Acetyleugenol from *Acacia nilotica* (L.) exhibits a strong antibacterial activity and its phenyl and indole analogs show a promising anti-TB potential

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Abstract: Acetyleugenol is a phytochemical compound with broad effect against infectious diseases and tumors. Here, we extracted, characterized and elucidated the structure of acetyleugenol, for the first time, from the leaves of *Acacia nilotica* (L)—a well-known medicinal plant. The broad antibacterial potential of acetyleugenol was first confirmed against seven bacterial pathogenic isolates with best activity against *Proteus* sp., *Salmonella typhi*, *Staphylococccus aureus*, and *Streptococcus pneumonia*, which showed similar or better zone of inhibition to that of the control amoxicillin. To further investigate its effect against *Mycobacterium tuberculosis*, acetyleugenol and its indole and phenyl analogs were subjected to molecular docking experiments against two potential tuberculosis drug targets—MtPknE and MtPknB Ser/Thr protein kinases. The results reveal that all of the analogs have improved docking scores comparing to the acetyleugenol. The indole analogs EUG-1 and EUG-3 were more effective with better docking scores for MtPknE with $-11.08$ and $-10.05$ kcal/mol, respectively. Similar results were obtained for the MtPknB. In contrast, only the EUG-2 phenyl analog has given rise to similar docking scores for both targets. This opens the door for further comprehensive studies on these acetyleugenol analogs with *in vitro* and *in vivo* experiments to validate and get more insights into their mechanisms of action.

Keywords: acetyleugenol; *Acacia nilotica*; anti-tuberculosis; Serine/Threonine kinases; multi-drug resistant TB.

1. Introduction

Tuberculosis (TB) caused by *Mycobacterium tuberculosis* is among the top 10 causes of death worldwide [1], and represents serious challenges that faces the global disease eradication policy owing to the growing antimicrobial resistance [2,3]. TB kills annually around 1.5 million people with over 80% in the developing countries including India, China, Indonesia, Philippines, Pakistan, Nigeria, Bangladesh and South Africa [4]. TB treatment regimen is a complex procedure that involves the use of several antibiotics for a long-period of time and requires special awareness from patients. The lack of such awareness leads often to the emergence of multi-drug resistant tuberculosis (MDR-TB), which is a consequence of the lack of good education and poor hygienic condition and other environmental factors especially in developing countries [5]. This has also increases the toll of death due to TB among HIV-positive patients [6,7]. The drug resistance developed by *M. tuberculosis* against first line anti-tuberculosis (anti-TB) drugs—Isoniazid and Rifampicin, has become a major problem worldwide [8-10]. This has been complicated by the emergence of resistance against other anti-TB
drugs including, for example, the quinolones and aminoglycosides [11,12]. The emergence of the MDR-TB has

driven the needs for the discovery of novel therapeutic agents to overcome the TB challenge worldwide [9-12].

Medicinal plants are considered a major source in drug discovery which have well-documented
applications as natural remedies for the treatment of many infectious diseases including those infecting the
respiratory system [13]. Plants produce many secondary metabolites in tremendous amounts and many of
which have been used as templates or scaffolds for synthesis of novel pharmaceutical drugs [13-15]. In the
developing countries plants remain the major source of medicines for centuries [16,17]. Sub-Saharan Africa is
rich in medicinal plants, which have been used extensively to treat infectious diseases especially those of the
respiratory system such as tuberculosis, asthma, influenza and pneumonia [17]. For example, in South Africa
alone there are over 300 medicinal plants, with ~6% species belong to the family Fabaceae, that have been used in
remedies of various respiratory infections [18]. The medicinal plant Acacia nilotica (L.)—native to
Sudan—from to the family Fabaceae is well-known for its medicinal applications in the treatment of various
respiratory system diseases and other diseases including malaria and Hepatitis C virus as well as wound
healing and disinfection [19-21]. A. nilotica (L.) is rich with several groups of active phytochemical constituents
including alkaloids, flavonoids, tannins and saponins, gallic acid, kaempferol, umbelliferone, and niloticanes
[22-24]. Its pharmacological activities are broad—mostly attributed to these phytochemicals—covering
antibacterial, antifungal, anti-mutagenic, anti-inflammatory and antioxidant activities [20,22,25].

