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Protective Effects of Mirazid on Gentamicin-induced Nephrotoxicity in Rats through Antioxidant, Anti-inflammatory, JNK1/iNOS, and Apoptotic Pathways; Novel Mechanistic Insights

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Abstract

Background: As the use of Gentamicin became more widespread, the drug's harmful effects, particularly nephrotoxicity, became increasingly well-known. Antibacterial and anti-inflammatory properties have long been associated with Mirazid. The goal of this research was to find out more about frameworks for the protection of Mirazid against nephrotoxicity triggered by Gentamicin.

Methods: Three groups of albino male rats were created; the normal group received only saline. In the second group, nephrotoxicity was produced for 10 days with Gentamicin (100 mg/kg; i.p.). In the third group; Mirazid (10 mg/kg; p.o.) was administered for 10 days before receiving Gentamicin. This was done to investigate the kidney/body weight index, serum creatinine, urea, lactate dehydrogenase (LDH), malondialdehyde (MDA), and Glutathione (GSH) levels. Moreover, immunohistochemical staining was done to study Jun N-terminal kinase 1 (JNK1), inducible nitric
oxide synthase (iNOS), and caspase3 expressions along with histopathological changes. Additionally, a molecular docking study was performed for the seventeen isolated and identified compounds from myrrh, JNK1 is inhibited by an oleo-gum resin derived from the Commiphora species of plants (Burseraceae).

**Results:** The Gentamicin group showed an increase in kidney/body weight index, serum creatinine, urea, LDH, and MDA, while decreasing GSH levels. Furthermore, immunohistochemical staining revealed increased JNK1, iNOS, and caspase3 expressions along with histopathological changes. All of these indicators were significantly reduced by mirazid, which also restored oxidant/antioxidant hemostasis. Furthermore, the histological architecture of tissues has been significantly conserved. Concerning the docking study, the isolated compound (12) was found to be superior to the co-crystallized inhibitor (18) with a binding score of -7.19 kcal/mol compared to -6.95, respectively.

**Conclusion:** Mirazid was found to be a potential method for suppressing the nephrotoxicity caused by Gentamycin by inhibiting the JNK1/iNOS pathways, therefore preserving kidney function. The antioxidant, anti-inflammatory, and anti-apoptotic properties of mirazid are thought to be responsible for its preventive efficacy.

**Keywords:** Gentamicin, Mirazid, Nephrotoxicity, *In vivo*, JNK1/iNOS, Molecular docking.
1- Introduction

Aminoglycosides have long been linked to drug-induced nephrotoxicity, which is one of the most common side effects\(^1\). Nephrotoxicity triggered by Gentamicin involves pathological conditions, such as altered intraglomerular hemodynamics, the toxicity of tubular cells, and inflammation\(^2\). Due to the disruption of oxidant-antioxidant systems, toxicity is induced by the generation of free radicals and protein oxidation\(^3, 4\). The production of reactive oxygen species (ROS), as well as the activation of a number of inflammatory mediators, have all been associated with gentamicin-induced nephrotoxicity\(^5\).

Oxidative stress-induced by gentamicin plays a vital role in the activation of pro-inflammatory cytokines, including Jun N-terminal Kinase (JNK1) which leads to kidney damage. Free radicals stimulate glomerulus lipid peroxidation and influence the normal physiological function of renal tissues, contributing to metabolic disorders\(^6\). Renal inflammatory cascades, renal oxidative stress, and pathogenic signaling systems are all exacerbated in nephrotoxicity\(^7\). Previous studies have indicated that medications with significant antioxidant and anti-inflammatory cellular characteristics can be employed to combat Gentamicin's nephrotoxicity\(^8-10\).

As the inflammatory system becomes more activated, pro-inflammatory mediators such as chemokines and cytokines are released, resulting in inflammatory signals. These signals enable the body to recognize, destroy, and eliminate foreign objects, resulting in an effective acute inflammatory response\(^11\). Immune suppression and the onset of chronic inflammatory diseases can be caused by the inappropriate production of pro- or anti-inflammatory mediators. Tissue injury and degeneration are connected to inflammation over time. This has been recognized as a necessary condition for the start of numerous neurological, autoimmune, and malignant diseases\(^12\). Macrophages are crucial participants in the immune and inflammatory responses that occur during a host's defense. Once activated, they start producing cytokines, oxygen, and nitrogen species. This stimulation causes cytokines to be released and enzymes like inducible nitric oxide synthase (iNOS) to be produced\(^13\).

