxide | −92.9       | −98.9       | −96.4       | −103.6      | −98.1       | −89.9       | −89.9       | −90.3       | −94.1       |
| Isoicensoleolol | −88.5       | −99.5       | −92.4       | −101.5      | −98.3       | −91.2       | −77.0       | −80.4       | −73.6       |
| Serratol     | −104.1      | −95.7       | −92.2       | −106.1      | −94.5       | −88.1       | −96.4       | −89.3       | −104.0      |

*TcAChE = *Torpedo californica* acetylcholinesterase. MmAChE = *Mus musculus* (murine) acetylcholinesterase. HsAChE = human acetylcholinesterase.
M06-2X/6-31G*/SM8 level [31]. Key interactions between boscartin G and TcAChE are hydrophobic interactions between the ligand and aromatic amino acid side chains of Trp84, Phe330, and His440 (Figure 4B). In addition, there are hydrogen-bonding interactions between the oxirane ring of the ligand and the phenolic -OH of Tyr121 and the C(11)-OH of the ligand and the peptide C=O of His440 (Figure 4B).

**Figure 4.** Lowest-energy docked pose of boscartin G with *Torpedo californica* acetylcholinesterase (TcAChE, PDB 1e66). (A): Ribbon structure of TcAChE with boscartin G in the active site; the co-crystallized ligand (huprene X) is shown as a green wire figure. (B): Key interactions of boscartin G with amino acids in the active site of TcAChE; hydrogen bonds are shown as blue dashed lines.

Boscartin A occupies the active site of TcAChE (Figure 5A, PDB 2cek). As observed for boscartin G with TcAChE, key interactions between the docked ligand and the protein are hydrophobic interactions with Trp84, Phe330, and His440, and a hydrogen-bond between the C(11)-OH of the ligand and the His440 peptide C=O. The conformation of the lowest energy docked pose of boscartin A (Figure 5A) is the same as the lowest-energy calculated (M06-2X/6-31G*/SM8, Figure 5B) [31] and not that found in the X-ray crystal structure [13].
Boscartin A occupies the active site of TcAChE (Figure 5A, PDB 2cek). As observed for boscartin G with TcAChE, key interactions between the docked ligand and the protein are hydrophobic interactions with Trp84, Phe330, and His440, and a hydrogen-bond between the C(11)-OH of the ligand and the His440 peptide C=O. The conformation of the lowest energy docked pose of boscartin A (Figure 5A) is the same as the lowest-energy calculated (M06-2X/6-31G*/*SM8, Figure 5B) [31] and not that found in the X-ray crystal structure [13].

Figure 5. Boscartin A. (A): Lowest-energy docked pose of boscartin A with Torpedo californica acetylcholinesterase (PDB 2cek). (B): Calculated lowest-energy conformation of boscartin A at the M06-2X/6-31G*/SM8 level of theory [31].

A number of bacterial proteins were targeted by Boswellia cembranoids (Table 3). Helicobacter pylori peptide deformylase (HpPDF) and Escherichia coli aspartate transaminase (EcAspTA) were particularly well targeted, while boscartin C and E and incensole oxide acetate showed remarkably exothermic docking energies. Boswellia resin extracts have shown in-vitro antibacterial activity [43–45], and frankincense resins have been used traditionally to treat wounds [6,46] and for oral hygiene [8]. Furthermore, B. papyrifera resin has shown activity against methicillin-resistant Staphylococcus aureus (MRSA) [47] and B. serrata resin showed activity in a clinical trial against plaque-induced gingivitis [48]. The selective targeting of bacterial proteins by Boswellia cembranoids corroborates the traditional medicinal uses and the demonstrated antibacterial activities of frankincense.

The potent docking properties of Boswellia cembranoids with HpPDF are particularly noteworthy. There is a strong association between colonization of the human stomach by Helicobacter pylori and gastrointestinal illnesses such as chronic gastritis and peptic ulcers [49]. Frankincense has been used traditionally to treat stomach disturbances [4] and ulcers [46]. In addition, Boswellia extracts have been shown in clinical studies to be helpful in treating ulcerative colitis [50].

Boscartin G is the strongest binding Boswellia cembranoid ligand with HpPDF (PDB 2ew5). The lowest-energy docked pose is shown in Figure 6. Boscartin G occupies the active site of HpPDF at the same location as the co-crystallized ligand, 4-[(1E)-3-oxo-3-[2-(phenylethyl)amino]prop-1-en-1-yl]-1,2-phenylene diacetate, a cavity surrounded by Ile45, Gly95, Glu94, His138, Cys96, and Gly46 (Figure 6B). The ligand forms two hydrogen-bonds with the peptide N-H groups of Ile45 and Gly46. The docked structure of boscartin G with HpPDF shows the same conformation (Figure 6C) as that predicted from DFT calculations (Figure 6D) [31].
Table 3. MolDock molecular docking energies (kJ/mol) of Boswellia cebranoids with bacterial target proteins. a

| Ligand           | EcAspTA  | SmChiB | SmChiB | SmChiB | SaQacR | SaQacR | SaQacR | SaQacR | HpPDF | BcPhzA/B | BcPhzA/B | MtBioA |
|------------------|----------|--------|--------|--------|--------|--------|--------|--------|-------|----------|----------|--------|
| Boscartin A      | −116.9   | −93.6  | −99.4  | −96.3  | −99.4  | −104.1 | −94.9  | −95.7  | −101.8| −100.3   | −107.8   | −104.9 |
| Boscartin B      | −98.2    | −107.8 | −103.7 | −91.5  | −98.1  | −106.5 | −94.7  | −96.5  | −104.6| −98.4    | −93.9    | −107.5 |
| Boscartin C      | −112.5   | −96.3  | −97.3  | −90.6  | −100.0 | −98.5  | −98.4  | −102.3 | −106.2| −99.3    | −106.4   | −100.4 |
| Boscartin D      | −114.4   | −99.7  | −105.3 | −91.5  | −95.2  | −106.2 | −100.1 | −96.3  | −104.9| −98.0    | −95.8    | −99.3  |
| Boscartin E      | −110.4   | −101.0 | −105.9 | −105.4 | −99.2  | −101.3 | −95.3  | −95.3  | −107.5| −84.7    | −96.7    | −107.7 |
| Boscartin F      | −99.7    | −97.3  | −103.7 | −89.9  | −101.0 | −95.7  | −94.4  | −85.2  | −109.5| −87.7    | −98.0    | −95.1  |
| Boscartin G      | −105.5   | −90.8  | −93.0  | −93.8  | −95.1  | −98.2  | −92.2  | −88.7  | −112.7| −95.4    | −92.6    | −108.5 |
| Boscartin H      | −104.1   | −103.6 | −106.0 | −104.4 | −97.2  | −98.8  | −97.4  | −94.5  | −100.9| −93.4    | −101.5   | −99.5  |
| Incensole        | −104.3   | −102.4 | −101.1 | −96.2  | −85.0  | −96.3  | −90.7  | −95.0  | −94.0 | −92.9    | −93.0    | −95.7  |
| Incensole acetate| −109.9   | −102.3 | −96.2  | −98.1  | −96.6  | −104.5 | −96.1  | −98.8  | −96.2 | −103.7   | −91.5    | −96.2  |
| Incensole oxide  | −108.6   | −92.0  | −93.5  | −90.3  | −97.4  | −99.8  | −94.2  | −95.6  | −101.4| −97.1    | −103.5   | −103.5 |
| Incensole oxide acetate | −119.0 | −101.1 | −98.9  | −96.4  | −96.7  | −99.2  | −100.0 | −106.9 | −105.5| −99.4    | −100.0   | −100.0 |
| Isoincensole oxide| −96.2   | −87.6  | −93.5  | −89.9  | −98.5  | −97.5  | −98.5  | −92.7  | −88.9 | −87.5    | −103.3   | −91.2  |
| Isoincensolol   | −87.3    | −91.2  | −88.9  | −87.9  | −107.9 | −90.0  | −99.6  | −87.8  | −99.7 | −91.5    | −100.0   | −95.2  |
| Serratol        | −90.2    | −90.1  | −89.9  | −82.2  | −90.1  | −91.4  | −97.6  | −89.1  | −103.2| −87.4    | −92.7    | −93.4  |
| Verticillatrine  | −74.0    | −69.5  | −69.0  | −81.2  | −87.2  | −85.5  | −78.5  | −77.3  | −76.5 | −89.9    | −84.4    |        |

a EcAspTA = *Escherichia coli* aspartate transaminase. SmChiB = *Serratia marcescens* chitinase. SaQacR = *Staphylococcus aureus* multidrug binding protein. HpPDF = *Helicobacter pylori* peptide deformylase. BcPhzA/B = *Burkholderia cepacia* phenazine biosynthesis protein A/B. MtBioA = *Mycobacterium tuberculosis* 7,8-diaminopelargonic acid synthase.
The potent docking properties of *Boswellia* cembranoids with HpPDF are particularly noteworthy. There is a strong association between colonization of the human stomach by *Helicobacter pylori* and gastrointestinal illnesses such as chronic gastritis and peptic ulcers [49]. Frankincense has been used traditionally to treat stomach disturbances [4] and ulcers [46]. In addition, *Boswellia* extracts have been shown in clinical studies to be helpful in treating ulcerative colitis [50].

