

#### **RESEARCH ARTICLE**

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# Prediction of Aflatoxin-B1 (AFB1) Molecular Mechanism Network and Interaction to Oncoproteins Growth Factor in *Hepatocellular Carcinoma*

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#### Abstract

Aflatoxin-B1 (AFB1) is a common contaminant for staple foods during the storage process. Chronic exposure to AFB1 is widely known to induce the development of hepatocellular carcinoma (HCC). However, there is a lack of understanding of AFBi role in HCC mechanism. This research aims to identify protein(s) in HCC that might interact with AFB1 and to predict the pathway effected by AFB1. Analyses were performed using bioinformatics tools. SMILES notation of AFB1 was submitted into Swiss Target Prediction. Interaction among predicted proteins were analyzed by using STRING. The 3D structure of target protein was constructed by homology modeling. Reverse docking was performed, and the result was ranked based on binding affinity score. Furthermore, protein interaction network was constructed and analyzed by using Cytoscape. Results showed that three protein groups were predicted as target of AFB1, such as kinases, phosphatases, and G protein-coupled receptor with probability of 46.7%, 20%, and 6.7%, respectively. Seven proteins of kinases were strongly related to HCC, including RAF1, MAPK1, MAPK3, AKT1, EGFR, GSK3B, and mTOR. Reverse docking considered the AKT1-AFB1 as the most potential complex with the lowest affinity score -10.2 kcal.mol<sup>-1</sup>. It has hydrophobic bonds in Trp80, Val270, Tyr272, Asp292, Thr211, Leu210, Leu264, and Lys268 residues, whereas hydrogen bond in Ser205 residues. Moreover, further analysis demonstrated that interaction of AKT1-AFB1 is related to the metastasis pathway in HCC mechanism.

Keywords: Cancer, Kinase Protein, Protein Interaction, Protein Pathway, Toxin

#### **INTRODUCTION**

Aflatoxin is one of secondary metabolites produced by Aspergillus flavus and Aspergillus parasiticus. In general, there are several types of aflatoxins, such as aflatoxin-B1 (AFB1), aflatoxin-B2 (AFB2), aflatoxin G1 (AFG1), and aflatoxin G2 (AFG2). Aspergillus flavus typically produces AFB1 and AFB2, while Aspergillus parasiticus produces all type of aflatoxins. Among those aflatoxins, AFB1 is considered as the most common type which contaminates the staple foods and agricultural product. This mycotoxin is highly produced within 25°-30°C of temperature and 85% of relative humidity during distribution and storage processes.1 The prolonged consumption of food contaminated by AFB1 is considered as one of the major factors for hepatocellular carcinoma (HCC) development.2

HCC is dedicated as a cancer with the highest mortality across the globe. It is also classified as the most common cancer in liver which is indicated by epithelial neoplasm in hepatocyte.<sup>3,4</sup> The highest occurrence of the disease is spread across Asia, Africa, and America. Recent studies reported that hepatic tumor is observed in mice exposed by AFB1.5 However, the role of AFB1 within this HCC mechanism remains unclear. Therefore, in this research, we aim to

identify and explore the protein(s) which might interact with AFB1 and to predict the pathway effected by AFB1. This is considered as an essential study which could potentially be developed as a foundation for further understanding of relation between AFB1 and HCC. In addition, the analysis of mechanism pathway is part of system biology which integrated to understand the system function in organism through protein-protein interaction network (PPIN).<sup>6</sup>

#### **MATERIALS AND METHODS**

#### **Collection of AFB1**

The Aflatoxin B1 (AFB1) molecule was obtained from the PubChem database (https://pubchem.ncbi.nlm.nih.gov/) with ID number CID186907. The notation of canonical simplified molecular-input line-entry system (SMILES) was copied, and its 3D structure was collected in SDF format. The minimization process was conducted by using the Open Babel plug-in in PyRx software (The Scripps Research Institute).<sup>7</sup> This process was essential for further analysis related to molecular docking.

#### **Target Protein Prediction**

The target protein of AFB1 was predicted by submitting the canonical SMILES

to the Swiss Target Prediction (http://www.swisstargetprediction.ch/). *Mus musculus* was selected as the organism target, which is defined as the representative research object. Furthermore, top three target class of protein were selected as potential target protein. All predicted target proteins which belong to each class of proteins were listed.

