In silico Targeting, inhibition and analysis of polyketide synthase enzyme in Aspergillus ssp

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A B S T R A C T
Aflatoxins are toxic and carcinogenic components produced by some Aspergillus species such as Aspergillus flavus. Polyketide synthases enzyme (PKS) plays a central role in aflatoxin s biosynthesis of in Aspergillus flavus, especially the product template (PT) domain, which controls the aldol cyclization of the polyketide forerunner during the biosynthesis of the aflatoxin pathway process. Here, we apply the in silico approaches to validate 623 natural components obtained from the South African Natural Compound Database (SANCDB), to distinguish the PT domain s prospected inhibitors. From the 623 compounds, docking results showed that there are 330 different compounds with energy binding lower than the natural substrate (palmitic acid or PLM) of the Product Templet domain (PT). Three factors were selected to determine the best 10 inhibiting components: 1) energy binding, 2) the strengthen chemical interactions, 3) the drug-likeness. The top ten inhibiting components are kraussianone 6, kraussianone 1, neodiospyrin, clionamine D, bromotopsentin, isodiospyrin, spongotine A, kraussianone 3, 14β-Hydroxy bufa-3,5,20,22-tetraenolide and kraussianone 7. The chemical interactions between 3HRQ domain and the natural substrate in the active site amino acids are highly similar to the 3HRQ with the top ten components, but the main differences are in the binding energy which is the best in the top ten ligands. Those ten components give successful inhibition with PT domain which will lead to the formula to be used for inhibition and control aflatoxin contamination of agriculture crop yields and lessen the degree of harming and sicknesses that are coming about because of acquiring measures of aflatoxin.

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1. Introduction

Aflatoxins are a family of toxins produced by some species of Aspergillus fungi, mainly Aspergillus flavus, Aspergillus parasiticus, and Aspergillus nomius. They contaminate many crops, such as tree nuts, wheat, maize, cottonseed, and peanuts (Williams et al., 2004). Exposure to aflatoxin causes many diseases such as vomiting, abdominal pain, nausea, convulsions acutely, hepatotoxicity, teratogenicity, hepatocellular carcinoma, and immunotoxicity, according to (Aayush and Evelyn, 2020). The DNA information of Aspergillus is organized in 8 chromosomes, including the 54th cluster of 30 genes that are responsible for aflatoxin production and are regulated by aflR and aflS transcription factors. Aflatoxins are considered polyketide-derivate, which demand hexanolate units to
biosynthesis pathway’s backbone in the early stages (Usha et al., 2017). There were previous strategies to inhibit aflatoxin production such as gene silencing and transgenic plants. (Jonathan and Guy, 2018) illustrated that RNAi was constructed against polyketide synthase to produce transgenic maize plants, resulting in a 93% reduction of aflatoxin production. This technique cannot be applicable in many crops such as maize and wheat because it may have a harmful effect on humans and animals and undergo many experiments. Also, (Jonathan and Guy, 2018) illustrated a different technique of spreading some Aspergillus strains to the soil of plants, which are capable of eliminating aflatoxin producers. However, this technique cannot conserve against postharvest contagion during grain storage. Using natural components as aflatoxin inhibitors are more secure for human rather than using pesticides or chemicals. The non-toxic natural components have been used as a thriving source for producing medicines for humans and animals (Daniel et al., 2012). In this investigation, 623 unique components were examined by in silico docking for the first time for their ability to inhibit the production of aflatoxin. These components had isolated from some African plant species, perennial herb, and South African marine.

2. Materials and methods
2.1. Biological data, domain sequence, and 3D structure

The information on the biological functions and interactions of Aspergillus flavus genes and aflatoxin pathway were obtained from the Kyoto Encyclopedia of Genes and Genomes KEGG pathway.
database (http://www.genome.ad.jp/kegg). The polyketide synthase gene of *Aspergillus parasiticus* (AflC), and its functional enzyme (Norsolorinic acid synthase) (https://www.uniprot.org/uniprot/Q12053) was searched for Norsolorinic acid synthase structure and PT domain using the NCBI database and EMBL-EBI database (Figs. 1 and 2) (Jason et al., 2009). This enzyme belongs to polyketide synthases with EC number 2.3.1.221. As our target PT domain, we employed the X-ray crystallographic structure derived from *Aspergillus parasiticus*, with a 1.80 Å resolution (PDB ID: 3HRQ) and a sequence with 357 amino acids (UniProtKB ID: Q12053) (https://www.ebi.ac.uk/pdbe/entry/pdb/3HRQ). The hexanoyl starter unit in the PT domain was provided to the acyl-carrier protein (ACP) domain by a dedicated fungal fatty acid synthase (Fig. 3).