**Boscartin G** is the strongest binding *Boswellia* cembranoid ligand with HpPDF (PDB 2ew5). The lowest-energy docked pose is shown in Figure 6. Boscartin G occupies the active site of HpPDF at the same location as the co-crystallized ligand, 4-{(1E)-3-oxo-3-[(2-phenylethyl)amino]prop-1-en-1-yl]-1,2-phenylene diacetate, a cavity surrounded by Ile45, Gly95, Glu94, His138, Cys96, and Gly46 (Figure 6B). The ligand forms two hydrogen-bonds with the peptide N-H groups of Ile45 and Gly46. The docked structure of boscartin G with HpPDF shows the same conformation (Figure 6C) as that predicted from DFT calculations (Figure 6D) [31].

**Figure 6.** Docking of boscartin G with *Helicobacter pylori* peptide deformylase (HpPDF, PDB 2cek). (A): Ribbon structure of HpPDF showing boscartin G in the active site; the co-crystallized ligand, 4-{(1E)-3-oxo-3-[(2-phenylethyl)amino]prop-1-en-1-yl]-1,2-phenylene diacetate, is shown as a green stick figure. (B): Key interactions of boscartin G in the active site of HpPDF; hydrogen-bonds are shown as blue dashed lines. (C): Conformation of boscartin G docked to HpPDF. (D): Lowest-energy conformation of boscartin G determined by density functional calculations (M06-2X/6-31G*/SM8) [31].

Several *Boswellia* cembranoids showed selective docking to HIV-1 reverse transcriptase (HIV-1-RT) (Table 4). In particular, incensole oxide acetate showed excellent docking ($E_{\text{dock}} < -100$ kJ/mol) to four of the seven HIV1-RT protein crystal structures. The lowest-energy docked pose of incensole oxide acetate with HIV-1 reverse transcriptase (PDB 3mee) is shown in Figure 7. Key interactions between the ligand and the protein are Tyr181, Tyr188, Leu100, Trp229, and Lys103 (Figure 7B). Interestingly, the docking energies for the cembranoids to PDB 3lal and 3t19 are, on average, lower than for the other protein structures. The differences in docking energies can be attributed to the arrangements of the amino acid residues at the binding sites, resulting in different orientations of the docked ligands. Thus, for example, the key amino acids interacting with incensole oxide acetate in PDB 3al are Tyr188, Leu100, Tyr181, Phe227, and Tyr318 (Figure 7C), while PDB 3t19 had Leu100, Tyr188, Val106, Tyr318, and Tyr181 (Figure 7D). That is, binding sites of the protein crystal structures are heavily influenced by
the co-crystallized ligands. Both methanol and aqueous extracts of *Boswellia carteri* have demonstrated HIV-1 reverse transcriptase activity [51].

**Table 4.** MolDock molecular docking energies (kJ/mol) of *Boswellia* cembranoids with human immunodeficiency virus type 1 reverse transcriptase (HIV-RT).

| Ligand       | 1eet | 2hnz | 3irx | 3is9 | 3mee | 3lal | 3t19 |
|--------------|------|------|------|------|------|------|------|
| Boscartin A  | −99.1| −101.7| −96.8| −93.8| −96.0| −69.4| −75.4|
| Boscartin B  | −94.1| −98.5| −100.2| −94.7| −95.6| −72.2| −97.8|
| Boscartin C  | −104.2| −91.5| −97.0| −101.3| −100.5| −87.8| −84.0|
| Boscartin D  | −89.2| −100.5| −97.1| −97.1| −96.4| −78.7| −83.2|
| Boscartin E  | −97.2| −100.6| −91.7| −97.0| −89.4| −60.8| −72.0|
| Boscartin F  | −88.9| −93.8| −98.1| −98.1| −95.1| −88.1| −69.4|
| Boscartin G  | −92.8| −88.7| −97.6| −95.4| −113.8| +9.3| −82.3|
| Boscartin H  | −98.2| −90.4| −96.9| −97.1| −98.2| −86.9| −56.2|
| Incensole    | −83.8| −101.5| −88.8| −88.3| −86.8| −84.0| −94.7|
| Incensole acetate | −95.6| −92.6| −97.9| −97.0| −84.6| −95.9| −91.6|
| Incensole oxide | −105.1| −93.2| −94.9| −98.7| −106.7| −81.7| −79.0|
| Incensole oxide acetate | −107.2| −84.5| −106.8| −106.2| −113.5| −87.8| −87.1|
| Isoincensole oxide | −97.8| −86.5| −89.4| −90.2| −98.7| −77.5| −57.7|
| Isoincensolol | −95.2| −85.6| −98.2| −85.4| −85.9| −70.6| −72.5|
| Serratol     | −90.4| −87.9| −86.8| −89.3| −90.1| −56.0| −48.9|
| Verticillatriene | −81.6| −67.1| −78.4| −81.0| −82.5| −76.9| −25.5|

**Figure 7.** Molecular docking of incensole oxide acetate with HIV-1 reverse transcriptase. (A): Ribbon structure of HIV1-RT (PDB 3mee) showing incensole oxide acetate in the active site. (B): Key interactions of incensole oxide acetate in the active site of HIV1-RT (PDB 3mee). (C): Key interactions of incensole oxide acetate in the active site of HIV1-RT (PDB 3lal). (D): Key interactions of incensole oxide acetate in the active site of HIV1-RT (PDB 3t19).
3.2. Cneorubenoid Diterpenoids

The cneorubenoid diterpenoids, boscartols A–I and olibanumol D, can be considered to be prenylated aromadendranes (Figure 2), and have been isolated from the oleo-gum resin of *Boswellia carteri* [21]. The absolute configuration of the C(15) of boscartol D was not experimentally determined [21]. Nevertheless, both diastereomers, (15R)-boscartol D and (15S)-boscartol D were used in the reverse docking. In addition, the stereochemistry of C(15) was determined theoretically using density functional theory (DFT) conformational analysis carried out at the M06-2X/6-31G* level of theory, including a non-polar (CHCl₃) solvent model. A complete conformational analysis of (15R)-boscartol D was carried out giving 20 low-energy conformations ($E_{\text{rel}} < 14.0$ kJ/mol, accounting for 100% of the Boltzmann distribution of conformers). Similarly, conformational analysis of (15S)-boscartol D returned 13 low-energy ($E_{\text{rel}} < 13.0$ kJ/mol). For each of the conformations, the H-C(15)-C(16)-H dihedral angle was determined and the corresponding vicinal coupling constants ($J_{HH}$) calculated using both the original Karplus equation [52] and the Haasnoot/Altona generalized Karplus equation that includes correction terms for the electronegativity of substituents [53]. Accounting for the Boltzmann distribution, (15R)-boscartol D is predicted to have $J_{HH}$ of 4.3 and 5.1 Hz, respectively. The (15S)-diastereomer, on the other hand, is calculated to have $J_{HH}$ of 6.3 and 6.5 Hz, respectively. The reported $J_{HH}$ coupling constant was 7.6 Hz [21]. Based on the calculated $J_{HH}$ coupling constants, the stereochemistry of boscartol D is predicted to be (15S).

The protein targets that showed the best docking properties with *Boswellia* cneorubenoids are listed in Table 5, along with median docking energies. The protein that was best targeted by *Boswellia* cneorubenoids was *Bacillus anthracis* nucleotide adenylyltransferase (BaNadD, PDB 3hfj) with seven of the 11 ligands showing docking energies $< -120$ kJ/mol. Human folate receptor β (HsFRβ, PDB 4kn0 and 4kn1) was also well targeted with 7/11 cneorubenoids with $E_{\text{dock}} < -120$ kJ/mol. The strongest docking ligands were boscartol E and boscartol I, and both of these ligands targeted BaNadD (PDB 3hfj) and HsFRβ (PDB 4kn0) very well.