Eugenol, 4-allyl-2-methoxybenzene, is a natural product that is produced by many angiosperms and
known for its aromatic flavor [26,27]. It is a phenolic compound that belongs to the group of phenylpropanes,
which is biosynthesized via the shikimic acid pathway [27]. Its derivative acetyl-eugenol, with the formula
C₁₀H₁₅O₃ and molecular weight 206.238 Da, is one of the major components of clove Syzygium aromaticum (L.)
essential oil and is also produced by many other plants such as cinnamon, pimento, ylang ylang and rose [28].
Eugenol has also been detected in the chloroform extracts of the A. nilotica (L.) leaves without further chemical
description [29]. Eugenol is slightly acidic and very reactive with a stronger action than alcohols [28]. It has a
wide range of applications in food and cosmetic industry as flavor and antioxidant as well as in the traditional
medicine [26,28,30]. Eugenol and its derivatives have also been reported to have potential activities as anti-
inflammatory, anti-tumor, antimicrobial, anti-pyretic, analgesic and anesthetic [31]. Its benzoate and 2-
nitrobenzoate derivatives as well as its synergistic effects with approved anti-TB drugs have been shown to
exhibits strong activity against Mycobacterium spp. and M. tuberculosis [32]. Its hydrazone derivatives were
studied by molecular docking against the enoyl-ACP reductase as a molecular target and validated by in vitro
assay which showed a docking score of −10.393 kcal/mol with minimum inhibitory activity at 25μg/ml [31,33].

The M. tuberculosis Serine/Threonine (S/T) protein kinases MtPknA-L, except PknC, are considered as
potential anti-TB drug targets [34]. The M. tuberculosis S/T protein kinases, belong to the Pkn2 family, are
eukaryote-like S/T kinases and function in phosphorylation-dependent signal transduction manner by
transducing extracellular stimuli into a cellular response to ensure M. tuberculosis cell division, transcription
regulation, stress response, regulation of multiple metabolic processes and pathogenesis [34,35]. These Mt
kinases can be categorized based on their similarity into five clades; Clade-1 (PknA, PknB, PknL), Clade-2
(PknH, PknE, PknD), Clade-3 (PknF, PknL, PknJ), Clade-4 (PknK) and Clade-5 (PknG) [36]. MtPknB and
MtPknE have been implicated in important biological roles in M. tuberculosis which include the determination
of cell shape, morphology, cell division and biofilm formation for PknB [37,38], and inhibition of apoptosis in
infected macrophages for PknE [39,40]. Thus, both PknB and PknE are considered as important molecular
targets for effective anti-TB drug search [34,41,42]. The MtPknB and MtPknE crystal structures were reported
previously and their structures have been used successfully as molecular targets for inhibitors screening and
drug discovery [43-45]. Here in, we applied bioactivity-guided screening of acetyl-eugenol isolated from A.
ilotica (L.) leaves against various pathogenic bacterial isolates and performed molecular docking studies using
MtPknB and MtPknE as targets. Based on the preliminary results of isolated acetyl-eugenol, we designed and in silico synthesised phenyl and indole derivatives using acetyl-eugenol as a chemical scaffold and studied them
with docking experiments against the same targets.

2. Materials and Methods
2.1 Plant collection and authentication

Fresh disease-free leaves of *A. nilotica* (L.) were separately collected from Bodinga, Sokoto State, Nigeria. The specimens were identified and authenticated by Botanists at the Department of Biological Sciences, Usmanu Danfodiyo University, Sokoto, Nigeria. The specimens identified as *A. nilotica* (L.) has given the voucher number UDUH/ANS/0247. Specimens were then shed-dried, ground finely and kept in air-tight containers until further use.

2.2 Preparation of ethyl acetate extracts

Crude methanolic extraction was initially performed followed by solvent partitioning on the residue (marc) to obtain the ethyl acetate extract. In brief, the methanol extraction was carried out by soaking 1 kg of the specimen in 6 L of 90% methanol for 72 h and was filtered using a clean cloth and Whatman No. 1 filter paper. The filtrate was then concentrated *in vacuo* at 30°C and stored in sterile containers at 4°C until further use. For the solvent partitioning, about 100 g of the crude extract of *A. nilotica* was dissolved in 500 ml sterile distilled water in a separatory funnel and extracted with *n*-hexane. The resulting *n*-hexane phase was concentrated to dryness and the powder was kept in a freezer in an air-tight container. The aqueous phase was further extracted with ethyl acetate. The fraction obtained from the ethyl acetate extraction was concentrated to dryness and the powder was kept at 4°C until used. The same procedure was applied to obtain the *n*-butanol fraction.

2.3 Isolation and purification of the bioactive compounds

Thin layer chromatography (TLC), column chromatography (CC) and preparative TLC techniques were used for the isolation and purification of the bioactive compounds as described previously [46,47].

2.4 Thin Layer Chromatography

TLC was carried out on aluminum TLC sheets pre-coated with silica gel (60 PF254) with a layer thickness of ~0.2 mm. The one-way ascending spotting and development technique was followed. Spots were applied manually on silica gel aluminum plates using a capillary tube and dried using an air blower. The TLC was developed at a room temperature using a Shandon chromatographic tank covered with a lid during the run. The solvent system contained a mixture of *n*-hexane and ethyl acetate solvent with a ratio of 70:30 (see Figure S1).

