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# Identification of potential inhibitors targeting DNA adenine methyltransferase of *Klebsiella pneumoniae* for antimicrobial resistance management: a structure-based molecular docking study

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## Abstract

**Background:** *Klebsiella pneumoniae* is an important opportunistic pathogen that frequently causes nosocomial infections. Notably, this bacterium has emerged as a major problem in hospital settings because of its acquisition of resistance to carbapenems. The majority of antibiotics act by targeting crucial pathways within bacterial cells. However, due to the development of resistance mechanisms, the efficiency of these antibiotics has decreased. Therefore, this study focused on a putative protein (DNA adenine methyltransferase; Dam) found in *K. pneumoniae* that encompasses a DNA methylation protein domain, indicating a novel potential target for pharmacological intervention. DNA methylation affects bacterial virulence attenuation.

**Methods:** In the unavailability of a 3D structure for Dam protein in protein database, a 3D model was generated using SWISS-MODEL server and validated using computational tools. Following that, screening was performed against the Dam protein using a set of 2706 phytochemicals obtained from the ZINC database using PyRx0.8. ProTox-II platform was used to predict the physicochemical properties and various toxicity endpoints.

**Results:** Among the screened compounds, ZINC4214775, ZINC4095704, and ZINC4136964 had higher binding affinity for the Dam and interacted with its active site residues. The computational analyses of these three identified hits indicate that their predicted properties were within an acceptable range for evaluating toxicity. In addition, a toxicity radar chart showed that these hits were within an acceptable range.

**Conclusions:** These compounds have the potential to act as Dam inhibitors and could be investigated further for managing antimicrobial resistance in *K. pneumoniae*.

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## Keywords:

*Klebsiella pneumoniae*;  
Dam; Antimicrobial  
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## Introduction

*Klebsiella pneumoniae*, a member of the Enterobacteriaceae family, is an indigenous bacteria of the human and animal gastrointestinal microbiota [1]. This bacterium has multiple antibiotic resistance mechanisms and frequently acts as a pathogen, causing nosocomial infections such as surgical wound infections, gastrointestinal infections, and community-onset infections, and frequently causing outbreaks within healthcare facilities. The global prevalence of drug-resistant *K. pneumoniae* has increased significantly, reaching up to 70%, and the associated mortality rate from infections caused by this pathogen ranges between 40% to 70% [2]. The emergence of multidrug-resistant as well as carbapenem-resistant *K. pneumoniae* has become a major global public health concern [3].

DNA adenine methyltransferase (Dam) is a promising target for therapeutic intervention because it plays an important role in the epigenetic regulatory apparatus that is responsible for bacterial pathogen survival and pathogenicity [4]. Dam has been implicated in a variety of cellular processes such as mismatch repair [5], replication regulation, transposition [6], and gene expression control [7]. Furthermore, studies have found a link between Dam and antibiotic resistance. Elevated resistance to ampicillin, tetracycline, and nalidixic acid has been observed in bacteria due to epigenetically induced changes in the expression of resistance-associated genes [8].

The process of introducing a new pharmaceutical medication to the market necessitates a significant financial and time investment. The average time for drug development is 10-15 years, with a cost estimate of approximately \$800 million [9,10]. Computer-aided drug discovery (CADD) has emerged as an essential component of numerous projects conducted in a variety of settings and research environments. CADD has significantly aided in the identification and optimization of hit compounds, propelling them to advanced stages of the drug discovery pipeline or eventual market entry [11]. This study aimed to find natural Dam inhibitors to fight the antimicrobial resistance in *K. pneumoniae*.

## Methods

### Sequence retrieval and homology modeling of Dam protein

The UniProt database was used to find *K. pneumoniae* protein sequences containing the Dam domain (UniProt ID: A0A2U0NNR3). The SWISS-MODEL server was used to build the targeted 3D protein structure, which involved fully automated procedures. The structure of the protein of interest was first generated using

homology modeling, which was then compared to a suitable template protein structure [12]. The structural properties of the Dam-containing protein in *E. coli* (PDB ID: 4RTL) were used to select the template protein [13].

### Validation of modeled structure

To validate the modeled structure, we initially conducted energy minimization. Subsequently, we uploaded the modeled structure to the SAVES server (<http://nihserver.mbi.ucla.edu/SAVES/>), which assessed its overall correctness using ERRAT and evaluated the stereochemical quality with PROCHECK. Additionally, we employed Verify3D to determine the compatibility of the atomic model with its corresponding amino acid sequence. PDBsum tool was also employed to access the Ramachandran plot and secondary structure of the model protein. These comprehensive evaluations were performed to ensure the accuracy and reliability of the modeled structure.