Mirazid, a drug that has been on the Egyptian market for over a decade, is made from (Arabian or Somali) myrrh, an oleo-gum resin derived from plants of the Commiphora species (Burseraceae)\(^14\). Because of its antimicrobial activity, infections and inflammation are treated with Mirazid\(^15\). Also, it treated blood stagnation, inflammatory diseases, and reduced swelling and pain\(^16\). In clinical trials, some myrrh-based recipes were utilized as anticancer medicines in the treatment of liver, pancreatic, and nasopharyngeal malignancies and have shown adequate curative efficacy\(^17, 18\). Extracts of these plants' resinous exudates and/or their constituents showed analgesic\(^19\) anti-
inflammatory, lipid-lowering, neuroprotective, and antibacterial properties. Mirazid's influence on mucus formation and up-regulation in sulphydryl concentrations of nucleic acid, as well as its free radical-scavenging, thyroid-stimulating, and prostaglandin-inducing capabilities are hypothesized to play a role in the treatment of stomach ulcers.

One of the most essential technologies for drug discovery is computational drug design and development is molecular docking. It helps scientists to design new drugs, repurpose existing candidates, or study their mechanisms of action. Molecular docking is a crucial tool in molecular biology and computer-aided drug design. Its main goal is to predict the most important binding mode(s) of a ligand with a protein of known 3D structure. Successful docking uses a scoring function that correctly ranks candidate dockings.

The study of ways to reduce the toxicity of aminoglycosides continues to pique clinical interest. As a result, the goal of this study was to see how Mirazid affected oxidative stress, inflammation, and apoptotic pathways in Gentamicin-induced nephrotoxicity.

2- Materials and methods

2.1. Experimental animals

In this investigation, twenty-four male rats (albino Wistar) weighing 190-220 g were employed. The rats were provided by the Modern Veterinary Office For Laboratory Animals (Cairo, Egypt). Rats were kept in a temperature-controlled environment (25°C) with a 12-hour light/dark cycle. Food and water were allowed ad libitum during the study period. Before the trial, the rats were given two weeks to acclimate in the laboratory. The protocol for the experiment was approved by the Research Ethics Committee, Faculty of Pharmacy, Delta University (FPD4 15/2018).

2.2. Drugs

Gentamicin (Gentamicin Sulphate) vials were purchased from Sigma company, U.S.A. Gentamicin sulfate is 80 mg per ml in each vial. Mirazid capsules were purchased from the producing company Pharco Pharmaceuticals (Alexandria, Egypt).

2.3. Induction of nephrotoxicity

The intraperitoneal administration of Gentamicin (100mg/kg/body weight) for 10 days resulted in nephrotoxicity.

2.4. Experimental protocols

Twenty-four rats were allocated into three groups at random (8 rats each). Control group, rats did not receive any drug or solvent, Gentamicin group; rats were injected with Gentamicin (100
mg/kg; i.p.) for 10 days and Mirazid prophylactic group; Mirazid (10mg/kg; p.o.) was administrated to rats starting 10 days before Gentamicin administration\(^5\).

### 2.5. Sacrification and biological samples collection

Under light ether anesthesia, a clean sterile capillary tube was inserted in the inner canthus of the eye to collect blood samples from the orbital sinus (retro-orbital plexus). After allowing the blood to coagulate for 20 minutes, it was centrifuged for 15 minutes at 4000 rpm. The serum samples were then separated, collected in clean tubes, and maintained at -20°C until they were used to determine serum creatinine, urea levels, and lactate dehydrogenase (LDH) levels using a colorimetric kit, as directed by the manufacturer. Rats were sedated with thiopental sodium (50 mg/kg) and killed by cervical dislocation at the end of the experiment. The kidneys were separated and rinsed in ice-cold phosphate-buffered saline (pH = 7.4). Body weights and kidney weights were measured for the calculation of kidney/body weight index. The right kidney was rinsed in ice-cold saline, sliced lengthwise, and preserved in 10% buffered formalin for histological investigation. The left kidney was submerged in liquid nitrogen and stored at 80 °C for tissue homogenate preparation.

### 2.6. Preparation of kidney homogenate

Kidney homogenate (10% w/v) was made using roughly 2 cm of kidney tissue in ice-cold KCl (1.15 percent, pH 7.4). The homogenate was centrifuged for 10 minutes at 3000 rpm at 4 °C, and the supernatants were decanted and utilized to measure kidney GSH and MDA levels.

### 2.7. Determination of serum creatinine, urea, and lactate dehydrogenase (LDH)

The assay is based on Jaffe's description of the reaction of creatinine with sodium picrate. Creatinine forms a crimson complex with alkaline picrate. Interferences from other serum constituents are avoided by the time interval used for measurements. The amount of colour generated is proportional to the amount of creatinine in the sample. Serum urea was determined enzymatically according to the previously described method of Kaplan and Kohn\(^5\). The level of LDH in the serum was determined using a spectrophotometer and commercially available kits (Biomed Diagnostics test kits), as directed by the manufacturer (Egypt).