2.5 Column Chromatography

The ethyl acetate extract was loaded on a stationary phase containing silica gel (60-120 mesh size) using the wet loading method in a glass column (75 × 3.5 cm dimensions) packed with a wet slurry silica gel [46]. The extract was first dissolved in a small amount of *n*-hexane mixed with a small quantity of silica gel and then loaded on top of the pre-packed column. Different solvent systems were used to elute the silica gel column by gradient elution method— i) 100% *n*-hexane, ii) *n*-hexane/ethyl acetate mixtures (5:95, 10:90, 15:85, 20:80, 25:75, 30:70, 35:65, 40:60, 45:55, 50:50, 55:45; 60:40, 65:35, 70:30, 75:25, 80:20, 85:15, 90:10, 95:5), iii) 100% ethyl acetate, or iv) ethyl acetate/methanol mixtures with similar ratios to that of *n*-hexane/ethyl acetate. The best solvent system from which the fraction was eluted is *n*-hexane/ethyl acetate (75:25), which was confirmed by TLC using *n*-hexane/ethyl acetate solvent with a ratio of 70:30.

Theethyl acetate extract—the most effective in the anti-bacterial sensitivity assays—was further subjected to CC fractionation as described above. In brief, 3.0 g of the extract was chromatographed on a silica gel column eluted with a mobile phase containing: i) 100% *n*-hexane, ii) *n*-hexane/ethyl acetate mixtures (90:10, 80:20; 70:30, 60:40, 50:50, 40:60, 30:70, 20:80, 10:90), iii) 100% ethyl acetate, or iv) ethyl acetate/methanol mixtures (98:2, 95:5, 90:10, 80:20), which were gave rise to a total of 110 fractions. These 110 fractions were pooled together based on similarity in their TLC profile and their retention factors (Rf), calculated according to equation (1), to give rise into 5 sub-fractions. The sub-fractions were then subjected to TLC and a single spot was observed for each fraction confirming the similarity and assuring the pooling of sub-fractions. Sub-fractions were dried, weighed and labelled before used for bioactivity assay.
2.6 Structural elucidation of the bioactive compounds

The structures of the bioactive compounds based on the antibacterial assays were elucidated and characterized by different spectroscopic and spectrometry methods including Gas chromatography/mass spectrometry (GC/MS), nuclear magnetic resonance (NMR), Fourier Transform-Infrared spectroscopy (FT-IR) and UV-visible spectrophotometry.

2.7 Gas Chromatography/Mass Spectrometry analysis

For GC/MS analysis, 0.5 mg of each pure compound was placed in a labelled vial and diluted with an appropriate solvent of HPLC grade up to a final concentration of 1.0 μM. The solution in each case was filtered to remove any insoluble material before injecting into the column. An aliquot of 2.0 ml of each sample solution was then injected into a GC/mass spectrometer equipped with a GC-MSD detector (Agilent GC 7890B, MSD 5977A, Agilent Technologies, USA) and ran for 27 min. The column oven and injection temperature were set at 80°C and 250°C, respectively, and the column pressure was maintained at 108 kPa during the run. The spectra were analyzed using the program GCMS-QP2010 PLUS.

2.8 Nuclear Magnetic Resonance spectroscopy

Proton (1H) and carbon-13 isotope (13C) NMR experiments alongside two-dimensional (2D) NMR spectroscopy including correlation spectroscopy (COSY), nuclear Overhauser effect Spectroscopy (NOESY), heteronuclear single-quantum correlation spectroscopy (HSQC), and heteronuclear multiple-bond correlation spectroscopy (HMBC) were employed to get structural information on the isolated compounds. For each sample, 10 mg of the pure compound was dissolved in 1.0 ml of 100% dimethyl sulfoxide (DMSO, Sigma-Aldrich) as a solvent and filtered through a Pasteur pipette equipped with a glass wool plug that was discharged into a labelled 5.0 mm NMR tube. The filtration serves to remove undissolved residues and dust from the solution that affects the resolution and line shape of NMR spectra. NMR spectra were recorded on 400 MHz Agilent-VNMRS-NMR spectrometer (Agilent Technologies, USA). Chemical shift values were reported in parts per million (δ-ppm). All spectra were analyzed and the results obtained were compared with published data in order to elucidate the structures of the isolated compounds.

2.9 Bioactivity antibacterial sensitivity assays

2.9.1 Microorganisms, culture media and inoculum preparations

The microorganisms used in the present study were locally isolated and identified by the Department of Microbiology, Faculty of Science, Kebbi State University of Science and technology, Aliero, Nigeria. All isolates were purified and maintained in slant cultures of Mueller Hinton Agar (MHA). These isolates are *Escherichia coli*, *Klebsiella pneumoniae*, *Proteus* sp., *Pseudomonas aeruginosa*, *Staphylococcus aureus*, *Salmonella typhi* and *Streptococcus pneumonia*. The sensitivity tests were performed on either Mueller Hinton Agar or Mueller Hinton Broth (MHB), which were all prepared freshly under aseptic condition.