| Ligand        | PDB  | $E_{\text{dock}}$ | Target Protein                                      |
|---------------|------|-------------------|-----------------------------------------------------|
| Boscartol A   | 3hfj | −124.2            | *Bacillus anthracis* nucleotide adenylyltransferase (BaNadD) |
|               | 4kn0 | −125.6            | human folate receptor β (HsFRβ)                      |
|               | 4kn1 | −127.5            | human folate receptor β (HsFRβ)                      |
|               |      | −89.7             | Median docking energy                                |
| Boscartol B   | 3hfj | −124.2            | *Bacillus anthracis* nucleotide adenylyltransferase (BaNadD) |
|               | 4kn0 | −127.8            | human folate receptor β (HsFRβ)                      |
|               | 4kn1 | −126.8            | human folate receptor β (HsFRβ)                      |
|               |      | −91.6             | Median docking energy                                |
| Boscartol C   | 3bt9 | −120.2            | *Staphylococcus aureus* multidrug binding protein (SaQacR) |
|               |      | −85.8             | Median docking energy                                |
| (15R)-Boscartol D | 2cek | −120.9            | *Torpedo californica* acetylcholine esterase (TcAChE) |
|               | 3bt9 | −122.7            | *Staphylococcus aureus* multidrug binding protein (SaQacR) |
|               | 3t19 | −125.5            | HIV-1 reverse transcriptase                          |
|               |      | −123.1            | HIV-1 reverse transcriptase                          |
|               |      | −87.6             | Median docking energy                                |
| (15S)-Boscartol D | 3lal | −120.6            | HIV-1 reverse transcriptase                          |
|               |      | −81.8             | Median docking energy                                |
| Boscartol E   | 1s9d | −125.9            | bovine guanine nucleotide exchange factor (BtGEF)    |
|               | 3hfj | −135.1            | *Bacillus anthracis* nucleotide adenylyltransferase (BaNadD) |
|               | 4b80 | −124.1            | murine acetylcholinesterase (MmAChE)                |
|               | 4kn0 | −124.2            | human folate receptor β (HsFRβ)                      |
|               |      | −85.0             | Median docking energy                                |
Nicotinate mononucleotide adenylyltransferase (NadD) has been identified as a target for development of antibacterial agents. The excellent docking of cneorubinoids with BaNdD, along with the known antibacterial activity of frankincense [43–45], corroborates the traditional uses of frankincense to treat wounds [6,46].

*Bacillus anthracis* NadD (PDB 3hfj) is a dimeric structure with the active site at the interface of the two protein monomers (Figure 8). The active site is a hydrophobic pocket formed by Trp116A, Trp116B, Tyr112A, Tyr112B, Lys115A, and Lys115B (Figure 8B).

**Figure 8.** *Bacillus anthracis* nucleotide adenylyltransferase (BaNdD, PDB 3hfj). (A): Lowest-energy docked poses of boscartol E (magenta) and boscartol I (green) in the active site of the dimeric enzyme. (B): Boscartol I in the hydrophobic pocket formed at the interface of the two protein monomers.

Human folate receptor β (HsFRβ) is overexpressed in activated macrophages associated with pathogenesis of inflammatory and autoimmune diseases [54] as well as neoplastic tissues [55]. Thus, antifolates that target folate receptors could be useful for the treatment of cancer and inflammatory diseases [56]. Clinical trials have demonstrated the encouraging results of frankincense treatment for inflammatory and autoimmune diseases such as rheumatoid arthritis, osteoarthritis, Crohn’s disease,
and collagenous colitis [57]. *Boswellia* cneorubenoids may be playing a role in the anti-inflammatory activity of frankincense.

The boccartols docked with HsFRβ in the folate binding site (Figure 9). The cyclopropazulane ring is surrounded by aromatic amino acids Trp187, Tyr101, Tyr76, and Phe78 (Figure 9B). In the case of boccartol A and boccartol B, the terminal –OH group is held in place by hydrogen bonds to Ser73 and Phe78 (Figure 9B).

![Figure 9. Human folate receptor β (HsFRβ, PDB 4kn1). (A): Ribbon structure of protein showed the docked poses of boccartin A (yellow) and boccartin B (purple). (B): Lowest-energy docked pose of boccartin A in the binding site of HsFRβ. Hydrogen-bonds are shown as blue dashed lines.](image)

### 3.3. *Boswellia* Triterpenoids

The *Boswellia* triterpenoids examined in this reverse docking study are shown in Figure 3 and the target proteins with the best docking energies for each triterpenoid ligand are summarized in Table 6. The most receptive protein targets for *Boswellia* triterpenoids were *Staphylococcus aureus* multidrug binding protein (SaQacR, PDB 3bt9) with an average docking energy ($E_{dock}$) of $\text{HsFRβ}\text{−}111.9$ kJ/mol and human fibroblast collagenase (PDB 1cgl) with an average docking energy ($E_{dock}$) of $\text{HsFRβ}\text{−}110.5$ kJ/mol.