| Compound               | CAS No.      | Chemical structure | Source of isolation  |
|------------------------|--------------|--------------------|----------------------|
| Kraussianone 6         | 761456–86-4  |                    | Eriosema kraussianum |
| Kraussianone 1         | 497858–65-8  |                    | Eriosema kraussianum |
| Neodiospyrin           | 33916–25-5   |                    | Euclea natalensis    |
| Clionamine D           | 1042138–30-6 |                    | Cliona celata        |
| Bromotopsentin         | 112515–44-3  |                    | Topsentia genetrix   |
| Isodiospyrin           | 20175–84-2   |                    | Euclea natalensis    |
| Spongotine A           | 116747–40-1  |                    | Topsentia pachastrelloides |
| Kraussianone 3         | 497858–67-0  |                    | Eriosema kraussianum |
| 14β-Hydroxybufa-3,5,20,22-tetraenolide | 545–51-7 |                       | Urginea epigea |
| Kraussianone 7         | 761456–87-5  |                    | Eriosema kraussianum |
2.2. Ligand preparation

About 623 natural compounds were obtained from the South African Natural Compound Database (SANCDC). Energy minimization has been performed using Avogadro, an open-source molecular builder and visualization tool (Version 1.XX, http://avogadro.cc/), followed by ligand preparation using Open Babel (Version 2.3.1, http://openbabel.org) following the analysis pipeline described by (Patrick et al., 2019) for preparing small-molecule libraries. The Swiss ADME database (http://www.swissadme.ch/index.php) was used for validating the toxicity of the components. The CAS number ID and the chemical structures of the top ten selected compounds, based on their binding energies, chemical interactions, and drug-likeness are shown in Table 1.

2.3. Molecular docking

Initially, the protein structure was modified by removing chain B and water molecules, adding hydrogen atoms to protein and ligand, and metals were treated using the Discovery Studio software (version 2019). The structure file was saved in PDBQT format. The random setting was used for rudimentary positioning, orientation, and torsions of the ligand. Then, Auto Dock Vina (Version 2.0, http://www.vina.dock.com) was used for the grid box. Auto Grid program was utilized to generate affinity grid maps of 48 × 62 × 74 XYZ Å points and 1.00 Å spacing. After that, the polyketide synthase’s active sites are identified and docked with ligands and metals were treated using the Discovery Studio software (version 2019). The structure file was saved in PDBQT format. The results included binding energy and inhibition constants, which implicate chemical interactions, hydrogen bonds, and hydrophobic regions. LIGPLOT was utilized for analyzing the hydrophobic and H-bond interactions between the ligands and domain complexes. They were further visualized in 3D using PyMOL. Interactions complex of each ligand – 3HRQ domain were analyzed to define binding efficiency, according to (Shraddha et al., 2019).

3. Results and discussion

3.1. Drug-likeness and analogs properties of the candidate inhibitors

The early detection and filtering of all the candidates’ inhibitors that have toxic effect, disadvantageous constitutional and physico-chemical characteristics that may lead to disturbed bounce on multiple protein targets and indigent properties such as distribution, absorption, metabolism, toxicity, and excretion is a master agent in remission limpness rate in the medication improvement process as it had been described by (Amit et al., 2016). The suitable properties of the identified candidate inhibitors were determined. All components were assessed and checked for toxicity and drug-likeness employing SWISS-ADME (Thommas and Özlem, 2019). As exhibited in Table 2, all the selected hits had adequate molecular weight, number of H-bond donor, number of H-bond acceptors, molar refractionity, lipophilicity, water-solubility, pharmacokinetics, drug-likeness, medicinal chemistry, and colored zone.