## Analysis of AFB1 Interaction to the Predicted Target Protein

The obtained target proteins candidates of AFB1 were used as input to the webserver Search Tool for the Retrieval of Interacting Genes/Proteins (STRING) (http://string-db.org). Collection was performed using multiple protein category with *Mus musculus* as an organism target. The result was analysed based on the KEGG pathway enrichment analysis by selecting hepatocellular carcinoma. Furthermore, data was linked and further analysed by using Cytoscape software (Institute for System Biology, USA). This process aimed to determine the protein interaction network (PIN) that involved in hepatocellular carcinoma (HCC) disease.

#### **Protein Modeling of Predicted Target Protein**

The proteins involved in HCC disease interactions were accessed through the Uniprot database (www.uniprot.org). Those amino acid sequences of each protein were retrieved in FASTA format for homology modelling process. Furthermore, it was submitted into SWISS MODEL (http://swissmodel.expasy.org) to get the 3D structure of each protein. The water molecules and other ligands were eliminated by using PyMOL so that the pure protein structure was obtained.

### Molecular Docking of AFB1 with Predicted Target Protein

The screening process by molecular docking was carried out between each predicted target protein with AFB1 on the specific site. The process was performed by using Autodock Vina on the PyRx software (The Scripps Research Institute). Furthermore, the binding affinity score and the position of each screening result were collected and compared. The results of molecular docking were then analyzed by using LigPlot

software to identify the chemical bonds of amino acid residues.<sup>8</sup>

#### Visualization of AFB1 and Selected Target Protein

The representative results of AFB1 with each of predicted target protein were visualized by using PyMol software. It aimed to determine the detailed structure of proteins. Moreover, it depicted the position interaction between AFB1 with the protein.

# Protein Interaction Network (PIN) Analysis of Predicted Target Protein of AFB1 in HCC Mechanism

Protein interaction network (PIN) which is constructed from predicted target protein of AFB1 was analysed by using topological analysis. It was performed by Network Analyzer application which is integrated into Cytoscape software (Institute for System Biology, USA). There were various parameters which could be selected for analysis. However, number of edges and betweenness parameters were prominent selected to identify the most essential protein in HCC mechanism. Moreover, predicted target protein of AFB1 was also subjected for analysis of protein interaction pathway in HCC mechanism.

#### **RESULTS**

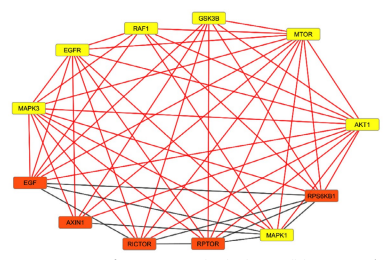
Aflatoxin B1 (AFB1) was retrieved from PubChem database with ID number ID186907. It described as a small molecule with molecular formula  $C_{17}H_{12}O_6$  and molecular weight 312.27 g/mol. The notation of canonical SMILES COC1=C2C3=C(C(=O)CC3)C(=O) OC2=C4C5C=COC5OC4=C1 was collected from database and utilized as a sample for analysis.

Protein target of AFB1 was predicted by submitting the notation of canonical SMILES into web server Swiss Target Prediction. The process was performed based on the ligand-based target prediction, whereas AFB1 as the center of query molecule. Gfeller et al., stated that predicted protein target is identified based on the similarity of another protein with its known ligand that resemblance to the query molecule. As it is shown in Table 1, there were three groups of protein which is predicted as a target of AFB1,

such as kinase, phosphatase, and G protein-coupled receptor with probability score 46.7%, 20%, and 6.7%, respectively. In detail, there are 22 kinase proteins predicted as the compatible target protein of AFB1. According to protein target prediction, it showed that AFB1 had several target proteins such as AKT/mTOR and MAPK.

The STRING analysis was conducted to identify the specific candidate of kinase protein for

AFB1. The result exhibited that 29 predicted AFB1 target proteins interacted to each other through 97 edges. However, there are only seven proteins which are closely involved in HCC mechanism, such as RAF1, MAPK1, MAPK3, AKT1, EGFR, GSK3B, and mTOR. Furthermore, protein interaction was constructed by using Cytoscape software. Seven proteins related to HCC mechanism were indicated by yellow-colored nodes and the interactions



**Figure 1.** Interaction of seven proteins related to hepatocellular carcinoma (HCC) based on Cytoscape software. Predicted target protein (yellow).