| Ligand | PDB  | $E_{dock}$ | Target |
|--------|------|------------|--------|
| (20R)-3,7-Dioxytocall-24-sien-21-oic acid | 3l3m | $-141.8$ | Human poly(ADP-ribose) polymerase-1 (HsPARP-1) (anticancer target) |
| | 1cgl | $-140.3$ | Human tumour necrosis-2 (HsTNF-2) + human poly(ADP-ribose) polymerase-2 (HsPARP-2) (antitumor target) |
| | 2b03 | $-132.4$ | Cavia porcellus 11β-hydroxysteroid dehydrogenase type 3 (Cp11βHSD3) (diabetes target) |
| | | | Porcine pancreatic phospholipase A2 (HsPLA2) (anti-inflammatory target) |
| | | | -$91.3$ | Medium docking energy |
| 11-Ethoxy-β-boswellic acid | 3l3m | $-124.6$ | Torpedo californica acetylcholinesterase (TcAChE) (Alzheimer’s target) |
| | | | -$63.0$ | Medium docking energy |
| 11-Keto-β-boswellic acid | 4b84 | $-125.3$ | Mouse acetylcholinesterase (MmAChE) (Alzheimer’s target) |
| | 36m | $-120.8$ | Torpedo californica acetylcholinesterase (TcAChE) (Alzheimer’s target) |
| | | | -$59.9$ | Medium docking energy |
| 12-Ursene-3,11-dione | 1h22 | $-118.8$ | Torpedo californica acetylcholinesterase (TcAChE) (Alzheimer’s target) |
| | | | -$64.2$ | Medium docking energy |
| 12-Ursene-3,24-diol | 1h22 | $-123.8$ | Torpedo californica acetylcholinesterase (TcAChE) (Alzheimer’s target) |
| | | | +$17.4$ | Medium docking energy |
| 2,3-Dihydrour-12-ursene-24-oic acid | 4b84 | $-128.0$ | Mouse acetylcholinesterase (MmAChE) (Alzheimer’s target) |
| | | | +$62.2$ | Medium docking energy |
| 20,22-Epoxytocall-24-en-3-one | 3l3m | $-133.5$ | Human poly(ADP-ribose) polymerase-1 (HsPARP-1) (anticancer target) |
| | 3g49 | $-125.0$ | Cavia porcellus 11β-hydroxysteroid dehydrogenase type 1 (Cp11βHSD1) (diabetes target) |
| | | | -$88.9$ | Medium docking energy |
| 24-Nor-3,12-ursadien-11-one | 1h22 | $-121.0$ | Torpedo californica acetylcholinesterase (TcAChE) (Alzheimer’s target) |
| | 4b84 | $-120.9$ | Mouse acetylcholinesterase (MmAChE) (Alzheimer’s target) |
| | | | -$90.9$ | Medium docking energy |
| 24-Nor-3,9(11),12-olsatriene | 3l6 | $-112.6$ | Cavia porcellus 11β-hydroxysteroid dehydrogenase type 1 (Cp11βHSD1) (diabetes target) |
| | | | -$31.2$ | Medium docking energy |
| 24-Nor-3,9(11),12-olsatriene | 1h22 | $-118.4$ | Torpedo californica acetylcholinesterase (TcAChE) (Alzheimer’s target) |
| | | | +$52.1$ | Medium docking energy |
| Ligand | PDB | F_value | Target |
|--------|-----|---------|--------|
| 3-Acetoxy-12,20(29)-lupadien-24-oic acid | 3bt9 | −118.4 | Human poly(ADP-ribose) polymerase-1 (anti-cancer target) |
|        | 3bt | −117.9 | Staphylococcus aureus multidrug binding protein (SaQaR) |
|        | +16.4 | Median docking energy |
| 3-Acetoxy-20(29)-lupen-24-oic acid (= Lupeolic acid) | 3bt9 | −117.3 | Human fibroblast collagenase (HMMF-1) (arthritis target) |
|        | +8.0 | Median docking energy |
| 3-β-Hydroxy-20(29)-lupen-24-oic acid (Lupeolic acid) | 3bt9 | −144.7 | Enterococcus cloaca pentaerythritol tetranitrate reductase (ECPETR) (antibacterial target) |
|        | +141.8 | Median docking energy |
|        | −136.4 | Median docking energy |
|        | −136.6 | Median docking energy |
|        | −64.6 | Median docking energy |
| 3-β-Hydroxy-20(29)-lupen-24-oic acid | 3bt9 | −149.9 | Enterococcus cloaca pentaerythritol tetranitrate reductase (ECPETR) (antibacterial target) |
|        | −137.7 | Median docking energy |
|        | −136.2 | Median docking energy |
|        | −134.5 | Median docking energy |
|        | −73.7 | Median docking energy |
| 3-Acetoxy-5,12-ursadien-24-oic acid | 3i6m | −131.7 | Torpedo californica acetylscholastic enzyme (TaACME) (Alzheimer’s target) |
|        | +66.7 | Median docking energy |
| 3-Acetyl-9,11-dehydro-β-hydroxytirucalla-8,24-dien-21-oic acid | 3i6m | −129.0 | Torpedo californica acetylscholastic enzyme (TaACME) (Alzheimer’s target) |
|        | −75.2 | Median docking energy |
| 3-Acetyl-9,11-dehydro-β-hydroxytirucalla-8,24-dien-21-oic acid | 3i6m | −124.0 | Torpedo californica acetylscholastic enzyme (TaACME) (Alzheimer’s target) |
|        | −64.4 | Median docking energy |
| 3-Acetyl-α-boswellic acid | 2b03 | −112.7 | Porcine pancreatic phospholipase A2 (sPLA2) (anti-inflammatory target) |
|        | −64.8 | Median docking energy |
| 3-Acetyl-β-boswellic acid | 3i6m | −128.3 | Torpedo californica acetylscholastic enzyme (TaACME) (Alzheimer’s target) |
|        | −71.5 | Median docking energy |
| 3-Hydroxy-20(29)-lupen-24-oic acid (= Lupeolic acid) | 3bt9 | −119.3 | Staphylococcus aureus multidrug binding protein (SaQaR) |
|        | −55.8 | Median docking energy |
| 3-Oestrus crassipes 7,11,24-trien-21-oic acid | 3i6m | −151.0 | Human tankyrase-2 (HstANK2) = Human poly(ADP-ribose) polymerase-5β (HfPARP-5b) (anticancer target) |
|        | −137.6 | Median docking energy |
|        | −139.9 | Median docking energy |
|        | −127.5 | Median docking energy |
|        | −96.8 | Median docking energy |
| 3-Salvia miltiorrhiza 3,4-dien-21-oic acid | 3i6m | −144.5 | Human tankyrase-2 (HstANK2) = Human poly(ADP-ribose) polymerase-5β (HfPARP-5b) ( anticancer target) |
|        | −130.0 | Median docking energy |
|        | −123.3 | Median docking energy |
|        | −129.1 | Median docking energy |
|        | −127.7 | Median docking energy |
|        | −127.3 | Median docking energy |
|        | −85.6 | Median docking energy |
| 3-Acetylosaurus 7,24-dien-21-oic acid | 3i6m | −147.1 | Human tankyrase-2 (HstANK2) = Human poly(ADP-ribose) polymerase-5β (HfPARP-5b) ( anticancer target) |
|        | −134.1 | Median docking energy |
|        | −128.9 | Median docking energy |
|        | −128.4 | Median docking energy |
|        | −128.4 | Median docking energy |
|        | −102.1 | Median docking energy |
| 3-Hydroxytruxarraca 7,8-diaminopelargonic acid synthase | 3i6m | −154.3 | Human tankyrase-2 (HstANK2) = Human poly(ADP-ribose) polymerase-5β (HfPARP-5b) (anticancer target) |
|        | −128.9 | Median docking energy |
|        | −126.7 | Median docking energy |
|        | −126.4 | Median docking energy |
|        | −98.7 | Median docking energy |
| 3-β-Hydroxytruxarraca 7,8-diaminopelargonic acid synthase | 3i6m | −149.6 | Cavia porcellus β-hydroxysteroid dehydrogenase type 1 (CpHSD1) (diabetes target) |
|        | −132.2 | Median docking energy |
|        | −119.2 | Median docking energy |
|        | −144.7 | Median docking energy |
|        | −111.8 | Median docking energy |
|        | −113.7 | Median docking energy |
| 3-β-Hydroxytruxarraca 7- to 11-carboxylic acid | 3i6m | −121.9 | Human poly(ADP-ribose) polymerase-1 (HfPARP-1) (anticancer target) |
|        | −50.4 | Median docking energy |
| 4,23-Dihydroxybutaric acid | 3i6m | −135.4 | Torpedo californica acetylscholastic enzyme (TaACME) (Alzheimer’s target) |
|        | −127.8 | Median docking energy |
|        | −125.1 | Median docking energy |
|        | −11.7 | Median docking energy |
| 6,7-Epoxide-9(11)-oleic acid | 3i6m | −118.6 | Human soluble epoxide hydrolase (HHEFH2) (anti-inflammatory target) |
|        | −60.7 | Median docking energy |
| 6,7-Epoxide-9(11)-oleic acid | 3i6m | −113.7 | Human prostaglandin F synthase (HfPGF) (hypertension target) |
|        | −49.1 | Median docking energy |
| 9,11-Dehydro-β-boswellic acid | 3i6m | −114.3 | Staphylococcus aureus multidrug binding protein (SaQaR) |
|        | −60.8 | Median docking energy |
| Boscarnene L | 3i6m | −131.0 | Human poly(ADP-ribose) polymerase-1 (HfPARP-1) (anticancer target) |
|        | −56.6 | Median docking energy |
| Boscarnene M | 3i6m | −133.7 | Human poly(ADP-ribose) polymerase-1 (HfPARP-1) (anticancer target) |
|        | −84.9 | Median docking energy |
| Boscarnene N | 3i6m | −157.2 | Human tankyrase-2 (HstANK2) = Human poly(ADP-ribose) polymerase-5β (HfPARP-5b) (anticancer target) |
|        | −130.7 | Median docking energy |
|        | −133.3 | Median docking energy |
|        | −125.2 | Median docking energy |
|        | −95.1 | Median docking energy |
| Dammarenediol II | 3i6m | −135.1 | Human oxidasedehydrogenase (HfODH) (hypercholesterolemia target) |
|        | −134.6 | Median docking energy |
|        | −129.2 | Median docking energy |
|        | −72.1 | Median docking energy |
Table 6. Cont.