To prepare the inoculum of each microorganism a 0.5 McFarland turbidity standard was used to standardize the microorganism. The scale was prepared by adding 50 μl of 1% BaCl2 to 9.95 ml of 1% H2SO4. The suspensions of the microorganisms were prepared in normal saline and compared with 0.5 McFarland turbidity standard against a white background with contrasting black lines in the presence of good lighting [48]. The suspensions were diluted with normal saline if the density is higher than that of the standard and additional microorganism suspension was added in case of lower density. This continues until the density of the suspension matches with that of 0.5 McFarland turbidity standard, which corresponds to ~ 1.5 × 10^{18} CFU/ml.

2.9.2 Antibacterial sensitivity assay of the extracts
Agar well diffusion method was used to test the antibacterial activity of column fractions as reported previously [49,50]. The assay was performed using stock concentrations of 10 mg/ml. The standardized inoculums of the isolates were uniformly inoculated on a freshly prepared MHA plates by streaking the swab extensively over the entire surface of the plates. A sterile Cork borer of 6.0 mm diameter was used to make two wells per plates and labelled appropriately. An aliquot of 20 μl of the extract was added in the well and allowed to diffuse into the agar and incubated at 37°C for 24 h. The zone of inhibition diameter was measured in millimeter (mm) as an indication of antibacterial activity of the extract against the bacterial isolate. The antibiotic amoxicillin was used as a positive control with a final concentration of 1.0 mg/ml.

2.9.3 Minimum inhibitory concentration (MIC) and Minimum bactericidal concentration (MBC)

The MIC determination was performed on the microorganisms that were sensitive to the fractions using the broth dilution method [51]. Different concentrations of the extracts were prepared and added to test tubes containing MHB. Bacterial isolates were inoculated into each tube and incubated at 37°C for 24 h. The minimum concentrations of the extract which showed no turbidity in the MHB was considered as the MIC.

The MBC was performed to test the viability of microorganisms upon treatment with the compound. The MIC cultures of different concentrations were inoculated on MHA plates and incubated at 37°C for 24 h. After incubation the plates were observed for the growth of bacterial colonies and data were recorded. The MBC represents the MHA plate with a minimum concentration of the extract that displays no bacterial growth [52].

2.10 Molecular docking experiment

2.10.1 Ligands and protein preparations

The 3D structure of acetyleugenol was obtained from the PubChem database (www.pubchem.ncbi.nlm.nih.gov). Other eugenol derivatives—a total of four molecules containing eugenol scaffold—were sketched and converted into 3D structures using ChemDraw (ChemDraw 2017 PerkinElmer Informatics Inc.). All molecules were prepared using the Molecular Operating Environment (MOE) program (www.chemcomp.com, Chemical computing group, Montreal, Canada). Partial charges were added to the compounds using MMFF94 force field before subjecting them to energy minimization to obtain optimized geometries.

For the protein targets preparation, X-ray crystal structures of the M. tuberculosis Serine/Threonine kinases enzymes MtPknE and MtPknB with PDB codes 2H34 and 2FUM, respectively, were retrieved from the protein databank (www.rcsb.org). The structural models were prepared by removing water molecules, adding hydrogen atoms, and applying energy minimization. The binding sites of the MtPknE and MtPknB were predicted using homology modelling. Target protein sequences were submitted to the SWISS-Model server and homolog templates were obtained by BLAST. The template selected for the next experiments was the plant Cipk family of protein kinases (PDB ID: 4CZU). Target proteins and templates were superimposed in the MOE software and the binding sites were searched for using the Site-finder function of the MOE. We purposely selected the binding sites that coincide with the co-crystallized ligand of the template proteins.

2.10.2 Docking procedure

The docking experiments were performed using the docking suite of the MOE software. The compound structures were transferred and saved in MOE database. A Gaussian contact surface was drawn around the binding sites of the target protein and the receptor was verified as a receptor and the site as ligand atoms. The Placement method used was Triangle Matcher. The first scoring function was set to London dG/MOE and the refinement to force field. The docking process was then started by retaining 100 poses. The final refined poses were ranked by the MM/GBV1 binding free energy estimation. The depiction of results was generated by PyMOL APBS tools and MOE.