Table 1. The top 3 predicted target proteins group of AFB1 in Mus musculus

Group of protein	Target protein	Probability score
Kinase	MAPK1, GSK3B, DYRK1A, ABL1, RAF1, FLT1, KIT, JAK3,	
	RET, MAPK14, LCK, MAPK3, AURKB, LYN, MAPK9,	
	AKT1, AURKA, PRKDC, EGFR, ALK, CDK1, mTOR	46.7%
Phosphatase	PTPN1, CDC25B	20.0%
G protein- coupled receptor	TAAR1, TRH-R2, TRHR, ADORA2A, HTR2C, ADORA, ADORA2B	6.7%

**Table 2.** Homology modelling and 3D structure of protein related to hepatocellular carcinoma based on the SWISS MODEL

Target Protein	Uniprot ID	Template ID	Similarity score (%)	Validation score (%)
mTOR	Q9JLN9	6bcu.1.A	96,63	91,94
EGFR	Q01279	3gop.1.A	88,73	93,55
GSK3B	Q9WV60	1j1b.1.B	99,05	95,58
AKT1	P31750	6hhg.1.A	97,53	94,77
MAPK3	Q63844	4qtb.1.A	98,56	97,36
MAPK1	P63085	3zuv	100,0	94,49
RAF1	Q99N57	3omv.1.A	98,31	86,13

among proteins were exhibited by the grey-colored edges (Figure 1). However, the association among proteins do not necessarily mean that those proteins are directly binding each other. It merely

showed that those candidate proteins are strongly related each other of the HCC mechanism.

In brief, to understand the role of seven proteins related to HCC mechanism, the function

**Table 3.** List of binding affinity based on reverse docking between AFB1 and seven predicted target proteins in hepatocellular carcinoma

Ligand	Target Protein		Position of grid docking					
			Center (Å)			Dimension (Å)		
		X	Υ	Z	X	Υ	Z	
AFB1	AKT1	269.5	214.5	239.8	150.0	99.1	170.4	-10.2
	MAPK3	33.2	46.5	56.8	61.2	48.6	69.0	-9.3
	mTOR	269.5	214.5	239.8	150.0	99.1	170.4	-9.1
	GSK3B	27.7	-6.5	-38.0	76.3	60.2	60.2	-8.9
	MAPK1	21.4	-25.5	-9.0	63.1	67.6	80.0	-7.8
	EGFR	-9.0	-21.0	-37.8	59.4	72.0	74.3	-7.4
	RAF1	6.2	18.1	43.9	58.8	55.8	54.7	-6.7

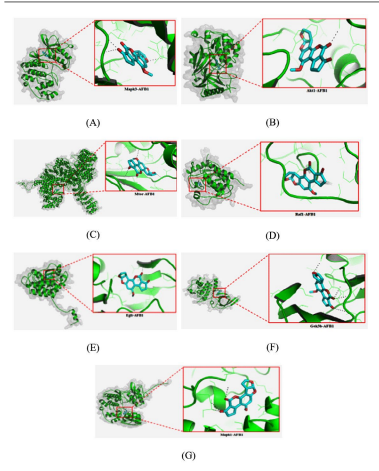


Figure 2. Interaction of AFB1 with several predicted protein: MAPK3 (A), AKT1 (B), mTOR (C), RAF1 (D), EGFR (E), GSK3B (F), and MAPK1 (G).

of each target protein is needed to be identified. Epidermal growth factor receptor (EGFR) belongs to receptor tyrosine kinase (RTK) which is found in plasma membrane, and it has a function to regulate the cell growth. <sup>10</sup> Generally, the downstream signaling cascade of EGFR includes two main pathways, such as RAS-RAF-MAP kinase pathway and PI3K-PTEN-Akt pathway. In cancer

**Table 4.** Number of edges of protein within protein interaction network in HCC mechanism

Protein	No. of edges	No. of	No. of
name		In-degree	Out-degree
AKT1	11	10	1
mTOR MAPK3	11 10	9	9
EGFR	9	2	7
EGF	9	3	6
MAPKAP1	9	7	2
RPS6KB1	9	5	4
RPTOR	8	3	5
RICTOR	8	8	0
RAF1	7	0	7
GSK3B	7		5
AXIN1	6	2	4

case, the level of EGFR is frequently elevated or mutated.