| Ligand | PDB* | F-dock | Target |
|--------|------|--------|--------|
| Dammarnediol II acetate | 1frt | -142.8 | Curcumin (HSD11) (diabetes target) |
| 1bx6 | -138.9 | Alpha-acid glycoprotein (AGP) (anti-inflammatory target) |
| 2aba | -136.4 | Enterobacter cloacae pentahydroxyethyl tetramine reductase (EcPETNR) (antibacterial target) |
| 4knl | -132.3 | Human folate receptor β (HFrRβ) (anticancer target) |
| - | -77.7 | Medium docking energy |
| Eupha-2,8,22-triene-20,24-diol<sup>3</sup> | 3lm | -139.8 | Human poly(ADP-ribose) polymerase-1 (HsPARP-1) (anticancer target) |
| 3aU | -135.3 | Human tankyrase-2 (HsTANK2) = human poly(ADP-ribose) polymerase-3b (HsPARP-3b) (antitumor target) |
| 4dbh | -124.0 | Human estrogentic 17β-hydroxysteroid dehydrogenase (17β-1SD) |
| - | -94.9 | Medium docking energy |
| Eupha-2,8,22-triene-20,24-diol<sup>b</sup> | 1uk1 | -137.6 | Human poly(ADP-ribose) polymerase-1 (HsPARP-1) (anticancer target) |
| 3aU | -135.8 | Human tankyrase-2 (HsTANK2) = human poly(ADP-ribose) polymerase-3b (HsPARP-3b) (antitumor target) |
| 4dbh | -127.4 | Human estrogentic 17β-hydroxysteroid dehydrogenase (17β-1SD) |
| - | -85.1 | Medium docking energy |
| Isofouqueryl acetate | 3lt6 | -134.1 | Curcumin (11β-hydroxyesteroid dehydrogenase type 1 (C11HSD)) (diabetes target) |
| 2aba | -131.3 | Enterobacter cloacae pentahydroxyethyl tetramine reductase (EcPETNR) (antibacterial target) |
| 4knl | -130.4 | Human folate receptor β (HFrRβ) (anticancer target) |
| - | -83.2 | Medium docking energy |
| Isofouquyral acetate | 3lt6 | -150.1 | Curcumin (11β-hydroxyesteroid dehydrogenase type 1 (C11HSD)) (diabetes target) |
| 2aba | -127.9 | Enterobacter cloacae pentahydroxyethyl tetramine reductase (EcPETNR) (antibacterial target) |
| 4knl | -127.0 | Human folate receptor β (HFrRβ) (anticancer target) |
| - | -135.7 | Medium docking energy |
| - | -135.5 | Medium docking energy |
| - | -123.7 | Medium docking energy |
| - | -131.0 | Rapamycin D receptor (RapDR) (target for psoriasis) |
| - | -79.5 | Medium docking energy |
| Lup-20(29)-ene-2a,3β-diol | 1ux0 | -112.3 | Human 11β-hydroxyesteroid dehydrogenase type 1 (HSD11β) (diabetes target) |
| 3w2d | -112.4 | Serratio marcescens citrate B (SmCitB) |
| 1xs1 | -112.2 | HIV-1 protease |
| - | -62.2 | Medium docking energy |
| Neolexonol | 1h22 | -118.2 | Torpedo californica acetylcholinesterase (TeAChE) (Alzheimer’s target) |
| - | -66.7 | Medium docking energy |
| Neolexonyl acetate | 2lht | -118.8 | Human 11β-hydroxyesteroid dehydrogenase (Hs11βHSD) (diabetes target) |
| 4dsh | -118.5 | Murine acetylcholinesterase (MmAChE) (Alzheimer’s target) |
| - | -74.8 | Medium docking energy |
| Nizwanone | 4ds4 | -123.2 | Murine acetylcholinesterase (MmAChE) (Alzheimer’s target) |
| - | -63.3 | Medium docking energy |
| Occtillyl acetate | 1gse | -144.3 | Alpha-acid glycoprotein (AGP) (anti-inflammatory target) |
| 2aba | -130.1 | Enterobacter cloacae pentahydroxyethyl tetramine reductase (EcPETNR) (antibacterial target) |
| 4knl | -127.7 | Human endoplasmic reticulum aminoacylase-2 (HsERAP2) (immune response) |
| - | -70.7 | Medium docking energy |
| Olibamunol E | 2lht | -109.5 | Human 11β-hydroxyesteroid dehydrogenase (Hs11βHSD) (diabetes target) |
| - | -63.5 | Medium docking energy |
| Olibamunol F | 1ogl | -110.1 | Human fibroblast collagenase (Mmp1) (arthritis target) |
| 3w49 | -108.4 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| - | -36.7 | Medium docking energy |
| Olibamunol G | 3w49 | -117.4 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| - | -53.0 | Medium docking energy |
| Olibamunol H | 3w49 | -121.6 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| - | -60.9 | Medium docking energy |
| Olibamunol I | 3w49 | -115.7 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| - | -63.2 | Medium docking energy |
| Olibamunol J | 1pw1 | -135.5 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| 1r6 | -133.3 | Murine soluble epoxide hydrolase (MmEPHX2) (anti-inflammatory target) |
| 3bi | -132.4 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| 3ys | -132.1 | Strongyloides stercoralis nuclear receptor SQR-12 (SsAF12) (antiparasitic target) |
| - | -55.5 | Medium docking energy |
| Olibamunol K | 3w49 | -152.3 | Human poly(ADP-ribose) polymerase-1 (HsPARP-1) (anticancer target) |
| 3w5e | -129.0 | Curcumin (11β-hydroxyesteroid dehydrogenase type 1 (C11HSD)) (diabetes target) |
| - | -128.8 | Human phosphodiesterase 4B (HsPDE4B) (anti-inflammatory target) |
| 3w49 | -123.4 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| - | -74.7 | Medium docking energy |
| Olibamunol L | 3w49 | -122.9 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| 3w49 | -122.1 | Torpedo californica acetylcholinesterase (TeAChE) (Alzheimer’s target) |
| - | -64.8 | Medium docking energy |
| 3w49 | -111.1 | Curcumin (11β-hydroxyesteroid dehydrogenase type 1 (C11HSD)) (diabetes target) |
| - | -27.6 | Medium docking energy |
| Olibamunol M | 1u22 | -128.4 | Torpedo californica acetylcholinesterase (TeAChE) (Alzheimer’s target) |
| - | -62.7 | Medium docking energy |
| Olibamunol N | 3w49 | -130.0 | Staphylococcus aureus multidrug binding protein (SaQacR) |
| - | -70.9 | Medium docking energy |
| Trameterolic acid B | 3aU | -131.6 | Human tankyrase-2 (HsTANK2) = human poly(ADP-ribose) polymerase-3b (HsPARP-3b) (antitumor target) |
| 3g4p | -127.1 | Curcumin (11β-hydroxyesteroid dehydrogenase type 1 (C11HSD)) (diabetes target) |
| 3bhb | -126.6 | Human tryptophan hydroxylase type 1 (HtPH1) (biosynthesis of serotonin) |
| 4tev | -126.5 | Torpedo californica acetylcholinesterase (TeAChE) (Alzheimer’s target) |
| - | -74.2 | Medium docking energy |
| Ura-12-ene-3a,11α-diol | 1b22 | -120.9 | Torpedo californica acetylcholinesterase (TeAChE) (Alzheimer’s target) |
| - | -61.2 | Medium docking energy |
| Ura-12-ene-3β,11α-diol | 3o4 | -123.7 | Torpedo californica acetylcholinesterase (TeAChE) (Alzheimer’s target) |
| - | -59.8 | Medium docking energy |
Enterobacter cloacae properties with HsPDE4B with docking energies of $-\text{psoriasis and atopic dermatitis}$ [63], and isofouquierol and olibanumol J showed good docking psoriasis [62]. Similarly, inhibitors of phosphodiesterase 4B have shown promise in the treatment of arthritis [61], and notably, acetate, olibanumol J, and olibanumol J’. There is a significant association of ERAP2 with psoriatic dihydroxydammar-25-ene, 3$\beta$ to human endoplasmic reticulum aminopeptidase 2 (HsERAP2), including 3$\beta$-acetoxy-20$\alpha$Boswellic acid 3g49$\beta$-Boswellic acid 3i6m, and 3$\beta$-acetoxydammar-24-ene-16$\alpha$-Boswellic acid 3g49. Several $\beta$acosytiirucalla-7,24-dien-21-oic acid with HsTANK2, $\beta$-acetoxytirucalla-7,9(11),24-trien-21-oic acid with human glucokinase (PDB 4ixc), $\beta$-acetoxy-20$\alpha$-Boswellic acid 3g49$\beta$-Boswellic acid 3i6m, 3$\beta$-hydroxytirucalla-7,24-dien-21-oic acid with human glucokinase (PDB 4ixc), E$_{\text{dock}}$ = -150.4 kJ/mol; isofouquieryl acetate with Guinea pig (Cavia porcellus) 11$\beta$-hydroxydihydroxydammar-25-ene or for eupha-2,8,22-triene-20,24-diol; both diastereomers for each of these compounds were examined in this reverse docking study.

The best overall triterpenoid ligand–protein target pairs were boscartene N with human tankyrase 2 (HsTANK2, PDB 3ua9), E$_{\text{dock}}$ = -157.2 kJ/mol; 3$\alpha$-hydroxytirucalla-7,24-dien-21-oic acid with HsTANK2, E$_{\text{dock}}$ = -154.3 kJ/mol; $\alpha$-elemolic acid with HsTANK2, E$_{\text{dock}}$ = -152.7 kJ/mol; 3-oxotirucalla-7(11),24-trien-21-oic acid with HsTANK2, E$_{\text{dock}}$ = -151.0 kJ/mol; 3$\alpha$-acetoxytirucalla-7,24-dien-21-oic acid with human glucokinase (PDB 4ixc), E$_{\text{dock}}$ = -150.4 kJ/mol; isofouquieryl acetate with Guinea pig (Cavia porcellus) 11$\beta$-hydroxydihydroxydammar-25-ene type 1 (11H3D1, PDB 3lz6), E$_{\text{dock}}$ = -150.1 kJ/mol; 3$\beta$-acetoxy-20$\alpha$-Boswellic acid 3g49$\beta$-Boswellic acid 3i6m-dihydroxydammar-25-ene with Enterobacter cloaceae pentaerythritol tetranitrate reductase (EcPETNR, PDB 2aba), E$_{\text{dock}}$ = -149.9 kJ/mol; and 3$\beta$-acetoxydammar-24-ene-16$\beta$,20R-diol with Guinea pig 11$\beta$-hydroxydihydroxydammar-25-ene type 1 (Cp11H3D1, PDB 3lz6), E$_{\text{dock}}$ = -149.6 kJ/mol.

Frankincense oleo-gum-resins have been used in traditional medicine to treat a variety of inflammatory conditions, including arthritis, colitis, and asthma [58,59]. Boswellic acids, including $\beta$-boswellic acid, 11-keto-$\beta$-boswellic acid, and acetyl-11-keto-$\beta$-boswellic acid, have been implicated in the anti-inflammatory properties of Boswellia resins; these triterpenoid components are involved in inhibition of 5-lipoxygenase (5-LOX), inducible nitric oxide synthase (iNOS), cyclooxygenase-1 (COX-1) and cyclooxygenase-2 (COX-2) [60].

In-silico screening of the Boswellia triterpenoids was carried out against molecular targets of inflammation, including human pancreatic secretory phospholipase A2 (HsPLA2), porcine pancreatic phospholipase A2 (SpPLA2), human phosphoinositide 3-kinase (HsPI3K), human interleukin-1 receptor associated kinase 4 (HsIRAK4), human glutathione transferase omega 1 (HsGSTO1), human 5-lipoxygenase (Hs5-LOX), mouse inducible nitric oxide synthase (MmiNOS), ovine COX-1 (OaCOX-1), murine COX-2 (MmCOX-2), human fibroblast collagenase (matrix metalloproteinase-1, HsMMP-1), murine soluble epoxide hydrolase 2 (MmEPHX2), human endoplasmic reticulum aminopeptidase 2 (HsERAP2), human soluble epoxide hydrolase 2 (HsEPHX2), and human phosphodiesterase 4B (HsPDE4B). The docking energies of the triterpenoid ligands with inflammatory target proteins are shown in Table 7.