3.2. Docking score

The previous in silico docking studies illustrated down-regulation in the expression levels of polyketide synthase A enzyme due to the inhibition by quercetin and directly causing depression of aflatoxin production and contamination levels

![Table 2: SWISS-ADME results: the molecular weight, number of H-bond donor, number of H-bond acceptor, Molar refractivity, Lipophilicity, Water Solubility, Pharmacokinetics, Drug-likeness, Medicinal Chemistry, and Colored Zone of each compound.](image-url)

| Molecule     | Molecular weight (g/mol) | No. of H-bond donors | No. of H-bond acceptors | Molar refractivity (cm³ mol⁻¹) | Lipophilicity (Log P or Log Po/w) | Water Solubility | Pharmacokinetics (GI absorption) | Drug-likeness | Medicinal Chemistry |
|--------------|--------------------------|----------------------|-------------------------|-------------------------------|----------------------------------|------------------|----------------------------------|---------------|---------------------|
| Kraussianone 6 | 436.45                   | 3                    | 7                       | 121.05                        | 3.85                             | Moderately soluble | High                             | Yes           | PAINS: 0 alert       |
| Kraussianone 1 | 418.44                   | 2                    | 6                       | 120.21                        | 3.99                             | Moderately soluble | High                             | Yes           | PAINS: 0 alert       |
| Neodiospyrin  | 374.34                   | 2                    | 6                       | 101.30                        | 2.53                             | Moderately soluble | High                             | Yes           | PAINS: 1 alert quinone_A |
| Clionamine D  | 401.54                   | 1                    | 5                       | 109.59                        | 2.81                             | Moderately soluble | High                             | Yes           | PAINS: 0 alert       |
| Bromotopsentin | 421.25                   | 4                    | 3                       | 107.33                        | 1.94                             | Moderately soluble | High                             | Yes           | PAINS: more_than_2_esters |
| Isodiospyrin  | 374.34                   | 2                    | 6                       | 101.46                        | 2.32                             | Moderately soluble | High                             | Yes           | PAINS: 1 alert quinone_A |
| Spongotine A  | 407.26                   | 3                    | 2                       | 113.76                        | 1.94                             | Moderately soluble | High                             | Yes           | PAINS: 0 alert       |
| Kraussianone 3 | 438.47                   | 4                    | 7                       | 123.50                        | 3.63                             | Moderately soluble | High                             | Yes           | PAINS: 0 alert       |
| 14β-Hydroxybufa-3,5,20,22-tetraenolide | 366.49                   | 1                    | 3                       | 107.75                        | 3.36                             | Moderately soluble | High                             | Yes           | PAINS: 0 alert       |
| Kraussianone 7 | 436.45                   | 4                    | 7                       | 122.99                        | 3.71                             | Moderately soluble | High                             | Yes           | PAINS: 0 alert       |

- **PAINS:** Patterson Anti-Narcotic Screener
- **Brenk:** Brenk index
- **Brenk:** 0 alert = no alert
- **Brenk:** 1 alert = alert
- **GI absorption:** gastrointestinal absorption
- **Water Solubility:** water solubility
- **Pharmacokinetics:** pharmacokinetics
- **Drug-likeness:** drug-likeness
- **Medicinal Chemistry:** medicinal chemistry
Sudharsan et al., 2019). So, in silico drug design parameters such as docking is very useful to discover potent inhibitors for aflatoxin, especially for polyketide synthase protein. The 3HRQ domain - interaction with the natural substrate (PLM) and the 623 compounds were analyzed. The binding energy scores of the 330 compounds were identified and showed powerful interactions with the plurality of the 3HRQ domain compared with PLM. So, ligands with its structures resort to setup more contacts with the receptor residues, leading to vigorous interaction binding energy (María et al., 2015).