Furthermore, AFB1 is estimated to have an ability to interact with one of seven predicted proteins within HCC mechanism. Therefore, molecular docking would be conducted between AFB1 and those selected proteins. Whilst the 3D structures of each protein were constructed by homology modelling. The result of homology modelling showed that all of protein is considered valid according to the similarity score and validation score. The highest similarity score was exhibited by MAPK1, while the lowest score was showed by EGFR (Table 2). However, both similarity

**Table 5.** Top three betweenness score of protein interaction within protein interaction network in HCC mechanism

Protein name	Edge Betweenness score	
GSK3B – AXIN1	4.33	
GSK3B – EGF	4.00	
GSK3B - RPTOR	3.50	
GSK3B - RPS6KB1	2.50	

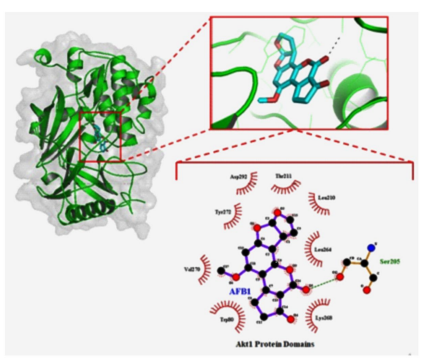


Figure 3. Interaction of amino acid residues within AKT1-AFB1 complex.

scores are still in the requirement range about 80-90% similar with the protein template.

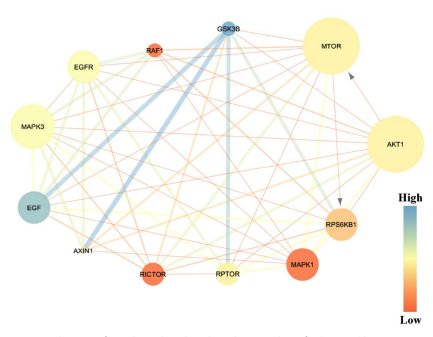
Seven proteins which are selected as predicted target protein of AFB1 and related to HCC pathway were selected to be identified about their interaction with AFB1. Each protein, such as RAF1, MAPK1, MAPK3, AKT1, EGFR, GSK3B, and mTOR was examined by reverse docking with AFB1. The molecular docking result showed that among seven proteins, complex of AKT1-AFB1, MAPK3-AFB1, and MTOR-AFB1 are demonstrated to have the lowest binding affinity about -10.2 kcal.mol<sup>-1</sup>, -9.3 kcal.mol<sup>-1</sup>, and -9.1 kcal.mol<sup>-1</sup>, respectively. However, the lowest binding affinity energy is exhibited by the interaction between AFB1 and AKT1 (Table 3).

The binding site position of AFB1 on each target protein was also listed in detail. The result showed that AFB1 has different site position on each protein (Table 3), which is vividly depicted in Figure 2. This different position of binding site is exhibited since AFB1 as a ligand was exposed to the whole surface of target protein without any prior specific site. This process is performed to explore the favourable site and pose of AFB1 on target protein. Complex of AKT1-AFB1 was depicted by

the lowest binding energy during the molecular docking analysis. Therefore, it is predicted to have a strong bond and can trigger the biological responses.

The interaction of protein-ligand in the AKT1-AFB1 complex is pursued for further analysis to find the detailed interaction by using LigPlus. The results showed that AFB1 interacted in the protein domain AKT1 by using hydrophobic and hydrogen bonds. In detail, the interaction of hydrophobic bonds is formed through Trp80, Val270, Tyr272, Asp 292, Thr211, Leu 210, Leu 264, and Lys 268 residues, whereas hydrogen bond is only found in Ser205 residues. On the results visualization, the AFB1 structure is showed in the purple colour, the green dotted line showed the hydrogen bonds with amino acid residue, and half circle pattern in red demonstrated the interaction of hydrophobic bonds (Figure 3).

Further analysis was conducted to identify protein pathway of AKT1-AFB1 in HCC mechanism. Constructed protein interaction network (PIN) in HCC mechanism (Figure 1) was analysed by topological analysis based on number of edges and betweenness parameters score.



**Figure 4.** Visualization of topological analysis based on number of edges and betweenness parameter. Predicted protein interaction of AKT1 in HCC mechanism related to metastasis pathway (grey arrow).

The number of edges is indicated by the size of protein node, whereas the betweenness score is indicated by colour and thickness of the edges (Figure 4). The result showed that AKT1 protein had a lower betweenness score compared to GSK3B protein. AKT1 protein had thin edges which interacted to several proteins, while GSK3B had the thickest blue edges which directly interacted to EGF, AXIN1, and RPTOR with score 4.33, 4.0, 3.5, and 2.5, respectively (Table 5). According to another parameter, the analysis demonstrated the different result. Compared to other protein, AKT1 and mTOR protein exhibited the biggest size of protein node within a network (Figure 4) with 11 edges interacted to various proteins (Table 4). Further analysis showed that AKT1 had an interaction to another protein especially mTOR and RPS6KB1 protein which is indicated by the arrow (Figure 4).