3. Results and Discussion

3.1 Isolation and structural elucidation of Eugenol and its antibacterial activity
A. nilotica (L.) is a medicinal plant that is well-known for its applications in the traditional medicines and has been used extensively to treat various respiratory diseases and other infectious diseases [19-21]. Here, we applied bioactivity guided approach to justify the wide-spread application of A. nilotica (L.) by conducting systematic phytochemical analysis on the leaves and tested its antimicrobial activity against several pathogenic bacteria including E. coli, K. pneumoniae, Proteus sp., P. aeruginosa, S. aureus, S. typhi and S. pneumonia (Figure S1). First, we extracted the chemical compounds from the leaves using methanol and used the methanol extract and marc for further fractionation with n-hexane, ethyl acetate and n-butanol consecutively (Scheme S1). The ethyl acetate extract was found to be the most effective extract against these bacterial isolates; hence it was selected for further phytochemical analysis in order to isolate the active compound(s). TLC and column chromatography were subsequently applied to fractionate A. nilotica (L.) leaves extracts by using solvent systems containing various ratios of n-hexane, methanol and ethyl acetate. Fractions were pooled and categorized based on their similarity in colors and total phytochemicals in the different parts of A. nilotica (L.) such as flavonoids, alkaloids, terpenes and others active ingredients [19,22,23]. To elucidate the chemical structure and properties of the active ingredient in fraction I; GC-MS, NMR spectroscopy, FTIR and UV-visible spectroscopy were applied. The GC-MS spectrum indicated that the most prominent compound in fraction I is acetylneugenol with the formula C₁₃H₂₀O₈—an eugenol derivative (Figure S2). This is the first report in the purification of acetylneugenol, to our knowledge, from the leaves of A. nilotica (L.). ¹H NMR, ¹³C NMR and 2D NMR analyses confirmed the structure of fraction I—acetylneugenol (Figure S3, S4 and S5). The ¹³C NMR spectral data of fraction I was compared with a reference ¹³C NMR spectrum of pure acetylneugenol compound [53], as well as theoretically predicted ¹³C NMR chemical shifts for the acetylneugenol (b-ppm) [54]. These comparisons revealed similar chemical shifts for the 12 carbon positions in the backbone of acetylneugenol structure (Table 3) [53,54]. In light of these results along with the GC-MS, we concluded that the major compound of fraction I is an acetylneugenol. Fraction I was homogenous and showed maximum absorption peak (λmax) at 365 nm (Figure S6) and its FTIR spectrum indicated the presence of characteristic peaks of well documented eugenol compounds (Figure S7) [55]. Previous results on

Table 1. Antibacterial activity of pure fractions (I-V) from the ethyl acetate extraction of the A. nilotica (L.) leaves.

<table>
<thead>
<tr>
<th>Bacterial isolates</th>
<th>Fraction-I</th>
<th>Fraction-II</th>
<th>Fraction-III</th>
<th>Fraction-IV</th>
<th>Fraction-V</th>
<th>Methanol 5%</th>
<th>Amox.</th>
</tr>
</thead>
<tbody>
<tr>
<td>E. coli</td>
<td>22.0±3.00</td>
<td>5.33±1.52</td>
<td>n.d</td>
<td>4.33±2.52</td>
<td>n.d</td>
<td>28.7±1.15</td>
<td></td>
</tr>
<tr>
<td>K. pneumoniae</td>
<td>15.0±4.58</td>
<td>5.67±2.31</td>
<td>n.d</td>
<td>5.33±1.53</td>
<td>n.d</td>
<td>26.7±0.58</td>
<td></td>
</tr>
<tr>
<td>Proteus sp.</td>
<td>29.0±2.00</td>
<td>4.67±2.08</td>
<td>n.d</td>
<td>3.33±1.53</td>
<td>n.d</td>
<td>30.7±1.15</td>
<td></td>
</tr>
<tr>
<td>P. aeruginosa</td>
<td>24.7±3.51</td>
<td>7.67±3.06</td>
<td>n.d</td>
<td>4.33±1.15</td>
<td>n.d</td>
<td>25.4±1.53</td>
<td></td>
</tr>
<tr>
<td>S. aureus</td>
<td>24.3±4.16</td>
<td>6.00±1.73</td>
<td>n.d</td>
<td>3.61±2.08</td>
<td>n.d</td>
<td>23.0±1.00</td>
<td></td>
</tr>
<tr>
<td>S. typhi</td>
<td>28.0±5.57</td>
<td>2.00±1.00</td>
<td>n.d</td>
<td>3.00±2.00</td>
<td>n.d</td>
<td>28.0±2.00</td>
<td></td>
</tr>
<tr>
<td>S. pneumonia</td>
<td>26.7±1.53</td>
<td>5.33±1.53</td>
<td>n.d</td>
<td>3.33±1.52</td>
<td>n.d</td>
<td>21.7±1.53</td>
<td></td>
</tr>
</tbody>
</table>