### Active site prediction

The active site of a protein is a distinct binding site that plays a crucial role in its specific function. Identifying these active sites is a fundamental step in structure-based drug design, as it helps determine potential targets for therapeutic interventions [14]. Precise prediction of active sites is an invaluable tool in the field of bioinformatics. In this study, we employed Discovery Studio software to predict the active site of the modeled protein. By utilizing this software, we were able to identify the key binding site, providing valuable insights for further docking analysis.

### Natural compound library retrieval and preparation

'Biopurify Phytochemicals' subsets from the ZINC database consisting of 2706 compounds were downloaded in sdf format. These compounds were subjected to energy minimization using 'uff' force field and prepared as converting into pdbqt format for further docking-based virtual screening (VS).

### Virtual screening

VS was performed with the PyRx 0.8 [15] software tools to evaluate the interaction of the natural compound library with Dam protein. The binding affinity score was used as a metric for the evaluation.

### Physicochemical properties and toxicity endpoints prediction

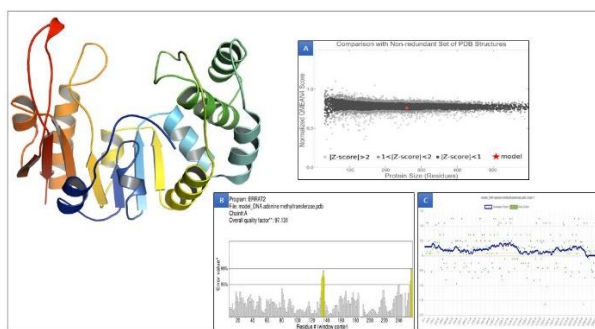
The top three selected compounds were subjected to evaluation of their physicochemical properties and predictions of various toxicity endpoints using the ProTox-II platform [16].

## Results

Initially, protein sequences of *K. pneumoniae* containing the Dam domain were retrieved from the

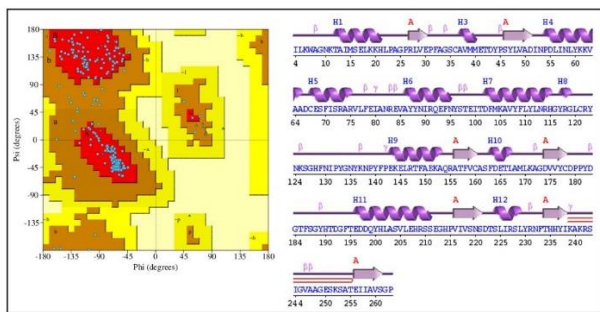
UniProt. Since there was no existing 3D structure available for this protein in the protein database, a 3D model was constructed using the SWISS-MODEL server. The structural properties of the Dam-containing protein in *E. coli* (PDB ID: 4RTL) were utilized as a template for the modeling process.

Several tools and web servers were used to validate the accuracy of the modeled structure. For this, the SAVES server and PDBsum were employed. According to PROCHECK analysis, 89.2% of the phi/psi angles of residues in the modeled structure were within the most favored regions, indicating a favorable stereo-chemical quality. Furthermore, the ERRAT graph produced a quality factor of 97.131, indicating a good model quality (a score greater than 50 is considered acceptable). The VERIFY3D server confirmed the model's high quality, with 95.24% of residues in the modeled protein scoring higher than 0.2. A normalized QMEAN4 score was also predicted, and VERIFY3D analysis revealed that 85.38% of the residues had an average 3D-1D score of  $>=0.1$ , indicating a "Pass" result for the structure (Figure 1).



**Figure 1:** 3D structure of modeled protein with validation metrics. The normalized QMEAN4 score is used to assess model quality (A). PROCHECK analysis elucidates the overall quality of the protein structure (B). The VERIFY3D assessment validates the model's reliability (C).

PDBsum tool predicted the secondary structure of the modeled structure, which consisted of the motif having 1 sheet, 3 beta alpha beta units, 1 beta-hairpin, 7 strands, 12 helices, 12 helix-helix interactions, 17 beta turns, and 4 gamma turns. Additionally, a Ramachandran plot was also generated (Figure 2).