The Structure view of the top ten compounds was shown in Fig. 4. From the obtained docking results, the docked natural substrate binding energy result was −7.3. In contrast, the docked kraussianone 6 component result was found to has the highest docking score of −11.1 and interacted with the 3HRQ domain by two conventional hydrogen bonds with ASN A: 1568 and ASP A: 1395 amino acids and other chemical interactions were Alkyl, Pi-Alkyl, Pi-Sigma and van der waals forces (Table 2). Other compounds showed different docking results ranged from −10.9 to −9.5. Thus, kraussianone 6 was a remarkable component. From the list of compounds in Table 3, kraussianone 6 shows high biological inhibition activity against fungal protein PKS 3HRQ domain and docking score of −11.1 and two conventional hydrogen bonds. Kraussianone 6, kraussianone 1, kraussianone 3, and kraussianone 7 were naturally isolated from the perennial herb *Eriosema krausianum*, neodiospyrin and isodiospyrin were naturally isolated from a dioecious African plant species *Euclea natalensis*, clionamine D was naturally isolated from *Cliona celata* (called the red boring sponge), bromotopsentin was naturally isolated from Mediterranean shallow-water sponges *Topsentia genetrix*, spongotine A was naturally separated from the intertidal South African marine *Topsentia pachastrelloides*, 14β-Hydroxybufa-3, 5, 20, 22-tetraenolide was naturally isolated from *Urginea epigea*. Those unique components had been extracted and identified recently from South Africa and were examined for the first time by in silico docking against PKS enzyme and aflatoxin production.

Table 3
Docking score for the top ten compounds and the number of conventional hydrogen bonds in each component.

| Name                | Binding energy | Number of Covalent hydrogen bond with 3HRQ domain |
|---------------------|----------------|--------------------------------------------------|
| Kraussianone 6      | −11.1          | 2                                                |
| Kraussianone 1      | −10.9          | 1                                                |
| Neodiospyrin       | −10.6          | 2                                                |
| Clionamine D       | −10.5          | 2                                                |
| Bromotopsentin     | −10.0          | 1                                                |
| Isodiospyrin       | −9.8           | 1                                                |
| Spongotine A       | −9.7           | 1                                                |
| Kraussianone 3      | −9.6           | 1                                                |
| 14β-Hydroxybufa-3, 5, 20, 22-tetraenolide | −9.6 | 1 |
| Kraussianone 7      | −9.5           | 2                                                |
| Substrate           | −7.3           | 2                                                |
Fig. 5. The 2D PKS (PT domain) – chemical interaction with the substrate and the top ten inhibitors. 1) The interaction with the substrate. 2) The interaction with the Kraussianone. 3) The interaction with the kraussianone 1. 4) The interaction with the neodiospyrin. 5) The interaction with the clionamine D. 6) The interaction with the bromotopsentin. 7) The interaction with the isodiospyrin. 8) The interaction with the spongote A. 9) The interaction with the kraussianone 3. 10) The interaction with the 14β-Hydroxybufa-3,5,20,22-tetraenolide. 11) The interaction with the Kraussianone 7.
3.3. Chemical interaction with 3HRQ domain

The computational assessment of the protein–ligand interaction of all of the ligands (natural components) with PKs’ 3HRQ domain was visualized using Discovery Studio software version 2019. It shows the 2D of the chemical interaction with substrate and inhibitors such as the number of conventional hydrogen bonds which ranged from two in the substrate, kraussianone 6, neodiospyrin, clionamine D, and kraussianone 7 and one in the others inhibitors (Hong-Lian et al., 2017). Other chemical interactions such as carbon-hydrogen bond, van der waals, Pi-sigma, Pi-sulfur, Alkyl, Pi-Alkyl, Pi-pi sigma, Pi-pi stacked, and Amide-pi stacked are shown in Fig. 5. The amino acids interact with palmitic acid (ligand) ASN A: 1568 and HIS A: 1345. The secondary structure between 3HRQ and the ligands supply the best possible association of both amino acid residues. Also, the amino acid residues that associate with kraussianone 6, kraussianone 1, neodiospyrin, clionamine D, bromotopsentin, isodiospyrin, spongotine A, kraussianone 3, 14β-Hydroxybufa-3,5,20,22-tetraenolide, and kraussianone 7 were ASN A: 1568, HIS A: 1345, and ASP A: 1395 and those interactions were by conventional hydrogen bond which is shown in Fig. 5.

Hydrophobic regions between PKS (PT domain) and the substrate and the top ten inhibitors were different from each inhibitor to the other and ranged from the brown color, which represents the hydrophobic regions, and the blue color, which represent the hydrophilic regions (Fig. 6). The number of hydrophobic atoms is significant in hydrophobic interactions in drug resolving due to the rising of the binding consanguinity between target drug mediators. The binding consanguinity and drug effectiveness correlated with hydrophobic interactions were advanced by the association at the hydrogen bonding site. On the other hand, water molecules’ subsistence in the hydrophobic locus is very important to make this region quite flexible. The augmentation in the number of hydrophobic atoms in the specific active core site of the drug-target complex will lead to an increase in the biological inhibition activity of the drug leadership (Rohan et al., 2010).