#### **DISCUSSION**

Protein target of AFB1 was identified as shown in Table 1. There were three groups of protein predicted as a target of AFB1, such as kinase, phosphatase, and G protein-coupled receptor. In detail, there are 22 kinase proteins predicted as compatible target protein of AFB1.

Kinases are a family of enzyme which has a main function to catalyze the phosphorylation process. In general, kinases transfer a phosphoryl group to the target protein and play essential role for maintaining various cellular functions, such as proliferation, cell growth, motility, and apoptosis. 11,12 According to protein target prediction, it showed that AFB1 had several target proteins such as AKT/mTOR and MAPK, identified as signaling pathway which regulate various cellular pathway related to malignancies. It is known that hundreds of kinases are commonly activated in any tumor development.

Based on the Bhullar et al. 13 kinase protein is a common targeted protein of cancer drug after G protein-coupled receptor. 13 Since 1980, there were more than 30 kinase inhibitors received by FDA for treatment. Moreover, about 150 drugs for kinase-targeted are developed following clinical trials. The target protein prediction of AFB1 provides the future potential for kinase-targeted

therapeutics agent. Therefore, a specific target protein is needed to be explored.

Based on STRING analysis, there were 29 predicted AFB1 target proteins interacted each other through 97 edges. However, only seven proteins were predicted to be closely involved in HCC mechanism, such as RAF1, MAPK1, MAPK3, AKT1, EGFR, GSK3B, and mTOR. Although there are association among proteins, it does not mean that those proteins are directly binding each other. It merely showed that those candidates' proteins is strongly related each other to the HCC mechanism.

To understand the role of seven proteins related to HCC mechanism, the function of each protein needs to be spotted. Epidermal growth factor receptor (EGFR) is belonged to receptor tyrosine kinase (RTK), which is found in plasma membrane, and it has a function to regulate the cell growth. 10 Generally, the downstream signaling cascade of EGFR includes two main pathways, such as RAS-RAF-MAP kinase pathway and PI3K-PTEN-Akt pathway. In cancer case, the level of EGFR is frequently elevated or mutated. The alteration of EGFR promotes the constitutive activation of downstream signaling and leads to the unlimited proliferation of cells during G1/S phase.14 In brief, once the EGFR is activated, it will phosphorylate its substrate, which led to the downstream pathway related to AKT activation. Then, AKT will phosphorylate the mammalian target of rapamycin (mTOR). It is one of tumour survival kinase which is commonly overexpressed in tumour for apoptosis escape and metastasis. 15,16 Moreover, after activation, AKT will also be translocated into nucleus and regulate the gene responsible for cellular proliferation.17

Furthermore, AFB1 is estimated to have the ability to interact with one of seven predicted proteins within HCC mechanism. Therefore, molecular docking would be conducted between AFB1 and those selected proteins. Result of homology modelling showed that all of protein is considered valid according to the similarity score and validation score. The highest similarity score was exhibited by MAPK1 while the lowest score was showed by EGFR (Table 2). However, both similarity scores are still in the requirement range about 80-90% similar with the protein template.

According to Park and Cho,<sup>18</sup> the reverse docking had the opposite process as compared to virtual screening docking.18 Commonly, docking is performed to screen the protein against several compatible ligands. Nevertheless, in reverse docking, target proteins are listed and ranked against selected ligand. It also commonly known as ligand-based docking. The molecular docking result showed that among seven proteins, complex of AKT1-AFB1, MAPK3-AFB1, and MTOR-AFB1 are demonstrated to have the lowest binding affinity about -10.2 kcal.mol<sup>-1</sup>, -9.3 kcal.mol<sup>-1</sup>, and -9.1 kcal. mol<sup>-1</sup>, respectively. However, the lowest binding affinity energy is exhibited by interaction between AFB1 and AKT1 (Table 3).

Result showed that AFB1 has different site position of AFB1 on each protein (Table 3) which vividly depicted in Figure 2. This different position of binding site is exhibited since AFB1 as a ligand was exposed to the whole surface of target protein without any prior specific site. This process is performed for exploring the favourable site and pose of AFB1 on target protein.