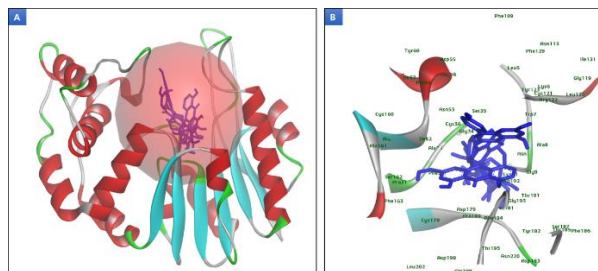


**Figure 2:** Ramachandran plot (left panel) and secondary structure (right panel) of the modeled structure.

After validating the modeled protein structure and predicting its binding sites using DS 2021 software, we utilized the 'define and edit binding site' option. Through this process, it was determined that the best site had the attributes  $X=0.451893$ ,  $Y=3.081089$ , and  $Z=90.425250$ . Subsequently, this site was utilized for the VS of prepared ligands, specifically natural compounds. We selected the top 10 hits for this study based on their binding affinity, which is listed in Table 1. Furthermore, we focused on the top 3 hits, providing a detailed description of their binding poses (Figure 3) within the active pocket and their interaction with active site residues.

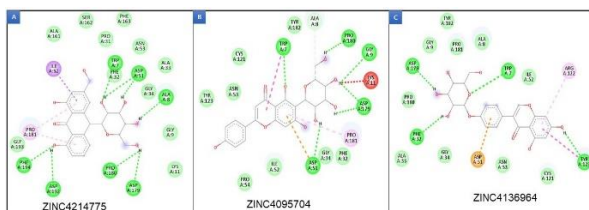
Natural compound hits	Binding affinity (kcal/mol)
ZINC4214775	-9.6
ZINC4095704	-9.6
ZINC4136964	-9.3
ZINC4098548	-8.9
ZINC3983877	-8.7
ZINC82046436	-8.7
ZINC238766490	-8.5
ZINC35457329	-8.5
ZINC36479659	-8.4
ZINC238809241	-8.2

**Table 1:** List of top ten screened natural hits and their binding affinity values.



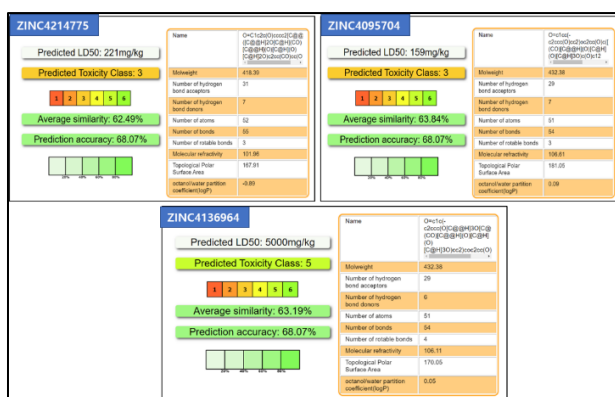
**Figure 3:** Interaction of top 3 ligands in the active pocket of target. Binding poses of ligands in site sphere (A), active site/binding site residues of the target protein (B).

The detailed interaction analysis of the top 3 hit compounds is as follows. ZINC4214775 was found to interact with Asp192, Phe194, Gly193, Pro181, Ile52, Ala161, Ser162, Pro31, Phe163, Trp7, Phe32, Asn53, Asp51, Ala33, Gly34, Ala8, Gly9, Lys11, Asp179, and Pro180 residues of Dam. The Phe194, Asp192, Trp7, Asp51, Ala8, Asp179, and Pro180 residues formed H-bond with ZINC4214775 (Figure 4A). ZINC4095704 interacted with Tyr123, Asn53, Cys121, Trp7, Tyr182, Ala8, Pro180, Gly9, Lys11, Asp179, Pro181, Phe32, Gly34, Asp51, Ile52, and Pro54 residues of Dam. The Trp7, Pro180, Gly9, Asp179, and Asp51 residues H-bonded with ZINC4095704 (Figure 4B). Further, ZINC4136964 was found to interact with Asp51, Gly34, Phe32, Ala33, Pro180, Asp179, Gly9, Tyr182, Pro181, Ala8, Trp7, Ile52, Arg122, Tyr123, Cys121, and Asn53 residues of Dam. The Phe32, Asp179, Trp7, and Tyr123 residues formed H-bond with ZINC4136964 (Figure 4C).