Amox. = Amoxicillin antibiotic (positive control). 5% Methanol = negative control. n.d = no detected activity. Data presented as mean ± std (n=3).
eugenol isolated from other plants indicated significant molecular interactions with Extended Spectrum Beta-Lactamase enzymes (ESBL) and showed significant antibacterial activity against *E. coli* and *K. pneumoniae* [56]. This might explain the strong susceptibility of these bacterial isolates to the acetyeleugenol (Table 1,2). Eugenol oil, which contains about 10% acetyeleugenol, has also been shown to exhibit a strong antimicrobial activity against *E. coli*, *S. aureus*, and *Candida albicans* and less activity against *Enterococcus fæcalis* [57]. These studies clearly indicate that acetyeleugenol may have broad spectrum antimicrobial activity. Thus, it is important to explore its potential with different drug targets to understand its mechanism of actions. Such knowledge can help to develop novel compounds based on the acetyeleugenol or eugenol scaffolds with stronger antimicrobial potential.

**Table 2.** The Minimum inhibitory concentration (MIC) and the minimal bactericidal concentration (MBC) of fraction I acetyeleugenol.

<table>
<thead>
<tr>
<th>Bacterial isolates</th>
<th>MIC (mg/ml)</th>
<th>MBC (mg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>E. coli</em></td>
<td>0.31</td>
<td>5.00</td>
</tr>
<tr>
<td><em>K. pneumoniae</em></td>
<td>1.25</td>
<td>10.0</td>
</tr>
<tr>
<td><em>Proteus sp.</em></td>
<td>0.63</td>
<td>5.00</td>
</tr>
<tr>
<td><em>P. aeruginosa</em></td>
<td>1.25</td>
<td>5.00</td>
</tr>
<tr>
<td><em>S. aureus</em></td>
<td>0.63</td>
<td>10.0</td>
</tr>
<tr>
<td><em>S. typhi</em></td>
<td>1.25</td>
<td>n.d</td>
</tr>
<tr>
<td><em>S. pneumonia</em></td>
<td>2.50</td>
<td>n.d</td>
</tr>
</tbody>
</table>

3.2 Molecular Docking of Acetyeleugenol and its analogues against MtPknB and MtPknE

To study the effect of acetyeleugenol at a molecular level and explore its antimicrobial potential, two important *M. tuberculosis* drug targets were chosen; i) *MtPknB* and ii) *MtPknE S/T* protein kinases, which play pivotal roles in *M. tuberculosis* development and pathogenesis [34,35]. First, we used the acetyeleugenol and conducted molecular docking experiments against the *MtPknE* protein kinase (PDB ID: 2H34). The results indicated relatively moderate docking score with a binding energy at —8.53 kcal/mol (Figure 1). The docking revealed that the acetyeleugenol interacts with *Ile96* residue of *MtPknE* via single hydrogen bond with 2.8 Å which is formed between the carbonyl oxygen of the ester group of acetyeleugenol and the hydrogen of the backbone NH group of *Ile96* residue (Figure 1a). The two alkyl groups of the acetyeleugenol were found to be embedded in a hydrophobic pocket that is formed by several hydrophobic residues including *Leu95, Ile96, Ala43, Val22, Val30, Val77, Val156, Phe158, Leu146*, and Ile160 (Figure 1b). The vinyl group (*—CH=CH2*) of the acetyeleugenol is located in a solvent accessible region of the binding cavity and show no interaction with any of the residues in this cavity. This result indicates that the active functional group of the acetyeleugenol in the interaction with *MtPknE* is the ester carbonyl group. A previous docking study on eugenol showed that both phenolic hydroxyl and methoxy oxygen of eugenol can interact with residues on the active site as demonstrated in the case of *C. albicans* squalene epoxidase [58]. Thus, the acetyeleugenol has smaller size relative to the binding cavity of the *MtPknE* kinases, which might potentially affect its orientation and binding. This might explains the relatively low binding energy and docking score.

**Table 3.** Comparison of the $^{13}$C NMR spectral data of fraction 1 (acetyeleugenol) obtained from the ethyl acetate extract fractionation of *A. nilotica* (L.) with literature and theory.

<table>
<thead>
<tr>
<th>Carbon position</th>
<th>$^{13}$C-NMR (δ-ppm)</th>
<th>Literature $^1$</th>
<th>Theoretical $^2$</th>
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<tbody>
<tr>
<td></td>
<td>Experimental</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C1</td>
<td>138.83</td>
<td>139.05</td>
<td>138.2</td>
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<tr>
<td>C2</td>
<td>166.78</td>
<td>150.88</td>
<td>151.0</td>
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<td>C3</td>
<td>52.06</td>
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<td>C4</td>
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<tr>
<td>C5</td>
<td>108.83</td>
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</tr>
</tbody>
</table>
Figure 1. Molecular docking of the eugenol acetate showing (a) the binding interactions in the binding site of MtPknE kinase. The eugenol acetate shown as stick (cyan), and (b) the 2D representation of the docking generated by MOE software; the amino acid residues are colored according to their chemical nature shown in the key. The 2D views shows the hydrogen bonding network and the Van der Waal contacts with the hydrophobic residues (green).