On the other hand, hydrogen bond interaction is shown in Fig. 7, which establishes more informatics of association’s structural view. The results ranged from the pink color, representing the H-donor, and green color representing the H-acceptor. The significance of hydrogen bonds, a target-drug complex’s binding alliance, was described previously (Rohan et al., 2010). It was clarified that

![Fig. 6. Hydrophobic Interaction between PKS (PT domain) and the substrate and the top ten inhibitors. 1) The interaction with the substrate. 2) The interaction with the kraussianone 6. 3) The interaction with the kraussianone 1. 4) The interaction with the nodiospyrin. 5) The interaction with the clionamine D. 6) The interaction with the bromotopsentin. 7) The interaction with the isodiospyrin. 8) The interaction with the spongotine A. 9) The interaction with the kraussianone 3. 10) The interaction with the 14β-Hydroxybufa-3,5,20,22-tetraenolide. 11) The interaction with the kraussianone 7.](image-url)
the domain-ligand complex interface increased the binding cogni-
tion of complex molecules and was upgraded for its hydrogen
bonds and the hydrophobic interactions. The enzyme’s standard
case changes, such as hydrophobic, hydrophilic, H-donor, and H-
acceptor, can lead to inactivation for this enzyme (Nicholls et al.,
2000).

3.4. 2D interaction map

The brief 2D interaction map was generated using the
LIGPLOT + program. The key residues, such as numbering according
to each protein’s catalytic domain protein–ligand complex and
protein-substrate complex interaction analysis, stabilize the
ligands inside the subsite pockets, performed on each complex
were determined (Sourav et al., 2019). The shared interactions
between the 3HRQ domain and substrate are highly similar to
the domain with the top ten ligands, but the main differences are
binding energy, which is the best in the top ten ligands. The com-
parison between the substrate and kraussianone 6 in the interac-
tions with the 3HRQ domain showed similarity in 11
interactions. Also, 11 interactions were shared with kraussianone
1, 12 interactions with neodiospyrin, 6 interactions with clion-
amine D, 10 interactions with bromotopsentin, 5 interactions with
isoroispyrin, 10 interactions with Spongotine A, 13 interactions
with Kraussianone 3, 6 interactions with 14β-Hydroxybufa-3,5,20
22-tetraenolide, and 14 interactions with Kraussianone 7 which
is shown in in red circles Fig. 8.

4. Conclusions

In conclusion, about 623 natural components were examined to
act as natural inhibitors for the 3HRQ domain to block the polyke-
Fig. 8. The interactions between the active site of the 3HRQ domain of PksA with the substrate and ten different ligands, kraussianone 6, kraussianone 1, neodiospyrin, clionamine D, bromotopsentin, isodiospyrin, spongotine A, kraussianone 3, 14β-Hydroxybufa-3,5,20,22-tetraenolide, and kraussianone 7 using LigPlot.
Fig. 8 (continued)
tide synthesis enzyme. Docking results showed that about 330 proved their ability to act as inhibitors. The 330 components were filtered to elicit the best ten components. The results showed kraussianone 6, kraussianone 1, neodiospyrin, clionamine D, bromotompsentin, isodiospyrin, spongotine A, kraussianone 3, 14β-Hydroxybufa-3,5,20,22-tetraenolide, and kraussianone 7 were the best depending on the lowest binding energy, the best in chemical interactions, and the best in drug-likeness. Those components would lead to the formula for inhibition and control aflatoxin contamination of agriculture crop yields and lessen the degree of harming and sicknesses.

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and materials**

All relevant data are within the manuscript.

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Authors’ contributions

The authors contributed to the work done in the manuscript as follows: ML conceived of the presented ideas, ML, SH, and OS developed the theory and performed the computations and conducted the experiments, presented the data in tables and graphs. MA, AZ, SH, and ME verified the analytical methods, revised the experimental design, guided the data analyses and interpretation and manuscript revision, and supervised the findings of this work. ML prepared the preliminary version of the manuscript. GO, IA, and IS contributed to the final version of the manuscript.

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