The Boswellia triterpenoids showed relatively weak docking to 5-LOX, iNOS, PI3K, IRAK4, or GSTO1, and no docking at all to either COX-1 or COX-2 (positive docking energies). However, several Boswellia triterpenoids showed relatively strong docking (E$_{\text{dock}}$ < $-120$ kJ/mol) to human endoplasmic reticulum aminopeptidase 2 (HsERAP2), including 3$\beta$-acetoxy-20$\alpha$,24S-dihydroxydammar-25-ene, 3$\beta$-acetoxydammar-24-ene-16$\beta$,20R-diol, isofouquieryl acetate, ocolityl acetate, olibanumol J, and olibanumol J’. There is a significant association of ERAP2 with psoriatic arthritis [61], and notably, Boswellia triterpenoids have shown promise for the treatment of psoriasis [62]. Similarly, inhibitors of phosphodiesterase 4B have shown promise in the treatment of psoriasis and atopic dermatitis [63], and isofouquierol and olibanumol J’ showed good docking properties with HsPDE4B with docking energies of $-128.3$ and $-128.8$ kJ/mol, respectively.

### Table 6. Cont.

| Ligand                  | PDB * | $E_{\text{dock}}$ | Target                                                                 |
|-------------------------|-------|-------------------|------------------------------------------------------------------------|
| $\alpha$-boswellic acid | 3g49  | -110.9            | Cavia porcellus 11$\beta$-hydroxydihydroxydammar-25-ene type 1 (11H3D1) (diabetes target) |
| $\alpha$-elemolic acid  | 3ua9  | -152.7            | Human tankyrase-2 (HsTANK2) = human poly(ADP-ribose) polymerase-5b (HsPARP-5b) (anticancer target) |
| $\alpha$-elemolic acid  | 3a6   | -136.9            | Apaf-1 and/or histone deacetylase (AdAHDAC) (anticancer target) |
| $\alpha$-elemolic acid  | 3g49  | -128.9            | Cavia porcellus 11$\beta$-hydroxydihydroxydammar-25-ene type 1 (11H3D1) (diabetes target) |
| $\beta$-boswellic acid  | 3tm   | -118.1            | Torpedo californica acetylcholinesterase (TcACHE) (Alzheimer’s target) |
| $\beta$-elemenic acid   | 3ua9  | -147.3            | Human tankyrase-2 (HsTANK2) = human poly(ADP-ribose) polymerase-5b (HsPARP-5b) (anticancer target) |
| $\beta$-elemenic acid   | 3tm   | -133.7            | Human poly(ADP-ribose) polymerase-1 (HsPARP-1) (anticancer target) |
| $\beta$-boswellic acid  | 3g49  | -133.9            | Cavia porcellus 11$\beta$-hydroxydihydroxydammar-25-ene type 1 (11H3D1) (diabetes target) |
| $\beta$-boswellic acid  | 3g49  | -106.7            | Cavia porcellus 11$\beta$-hydroxydihydroxydammar-25-ene type 1 (11H3D1) (diabetes target) |

* PDB: Protein Data Bank code.  The stereochemistry at C(24) was not experimentally determined for 3$\beta$-acetoxy-20$\alpha$,24S-dihydroxydammar-25-ene or for eupha-2,8,22-triene-20,24-diol; both diastereomers for each of

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The only other strong docking observed was olibanumol J with murine soluble epoxide hydrolase 2 (MmEPHX2), 3β-acetoxytirucalla-7,24-dien-21-oic acid with human matrix metalloproteinase-1 (HsMMP-1), (20S)-3,7-dioxotirucalla-8,24-dien-21-oic acid with porcine pancreatic phospholipase A2 (SwPLA2), and 3β-acetoxy-20S,24-dihydroxydammar-25-ene with human pancreatic secretory phospholipase A2 (HsPLA2). The targeting of matrix metalloproteinase-1 (fibroblast collagenase) is noteworthy; *B. serrata* extract has shown clinical efficacy as a treatment for osteoarthritis of the knee [64].

Note that β-boswellic acid, 11-keto-β-boswellic acid, and 3-acetyl-11-keto-β-boswellic acid had relatively weak docking with inflammation-relevant protein targets. It may be that these *Boswellia* triterpenoids, rather than inhibiting particular enzyme targets, are inhibiting the secretion of pro-inflammatory cytokines such as tumor necrosis factor α (TNFα), interleukin 1 (IL-1), IL-6, IL-12, IL-18, or interleukin γ (IFN-γ) [65].

Olibanumols G, H, I, and J all showed selective docking to *S. aureus* multidrug binding protein (SaQacR). The olibanumols occupy the binding site of SaQacR, same site as the co-crystallized ligand in PDB 3bti (Figure 10). The site is made up of aromatic amino acids Trp61, Tyr93, and Tyr123, forming a hydrophobic pocket. The triterpenoid ligand olibanumol J also has a hydrogen-bonding interaction between C(3)-OH of the ligand and the amide C=O of Ala153. Other key interacting amino acids in the binding site are Ser86, Glu90, and Asn157. In addition to the olibanumols, six additional triterpenoids also showed selective docking to SaQacR, 3-acetoxy-12,20(29)-lupadien-24-oic acid, lupeolic acid, 3-oxotirucalla-7,9(11),24-trien-21-oic acid, 4,23-dihydroburic acid, 9,11-dehydro-β-boswellic acid, and boscartene N.

Figure 10. Lowest-energy docking pose of olibanumol J with *Staphylococcus aureus* multidrug binding protein (SaQacR, PDB 3bti). (A): Ribbon structure of SaQacR showing olibanumol J (*E*$_{\text{dock}}$ = −132.6 kJ/mol) in the active site. The co-crystallized ligand, berberine (*E*$_{\text{dock}}$ = −113.8 kJ/mol), is shown as a green wire structure. (B): Key interactions of olibanumol J in the active site of SaQacR. The hydrogen-bond is indicated by a blue dashed line.

Another antibacterial target protein that showed good docking properties was *Enterobacter cloacae* pentaerythritol tetranitrate reductase (EcPETNR) with seven *Boswellia* dammarane triterpenoids showing selective docking to this target, 3β-acetoxy-20S,24-dihydroxydammar-25-ene (both diastereomers), 3β-acetoxydammar-24-ene-16β,20R-diol, dammarenediol II acetate, isofouquierol, isofouquieryl acetate, and ocofyll acetate. These dammarane triterpenoids occupy the active site of EcPETNR, near to the riboflavin monophosphate redox cofactor, with very similar docked poses (Figure 11). Key intermolecular contacts are hydrogen bonding of the C(3) acetoxygroup with Arg142, hydrogen-bonding of C(20)-OH with the riboflavin monophosphate cofactor and His184 (Figure 11B). In the case of 3β-acetoxy-20S,24S-dihydroxydammar-25-ene, there is also a hydrogen-bond between the C(24)-OH group and Asp274.
forming a hydrophobic pocket. The triterpenoid ligand olibanumol J also has a hydrogen-bonding interaction between C(3)-OH of the ligand and the amide C=O of Ala153. Other key interacting amino acids in the binding site are Ser86, Glu90, and Asn157. In addition to the olibanumols, six additional triterpenoids also showed selective docking to SaQacR, 3-acetoxy-12,20(29)-lupadien-24-oic acid, lupeolic acid, 3-oxotirucalla-7,9(11),24-trien-21-oic acid, 4,23-dihydroburic acid, 9,11-dehydro-β-boswellic acid, and boscartene N.

Figure 10. Lowest-energy docking pose of olibanumol J with Staphylococcus aureus multidrug binding protein (SaQacR, PDB 3bti). (A): Ribbon structure of SaQacR showing olibanumol J (E<sub>dock</sub> = −132.6 kJ/mol) in the active site. The co-crystallized ligand, berberine (E<sub>dock</sub> = −113.8 kJ/mol), is shown as a green wire structure. (B): Key interactions of olibanumol J in the active site of SaQacR. The hydrogen-bond is indicated by a blue dashed line.

Another antibacterial target protein that showed good docking properties was Enterobacter cloacae pentaerythritol tetranitrate reductase (EcPETNR) with seven Boswellia dammarane triterpenoids showing selective docking to this target, 3β-acetoxy-20S,24-dihydroxydammar-25-ene (both diastereomers), 3β-acetoxydammar-24-ene-16β,20R-diol, dammarenediol II acetate, isofouquierol, isofouquieryl acetate, and ocotillyl acetate. These dammarane triterpenoids occupy the active site of EcPETNR, near to the riboflavin monophosphate redox cofactor, with very similar docked poses (Figure 11). Key intermolecular contacts are hydrogen bonding of the C(3) acetoxy group with Arg142, hydrogen-bonding of C(20)-OH with the riboflavin monophosphate cofactor and His184 (Figure 11B). In the case of 3β-acetoxy-20S,24S-dihydroxydammar-25-ene, there is also a hydrogen-bond between the C(24)-OH group and Asp274.