Table 4. Molecular docking binding free energy of the acetyleugenol compound and its phenyl and indole analogues against MtPknB and MtPknE molecular targets.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Binding Energy (kcal/mol)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MtPknE</td>
<td>MtPknB</td>
</tr>
<tr>
<td>EUG-1</td>
<td>–11.08</td>
<td>–10.46</td>
</tr>
<tr>
<td>EUG-2</td>
<td>–10.29</td>
<td>–10.19</td>
</tr>
<tr>
<td>EUG-3</td>
<td>–10.05</td>
<td>–9.64</td>
</tr>
<tr>
<td>EUG-4</td>
<td>–9.25</td>
<td>–9.35</td>
</tr>
<tr>
<td>Acetyleugenol</td>
<td>–8.53</td>
<td>–7.60</td>
</tr>
<tr>
<td>Mitoxantrone</td>
<td>n.a*</td>
<td>–10.8</td>
</tr>
</tbody>
</table>

*n.a = not available

In order to optimize the molecular interactions of the acetyleugenol for best orientation and binding in the hydrophobic binding pocket of MtPknE, we performed in silico synthesis for new compounds based on acetyleugenol skeleton modifications by adding either phenyl ring or indole to the ester moiety or the vinyl sites. This gave rise to five acetyleugenol analogs including the acetyleugenol and named compounds 1-5 as depicted in Figure 2. The compounds 1 and 3, which were named EUG-1 and EUG-3, respectively, are indole...
analogs with indole group at either the vinyl double bond site (EUG-1) or at the ester site (EUG-3) and similarly the phenyl group was added to make compound 2 (EUG-2) and compound 4 (EUG-4). This is the first report, to our knowledge, to design and study the docking profile of these newly proposed acetyleugenol analogs.

To get insights into the mechanisms of interactions and docking based structure activity relationships of these newly designed compounds *M. tuberculosis* protein Ser/Thr kinases *MtPknB* and *MtPknE* were used as molecular targets [34]. The binding energy values of the docking experiments are listed in Table 4. The docking results of these compounds appear to be similar between *MtPknB* and *MtPknE* with slightly higher selectivity to *MtPknE* than *MtPknB*. The most selective ligand, ligand-protein affinity, is the EUG-1 which has the lowest binding energy — the best interaction and docking score. The results indicate that the addition of an indole group to the double bond of the vinyl site has significantly improved the docking score in comparison with the acetyleugenol (Table 4). This introduction of an indole ring to the acetyleugenol skeleton in the case of EUG-1 provides an additional hydrogen bond donor via the indole amino group to an existing hydrogen bond acceptor of the ester carbonyl group. The EUG-1 also provides a molecular bulk necessary for Van der Waals and hydrophobic interactions, which further favored the EUG-1 binding energetics and improved the docking scores with both *MtPknE* and *MtPknB* (Figure 3). The EUG-1 also reveals similar binding energies to *MtPknE* and *MtPknB* with −11.08 kcal/mol and −10.46 kcal/mol, respectively (Table 4). A recent study by Ali et al., (2018) has examined several propolis compounds as potential anti-TB drugs including flavonoids pachypodol and pinobanksin-3-(E)-caffeate, which showed lower affinity with docking scores at −9.1 kcal/mol binding energy against the *MtPknB* comparison to the control inhibitor mitoxantrone, which exhibited a better docking score with −10.8 kcal/mol binding energy [42]. When the same propolis were examined against other *M. tuberculosis* enzymes involving *MtPank*, *MtDprE1* and *MtKasA*, the results indicate similar or better docking scores than that of mitoxantrone [42]. This is a clear indication that the indole analogue EUG-1 of the acetyleugenol is a potential anti-TB that can act against both *MtPknE* and *MtPknB* kinases. Another compound named IMB-YH-8 has also provided promising clinical trials and was shown to act specifically against the *MtPkn* enzymes with the most effect against the *MtPknB* [59].