Complex of AKT1-AFB1 was depicted by the lowest binding energy during the molecular docking analysis. Therefore, it is predicted to have a strong bond and able to trigger the biological responses. Results showed that AFB1 interacted in the protein domain AKT1 by using hydrophobic and hydrogen bonds. In detail, the interaction of hydrophobic bonds is formed through Trp80, Val270, Tyr272, Asp 292, Thr211, Leu 210, Leu 264, and Lys 268 residues, whereas hydrogen bond is only found in Ser205 residues. On the results visualization, the AFB1 structure is showed in the purple colour, the green dotted line showed the hydrogen bonds with amino acid residue, and half circle pattern in red demonstrated the interaction of hydrophobic bonds (Figure 3).

Further analysis was conducted to identify protein pathway of AKT1-AFB1 in HCC mechanism. Constructed protein interaction network (PIN) in HCC mechanism (Figure 1) was analysed by topological analysis based on number of edges and betweenness parameters score. The number of edges indicated by the size of protein node, whereas the betweenness score indicated by colour and thickness of the edges (Figure 4). Result showed that AKT1 protein had lower betweenness score as compared to GSK3B protein. AKT1 protein

had thin edges interacted to several proteins, while GSK3B had the thickest blue edges which directly interacted to EGF, AXIN1, and RPTOR with score 4.33, 4.0, 3.5, and 2.5, respectively (Table 5). According to Simos et al.,19 protein with high score of betweenness score is considered to have the ability to control the other protein interaction within a network.19 However, the controlled interaction of protein still needs further studies. According to another parameter, the analysis demonstrated the different result. As compared to other protein, AKT1 and mTOR protein exhibited the biggest size of protein node within a network (Figure 4) with 11 edges interacted to various proteins (Table 4). It is widely known that protein with high number of edges or direct interaction is determined as protein which have the most critical role within network due to their abroad interaction.19 Based on that analysis, it showed that AKT1 and mTOR are considered as the essential protein within HCC mechanism. Therefore, interaction of AFB1 to AKT1 protein is suggested as potential strategy in promoting the HCC progression.

Further analysis showed that AKT1 had an interaction to another protein especially mTOR and RPS6KB1 protein which indicated by the arrow (Figure 4). The mTOR protein is phosphorylated by AKT1 protein, then it activated the protein downstream termed RPS6KB1, which is strongly involved in the cancer survival mechanism.20 RPS6KB1 is known as the major substrate of mTOR, which plays an essential role in cell growth, survival, and metastasis by cell cycle regulation. Recent research reported that RPS6KB1 is commonly high expressed in cancer related to the tumour size and metastasis progression. Moreover, overexpression of RPS6KB1 is also considered for exhibiting the worse condition of HCC case in patients.<sup>21</sup> Another in vitro result exhibited that AKT1 signalling had oncogenic effect for promoting the hepatocarcinogenesis. Akt is a central of PI3K/ Akt signaling pathway which regulates the cells proliferation and cells survival ability.

Based on all analysis, it is suggested that AFB1 have interaction to the most potential protein target in HCC mechanism that is AKT1 protein. The interaction of AKT1-AFB1 involved in the HCC progression by activated protein cascade of AKT1-mTOR-RPS6KB1, which generated the

metastasis mechanism. This indicated that the interaction between AFB1 and AKT had essential implications for HCC development.

#### **CONCLUSION**

AFB1 is predicted to interact with several proteins of hepatocellular carcinoma (HCC) major factors. Kinase's proteins, such as RAF1, MAPK1, MAPK3, AKT1, EGFR, GSK3B, and mTOR exhibited to have close related to HCC development. However, among those target proteins, Akt1 had strong interaction AFB1 with binding energy -10.2 kcal.mol<sup>-1</sup> and supported by hydrophobic bonds through Trp80, Val270, Tyr272, Asp292, Thr211, Leu210, Leu264, and Lys268 residues, also hydrogen bond with Ser205 residue. Further analysis demonstrated that the interaction AKT1-AFB1 is related to metastasis pathway in HCC mechanism. This finding would provide the understanding about HCC mechanism, which can be used as a basic development of cancer therapeutic agent.

#### **ACKNOWLEDGMENTS**

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#### **CONFLICT OF INTEREST**

The authors declare that there is no conflict of interest.

#### **AUTHOR'S CONTRIBUTION**

All authors listed have made a substantial, direct, and intellectual contribution to the work and approved the final manuscript for publication.

#### **FUNDING**

None.

#### DATA AVAILABILITY

The datasets generated during and/or analyzed during the current study are available

from the corresponding author on reasonable request.

#### **ETHICS STATEMENT**

Not applicable.

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