Drugs that reversibly inhibit acetylcholinesterase are currently being explored to treat Alzheimer’s disease [66], and Torpedo californica and murine acetylcholinesterases have been used as model enzyme targets for anticholinesterase inhibition [67]. β-Boswellic acid and its derivatives (11-keto-β-boswellic acid, 11-ethoxy-β-boswellic acid, 3-acetyl-β-boswellic acid, and 3-acetyl-11-keto-β-boswellic acid) showed selective docking to Torpedo californica acetylcholinesterase (TcACHe, PDB 3i6m). Interestingly, increasing oxygenation resulted in more exothermic docking energies (−118.1, −120.8, −124.6, −128.3, and −129.7 kJ/mol, respectively). These β-boswellic acid derivatives all adopt the same lowest-energy poses in the active site of the enzyme (Figure 12). It is tempting to suggest that the ordering of docking energies is due to increasing hydrogen-bonding of the more oxygenated ligands. However, the only hydrogen bonding seen is with the C(3)-substituent of the ligand (either –OH or –OAc) with Tyr121 (see Figure 10B). Note that the active site of the acetylcholinesterase is surrounded by aromatic amino acids (Trp84, Tyr334, Tyr121, Phe330, Phe331, and Trp279). The more important interactions, therefore, are van der Waals hydrophobic interactions between the ligand and the aromatic amino acids. The trend in docking energies for these β-boswellic acid derivatives is likely due to the increased number of heavy atoms and increasing molecular weight. Olibanumols K, L’, M, and N also showed selective docking to acetylcholinesterase proteins. The strongest-docking triterpenoid ligands to TcAChE were 4,23-dihydroburic acid (E<sub>dock</sub> = −135.8 kJ/mol) and 3-acetoxy-5,12-ursadien-24-oic acid (E<sub>dock</sub> = −131.7 kJ/mol); olibanumol N docked strongly with MmAChE (E<sub>dock</sub> = −133.0 kJ/mol). The excellent docking properties of Boswellia triterpenoid ligands to acetylcholinesterase supports the clinical use of frankincense to treat Alzheimer’s disease [41].

Figure 11. Enterobacter cloacae pentaerythritol tetranitrate reductase (EcPETNR, PDB 2aba). (A): Ribbon structure of EcPETNR with docked dammarane triterpenoids (stick figures). The riboflavin monophosphate cofactor is shown as a space-filling model. (B): Lowest-energy docked pose of 3β-acetoxy-20S,24S-dihydroxydammar-25-ene. Hydrogen-bonding interactions are indicated with blue dashed lines.
Table 7. MolDock docking energies (kJ/mol) of Boswellia triterpenoid ligands with inflammation-relevant protein targets.

| Ligand | MnEPHX2 | HsEPHX2 | HsMMMP-1 | SaPLA2 | HsPLA2 | Hs-LOX | Mn INOS | HsPKY | HsIRAK4 | HsERAP2 | HsGSTO1 | HsPDE4B |
|--------|----------|----------|-----------|--------|--------|--------|---------|--------|---------|---------|---------|---------|
| 1cr6   | 70.1     | 98.1     | 105.0     | 105.1  | 104.5  | 103.4  | 98.1    | 105.0  | 105.1   | 113.4   | 98.1    | 113.4   |
| 4hai   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 1cgI   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 2b03   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 11ja   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 3v99   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 1m8d   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 2a3u   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 51s    | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 4jbs   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 5v3q   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |
| 3w5e   | 98.1     | 105.0    | 105.1     | 104.5  | 103.4  | 98.1   | 105.0   | 105.1  | 113.4   | 98.1    | 113.4   | 105.0   |

**Note:** The table lists the docking energies of various Boswellia triterpenoid ligands with inflammation-relevant protein targets, measured in kilojoules per mole (kJ/mol). The ligands are compared across different proteins such as MnEPHX2, HsEPHX2, HsMMMP-1, SaPLA2, HsPLA2, Hs-LOX, Mn INOS, HsPKY, HsIRAK4, HsERAP2, HsGSTO1, and HsPDE4B.
Table 7. Cont.

| Ligand                          | MnEPHX2 | HsEPHX2 | HsMMP-1 | SaPLA2 | HsPLA2 | Hs5-LOX | Mm iNOS | HsPI3Kγ | HsIRAK4 | HsERAP2 | HsGSTO1 | HsPDE4B |
|--------------------------------|---------|---------|---------|--------|--------|---------|---------|---------|---------|---------|---------|---------|
|                                | 1cr6    | 4hai    | 1clg    | 2b03   | 1j1a   | 3v99    | 1m8d   | 2a5u   | 5t1s   | 4jbs   | 5v3q   | 3w5e   |
| Olibanumol E                   | −81.7   | −82.4   | −101.7  | −98.7  | −91.8  | −82.2   | −60.6  | −79.4  | −92.4  | −97.9  | −49.2  | −99.2  |
| Olibanumol F                   | −85.7   | −108.9  | −110.1  | −86.1  | −79.7  | −87.6   | −65.9  | −86.4  | −80.4  | −95.0  | −77.3  | −46.1  |
| Olibanumol G                   | −81.5   | −95.3   | −99.3   | −82.4  | −70.4  | −80.0   | −61.0  | −77.9  | −80.8  | −93.2  | −54.0  | −72.7  |
| Olibanumol H                   | −101.1  | −113.9  | −107.7  | −102.3 | −86.8  | −82.6   | −78.1  | −88.4  | −94.2  | −103.8 | −74.1  | −55.0  |
| Olibanumol I                   | −96.2   | −97.4   | −101.5  | −93.4  | −70.6  | −81.9   | −75.5  | −78.2  | −88.8  | −96.9  | −71.6  | −79.2  |
| Olibanumol J                   | −133.3  | −119.3  | −114.2  | −112.5 | −117.6 | −106.1  | −93.8  | −88.4  | −108.3 | −123.8 | −99.3  | −112.3 |
| Olibanumol L                   | −108.2  | −119.2  | −112.7  | −112.3 | −118.8 | −107.7  | −84.4  | −84.3  | −109.2 | −124.4 | −84.6  | −128.8 |
| Olibanumol M                   | −88.8   | −77.8   | −110.3  | −90.5  | −94.9  | −77.0   | −64.9  | −79.8  | −76.2  | −89.9  | −78.8  | −82.4  |
| Olidanumol N                   | −78.1   | −80.8   | −109.6  | −91.7  | −98.1  | −81.3   | −67.6  | −73.8  | −82.4  | −97.3  | −64.1  | −108.1 |
| Trametenolic acid B            | −85.0   | −74.1   | −111.4  | −97.1  | −83.9  | −84.7   | −67.6  | −77.1  | −72.7  | −92.4  | −70.0  | −37.9  |
| Urs-12-ene-3β,11α-diol         | −77.5   | −89.2   | −100.1  | −95.5  | −57.5  | −83.9   | −69.2  | −77.1  | −96.5  | −93.1  | −63.9  | −100.0 |
| Urs-12-ene-3β,11α-diol         | −85.0   | −76.5   | −103.3  | −84.5  | −70.8  | −86.3   | −73.6  | −79.7  | −87.9  | −96.8  | −70.4  | −50.1  |
| Urs-12-ene-3β,11α-diol         | −106.3  | −107.3  | −106.9  | −107.3 | −107.9 | −99.7   | −92.6  | −84.0  | −107.6 | −115.0 | −92.9  | −111.6 |
| Urs-12-ene-3β,11α-diol         | −106.3  | −110.5  | −113.9  | −124.4 | −104.1 | −99.7   | −107.1 | −84.0  | −108.7 | −111.1 | −97.3  | −110.3 |
| Urs-12-ene-3β,11α-diol         | −79.4   | −74.8   | −101.3  | −108.0 | −71.2  | −76.9   | −67.3  | −69.9  | −70.9  | −93.8  | −61.8  | −114.1 |
| Urs-12-ene-3β,11α-diol         | −77.9   | −87.8   | −103.7  | −100.2 | −95.0  | −80.7   | −64.4  | −79.4  | −85.7  | −94.8  | −62.5  | −62.0  |
| Urs-12-ene-3β,11α-diol         | −106.3  | −110.5  | −113.9  | −124.4 | −104.1 | −99.7   | −107.1 | −84.0  | −108.7 | −111.1 | −97.3  | −110.3 |
| Urs-12-ene-3β,11α-diol         | −79.4   | −87.8   | −108.9  | −94.2  | −90.0  | −82.5   | −7.8   | −76.7  | −91.1  | −94.4  | −85.2  | −95.5  |
| Urs-12-ene-3β,11α-diol         | −112.2  | −113.8  | −110.5  | −109.6 | −100.4 | −92.4   | −86.3  | −106.6 | −107.1 | −108.0 | −112.6 |
| Urs-12-ene-3β,11α-diol         | −110.0  | −114.6  | −120.2  | −120.4 | −106.3 | −99.8   | −83.5  | −109.2 | −109.0 | −103.6 | −116.2 |
| Urs-12-ene-3β,11α-diol         | −76.4   | −90.4   | −101.3  | −89.6  | −82.5  | −74.7   | −69.2  | −80.7  | −87.2  | −96.4  | −64.4  | −76.4  |
Figure 12. Lowest-energy docked poses of β-boswellic acid derivatives with Torpedo californica acetylcholinesterase (TcAChE, PDB 3i6m). (A): Ribbon structure with docked ligands, β-boswellic acid (brown), 11-keto-β-boswellic acid (magenta), 11-ethoxy-β-boswellic acid (yellow), 3-acetyl-β-boswellic acid (red), and 3-acetyl-11-keto-β-boswellic acid (aqua). (B): Molecular environment of docked β-boswellic acid in the active site of TcAChE. (C): Molecular environment of docked 3-acetyl-11-keto-β-boswellic acid in the active site of TcAChE.