![Chemical structures](image)

**Figure 2.** The acetyleugenol (5) from *A. nilotica* (L.) and the *in silico* synthesized phenyl and indole analogs (this study).
The docking results indicate that the EUG-1 interacts via its indole amino group and the carbonyl oxygen to form energetically favorable binding affinity with the drug target. It forms three hydrogen bonds with MtPkne; the first with NH group and the side chain of Asp100 with 3.07Å bond length, while the remaining hydrogen bonds are formed between the carbonyl oxygen and the backbone of Leu95 and Ile96 with 3.31Å and 2.87Å bond length, respectively (Figure 3a). The EUG-1 interacts with MtPknB via a hydrogen bond that is formed between the indole ring of EUG-1 and Val95 with 2.67 Å bond length and another hydrogen bond between the EUG-1 carbonyl group with Phe19 with 3.41Å bond length. Furthermore, the EUG-1 also interacts with MtPknB via a relatively weak H-π interaction between its indole pyrrole moiety and a hydrogen of Leu17 (Figure 3b).

Figure 3. Binding interactions of eugenol-indole analogs with MtPkne and MtPknB S/T kinases; a) EUG-1 with MtPkne, b) EUG-1 with MtPknB, c) EUG-3 with MtPkne and d) EUG-3 with MtPknB.

EUG-3, which has an indole group attached to the ester end of acetyleneugenol, reveals improved docking scores relative to the acetyleneugenol compound for both enzymes MtPkne and MtPknB with –10.29 kcal/mol and –10.19 kcal/mol binding energy, respectively (Table 1). These values are only slightly higher than that of mitoxantrone standard inhibitor for MtPknB that scores –10.8 kcal/mol with 0.8 μM IC50 [42]. Unlike EUG-1, the carbonyl group of EUG-3 doesn’t interact with any of the amino acid residues in the binding pocket owing most likely to the shielding provided by the bulky indole moiety that prevents the carbonyl from approaching any nearby residues as illustrated in the case of MtPkne (Figure 5c). EUG-3 interacts with MtPknB via one hydrogen bond that is formed between the amino group of the indole pyrrole and Val95. Additionally, we observed three weak H-π interactions among the EUG-3 aromatic benzyl and pyrrole rings and the enzyme.
Gly97, Leu17 and Met145 (Figure 5d). In contrast, the phenyl analogs EUG-2 and EUG-4, which have a phenyl ring attached either to the ester group site or to the double bond of the vinyl group of the acetylenogol skeleton, show weaker binding affinities for both enzymes comparing with that of the EUG-1 and EUG-3 (Table 1). EUG-2 has slightly better binding affinity comparing to EUG-4 with −10.05 kcal/mol binding energy for MtPknE and −9.64 kcal/mol for MtPknB. Altogether, all of the modifications applied in the present study have improved the binding affinity and docking score when compared with our starting compound acetylenogol. Generally, replacing the methyl group of the ester moiety by indole rather than phenyl group has robustly influenced the binding energetics and interactions with the hydrophobic pocket of the enzymes. On the other hand, the choice of the site of substitution has also played a paramount role in the improvement of the binding affinity, hence the docking scores especially with the MtPknE target. Therefore, these compounds are more likely to be good leads for the discovery of potential anti-tuberculosis drugs.

4. Conclusions

Here, we isolated and elucidated the molecular structure of acetylenogol from the leaves of A. nilotica (L.) and confirmed its antibacterial potential against seven clinical isolates. We further demonstrated for the first time the potential anti-TB of the acetylenogol analogs which were designed, in silico synthesized and used for molecular docking experiments against M. tuberculosis Ser/Thr protein kinases—the MtPknE and MtPknB. Our docking results indicate that the indole analogs EUG-1 and EUG-3 are potential inhibitors targeting these kinases with promising outcome. The effective interactions of EUG-1 and EUG-3 with both targets suggest a dual mechanism of actions for these compounds and proposing a possible interference with the pathogenesis and cell division of M. tuberculosis at different stages based on the functions of PknE and PknB in M. tuberculosis development. This urgently require chemical synthesis and more comprehensive in vivo and in vitro experiments to understand their mechanism of actions against M. tuberculosis or other mycobacterial models such as M. smegmatis to enable designing an effective anti-TB drug. The present study also encourages for further investigation of other acetylenogol derivatives against these kinases or other potential anti-TB targets.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Scheme 1: extraction and fractionation of the methanol crude extract of the A. nilotica leaves, Figure S1: phytochemical and antibacterial screening, Figure S2: GC-MS analysis of acetylenogol, Figure S3: 13C NMR spectrum of acetylenogol, Figure S4: 1H NMR and the DEPT experiments of acetylenogol, Figure S5: 2D COSY correlation of acetylenogol, Figure S6: UV-visible spectrum of acetylenogol, Figure S7: FT-IR analysis of the acetylenogol, Table S1: thin layer chromatography profile.

Author Contributions: Conceptualization, F.H.M.K. and A.M.I; proposed and designed compounds, F.H.M.K.; conducting experiments, A.A., T.A.A., and A.E.A.; writing—original draft preparation, F.H.M.K.; writing—review and editing, A.A., T.A.A., A.M.I, and F.H.M.K. All authors have read and agreed to the published version of the manuscript.

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References