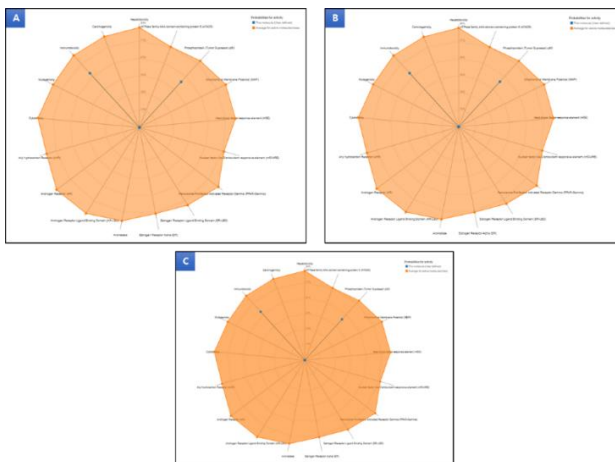


**Figure 4:** Interacting residues of Dam protein with ZINC4214775 (A), ZINC4095704 (B), and ZINC4136964 (C).

The predicted values for these 3 hits indicate that they fall within an acceptable range for toxicity assessment, as shown in Figure 5. Additionally, the toxicity radar chart, which demonstrates the confidence of positive toxicity results compared to the average of its class, was generated. It reveals that these hits fall within an acceptable range, as depicted in Figure 6.



**Figure 5:** Predicted oral toxicity assessment for the top 3 hits (ZINC4214775, ZINC4095704, and ZINC4136964).



**Figure 6:** Comparative toxicity radar chart for the top 3 hits, illustrating the toxicity profiles of ZINC4214775 (A), ZINC4095704 (B), and ZINC4136964 (C).

## Discussion

*K. pneumoniae* is a leading cause of hospital-acquired infections and is becoming increasingly drug-resistant [17]. The Dam is a protein that regulates genes associated with antibiotic resistance [8]. This study

identified natural Dam inhibitors that could be used to combat antimicrobial resistance in *K. pneumoniae*. Among the identified compounds, this study focused on the top three hits (ZINC4214775, ZINC4095704, and ZINC4136964) that had strong binding with DAM protein and interacted with active site residues of DAM protein. Furthermore, we utilized the ProTox-II web tool to conduct toxicity assessments on the top 3 compounds. Prediction of compound toxicities plays a vital role in the drug design and development process, as it helps determine the safety profile of potential drugs. The ProTox-II server offers predictions for a wide range of toxicity endpoints, including acute toxicity, hepatotoxicity, cytotoxicity, carcinogenicity, mutagenicity, immunotoxicity, adverse outcomes pathways, and toxicity targets. By utilizing these diverse techniques, ProTox-II provides comprehensive insights into the potentially toxic effects of compounds, facilitating the identification of safer and more effective drug candidates [16]. The three hits (ZINC4214775, ZINC4095704, and ZINC4136964) were found to have acceptable physicochemical properties and toxicity profiles. Using natural ingredients in medicine discovery and the food industry has various advantages. These contain unmatched chemical diversity characterized by complicated structures and powerful biological activities. Natural products occupy a different region within the chemical space, complementing synthetic chemicals. Furthermore, the creation of libraries including natural product analogs has the potential to improve drug-like qualities. By enhancing the regulation of natural product biosynthesis, it becomes able to investigate and comprehend targets and pathways involved in disease processes. Furthermore, using natural compounds allows for a more efficient transition from initial hits to medication development [18]. The hit compounds identified in this study are naturally occurring compounds.

This study focused on Dam protein found in *K. pneumoniae* that encompasses a DNA methylation protein domain. Screening of phytochemicals against the Dam protein was performed to identify potential compounds of interest. Notably, ZINC4214775, ZINC4095704, and ZINC4136964 had higher affinity for the Dam protein and interacted with its active site residues. The predicted physicochemical properties and toxicity profiles of these compounds were found to be within an acceptable range. These compounds have the potential to act as Dam inhibitors and could be studied further for antimicrobial resistance management in *K. pneumoniae*.

## Author Contributions

Conceptualization, Farah Anjum and Alaa Shafie; Methodology, Norah Ali Alshehri and Samia MUSAAD



Almutairi; Writing original draft, Ameerah Ali Alshehri and Ayshah Musad Almutairi; Review & editing, Farah Anjum.

## Conflicts of interest

The authors declare no conflict of interest.

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