Several protein targets related to cancer were targeted by Boswellia triterpenoids, including poly(ADP-ribose) polymerase-1 (PARP-1), tankyrase-1 (TANK1), tankyrase-2 (TANK2), folate receptor β (FRβ), and histone deacetylase (HDAC). Poly(ADP-ribose) polymerase-1 (PARP-1) is an important enzyme for the repair of single-strand DNA breaks, and inhibition of PARP-1 can cause multiple double strands DNA breaks to occur, leading to cell death [68]. Several types of cancers are more dependent on PARP-1 than normal cells, so PARP-1 has become an attractive target for cancer
chemotherapy [69,70]. At least 15 Boswellia triterpenoids showed selective docking to PARP-1 (see Table 6). The strongest-docking ligands for human PARP-1 were (20S)-3,7-dioxotirucalla-8,24-dien-21-oic acid ($E_{\text{dock}} = -141.8$ kJ/mol) and eupha-2,8,22-triene-20,24R-diol ($E_{\text{dock}} = -139.8$ kJ/mol).

Folate receptors are overexpressed in cancer cells, presumably due to the increased requirement of cancer cells for folic acid needed in cell proliferation [71] and folate receptor-β is overexpressed in lung, liver, skin, and soft tissue tumors, as well as associated stromal cells. Folate receptors, therefore, show promise as chemotherapeutic targets for cancer and other human pathologies [72]. Three of the Boswellia triterpenoids in this study showed excellent docking properties to human folate receptor β (HsFRβ), namely dammarenediol II acetate, isofouquierol, and isofouquieryl acetate, with docking energies of $-132.3$, $-130.4$, and $-137.0$ kJ/mol, respectively.

Histone deacetylases (HDAC) are enzymes that remove acetyl groups from lysine residues of histones, which allow the histones to envelope DNA more tightly. Thus, HDACs can affect cell growth and differentiation and cell death [73,74]. Histone deacetylase has been recognized as a promising target for cancer chemotherapy [75,76]. Reverse docking of Boswellia triterpenoids has revealed α-elemol to preferentially dock to Aquifex aeolicus histone deacetylase (AaHDAC).

Frankincense-containing formulations had been used in ancient Greece for treating various malignant tumors [77]. Frankincense has been used in the Indian traditional medicine (Ayurveda) and in Traditional Chinese Medicine (TCM) as a treatment for proliferative diseases [78]. In addition, extracts of frankincense oleo-gum resins have shown in-vitro cytotoxic activity on several human tumor-derived cell lines [58,79–81], and these activities have been attributed to boswellic acids [82]. Interestingly, although β-boswellic acid [83] and 3-acetyl-11-keto-β-boswellic acid [84] have shown antineoplastic activities, this reverse-docking study did not reveal particularly notable docking properties to cancer-relevant protein targets. It may be that the boswellic acids and derivatives are targeting inflammatory pathways [85,86] or multiple targets as their mechanisms of antineoplastic activities [82,84]

Several Boswellia triterpenoids showed good docking properties to 11β-hydroxysteroid dehydrogenase type 1 (11βHSD1). For example, the dammarane triterpenoids 3β-acetoxy-20S,24-dihydroxydammar-25-ene, 3β-acetoxydammar-24-ene-16β,20R-diol, dammarenediol II, dammarenediol II acetate, isofouquierol, and isofouquieryl acetate showed excellent docking energies with Cavia porcellus 11β-hydroxysteroid dehydrogenase type 1 (Cp11βHSD1, PDB 3lz6). These dammarane triterpenoids all occupy the same position in the active site of the enzyme, blocking access to the NADPH cofactor (Figure 13). The docked dammaranes are sandwiched between the NADPH cofactor and hydrophobic amino acids Tyr152, Tyr98, Tyr158, Leu192, and Tyr206 (Figure 13B). There are also close contacts, but no apparent hydrogen-bonds, with Thr197 and Asn99. 11β-Hydroxysteroid dehydrogenase type 1 mediates the interconversion of cortisone and cortisol and overexpression of 11βHSD1 can lead to metabolic disease, characterized by visceral obesity, hyperlipidemia, hypertension, glucose intolerance, insulin resistance, and type II diabetes [87,88]. Thus, inhibition of 11βHSD1 may serve as a treatment option for metabolic syndrome and type II diabetes [89,90].

Boswellia serrata is used traditionally by diabetic patients in Iran, and B. serrata supplementation has shown clinical benefit in blood lipid and glucose levels in type II diabetic patients [91,92]. Furthermore, B. serrata resin extract has been shown to prevent increase in blood glucose levels in streptozotocin-induced diabetic mice [93]. Similarly, B. glabra extracts have shown hypoglycemic effects in alloxan-induced diabetic rats [94]. The selective targeting of Guinea pig 11βHSD1 and human 11βHSD1 by Boswellia triterpenoids is consistent with the traditional use and anti-diabetic activities of Boswellia oleo-gum resin.

Oxidosqualene cyclases (OSCs) are enzymes that catalyze the cyclization of 2,3-epoxysqualene to form triterpenoids or steroids [95]. In mammals, cyclization of 2,3-epoxysqualene leads to lanosterol, which can then be converted to cholesterol [96]. Inhibition of oxidosqualene cyclase, therefore, has emerged as a viable therapeutic option to treat hypercholesterolemia and atherosclerosis [97,98].
The oxidosqualene-hopene cyclase from the thermophilic bacterium *Alicyclobacillus acidocaldarius* is homologous to the human enzyme and has been crystallized with OSC inhibitors [99]. Several triterpenoid ligands, most notably octilloyl acetate, 3β-acetoxydammar-24-ene-16β,20R-diol, and dammarenediol II acetate, docked well to AaOSC with docking energies of −144.6, −139.2, and −138.9 kJ/mol, respectively. These dammarane triterpenoids adopt the same positions in the active site of the enzyme (Figure 14). The active site of AaOSC is a hydrophobic pocket composed of Trp489, Trp169, Phe365, Ile261, Trp312, Phe601, and Tyr420. In addition, there is a hydrogen bond formed between the C(24)-OH of the ligand and the phenolic -OH of Tyr609. The structural similarities between triterpenoids and steroids are likely responsible for the docking properties of *Boswellia* triterpenoids to OSCs.

![Figure 13](image1.png)

**Figure 13.** Guinea pig 11β-hydroxysteroid dehydrogenase type 1 (Cp11β-HSD, PDB 3lz6). (A): Lowest-energy docked poses of 3β-acetoxy-20S,24R-dihydroxydammar-25-ene (magenta), 3β-acetoxydammar-24-ene-16β,20R-diol (green), dammarenediol II acetate (yellow), and isofouquieryl acetate (blue) with Cp11β-HSD. The NADPH cofactor is shown as a space-filling model. (B): Isofouqueryl acetate in the active site of Cp11β-HSD showing the molecular environment.

![Figure 14](image2.png)

**Figure 14.** *Alicyclobacillus acidocaldarius* oxidosqualene cyclase (AaOSC, PDB 1h36). (A): Lowest energy docked poses of octilloyl acetate (magenta), 3β-acetoxydammar-24-ene-16β,20R-diol (green), and dammarenediol II acetate (yellow) with AaOSC. (B): Molecular environment of docked octilloyl acetate in the active site of AaOSC.
4. Conclusions

Numerous *Boswellia* terpenoid components have shown selective docking to bacterial protein targets, antineoplastic molecular targets, diabetes-relevant targets, protein targets involved in inflammatory disease conditions, and the Alzheimer’s disease target acetylcholinesterase. The molecular docking properties of *Boswellia* terpenoid components corroborate the traditional uses of frankincense, the clinical efficacy of frankincense, and the biological activities of *Boswellia* oleo-gum resins and components. Furthermore, the biomolecular targets identified in this work should lead to further exploration of development and improvement of inhibitors to treat these various disease states.

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