### 2.8. Determination of kidney GSH concentration and MDA content

GSH concentration and MDA content were determined according to the manufacturer's instructions using a commercially available kit (Biodiagnostic, Giza, Egypt). According to a previously established approach, lipid peroxidation (LPO) was measured as thiobarbituric acid reactive substances (TBARS) in terms of generated MDA\(^5\). Glutathione (GSH) content was assessed according to a method illustrated earlier\(^5\).
2.9. Histopathological examination

For histological assessment, a 2 cm piece of the right kidney was removed, rinsed in cold saline, fixed in 10% buffered formalin solution, sliced transversely, paraffin-embedded, and 3m slices stained with hematoxylin and eosin (H&E). The tissues were evaluated using an Olympus CX21 microscope in a random order, with the histopathologist blinded to the experimental groups. For image analysis, slides were taken using an Olympus® digital camera set on an Olympus® microscope with 1/2 X photo adapter and a 40 X objective utilizing a computer-assisted digital image analysis (Digital morphometric study). The photos were examined using Video Test Morphology® software (Russia) with a built-in process for stain quantification and automated object analysis on an Intel® Core I3® based computer. All measurements are validated against a micrometer slide that was photographed with the same instrument at the same magnification using the same technique. This allows measurements to be taken in um² rather than square pixels. SO for Hx, the change in the number of inflammatory cells (cells/μm²).

2.10. Immunohistochemical evaluation of JNK 1, iNOS and Caspase3

For antigen retrieval, kidney slices were dewaxed and submerged in a solution of 0.05 M citrate buffer, pH 6.8. After that, the sections were treated with 0.3 % hydrogen peroxide and protein block. The sections were then treated with anti-JNK1, anti-iNOS, and anti-caspase3 polyclonal antibodies (Santa Cruz, Cat# (F-6): sc-8008, 1:100 dilution). After rinsing with phosphate-buffered saline, they were incubated for 30 minutes at room temperature with a goat anti-rabbit secondary antibody (Cat# K4003, EnVision+TM System Horseradish Peroxidase Labelled Polymer; Dako). Slides were visualized using a DAB kit before being counterstained with Mayer's hematoxylin. In a total of 1000 cells per 8 HPF, the staining intensity was evaluated and expressed as a percentage of positive expression. All measurements are calibrated against a micrometer slide that was photographed with the same instrument at the same magnification using the same technique. This allows measurements to be taken in um² rather than square pixels. JNK 1, iNOS, and Caspase-3 are scored based on the change in staining intensity.
2.11. Docking studies

Using GC–MS, and ICP–MS separation techniques, we previously identified seventeen chemicals from myrrh resin that were subjected to molecular docking studies using MOE 2019.012 suite to propose its mechanism of action as a promising JNK-1 inhibitor. JNK-1 inhibition is responsible for stopping both the apoptotic and iNOS pathways which are proposed to be the main mechanism of action responsible for the anti-inflammatory effects of myrrh. Also, thiophenecarboxamide urea (TCU) native co-crystallized inhibitor was used as a reference standard.

2.11.1. Preparation of the myrrh resin extract

The seventeen isolated and identified compounds from myrrh resin were downloaded from the PubChem database website. They were inserted into MOE and prepared for docking by applying the previously described steps. Then, they were imported together with the co-crystallized JNK-inhibitor (TCU) in a single database file and saved as an MDB file to be ready for the docking process.

2.11.2. Preparation of the target JNK-1 pocket

The JNK-1 X-ray structure (code: 3PZE) was downloaded from the Protein Data Bank and the full preparation steps for its preparation were applied as before.

2.11.3. Docking of myrrh isolated compounds to the binding pocket of JNK-1

The prepared database containing the isolated identified compounds was inserted in a general docking process and all the default methodology steps were performed as described earlier. By the end, we filtered the best poses according to their scores, RMSD, and amino acid interactions for all the examined compounds. Moreover, a process of redocking for the co-crystallized ligand inside the binding pocket of the JNK-1 receptor was applied to validate the MOE program. The validation was confirmed by obtaining low RMSD values (< 1) between the docked and native ligands.

2.12. Statistical analysis

For statistical comparison between parametric and nonparametric data, one-way analysis of variance (ANOVA) followed by Tukey–Kramer multiple comparison tests and Kruskal-Wallis test followed by Dunn's Multiple Comparison tests were employed, respectively. A $P$-value < 0.05 was the established level of significance

3- Results

3.1. Effect of prophylactic Mirazid on Gentamicin triggered a modification in the serum level of creatinine, urea levels, and LDH
When compared to the control group, serum creatinine and urea levels were considerably ($P < 0.05$) higher following Gentamicin administration. Prophylactic daily oral Mirazid (10 mg/kg) for 10 days resulted in a substantial ($P < 0.05$) decrease in serum creatinine and BUN as compared to the Gentamicin group. The nephrotoxicity marker LDH is used to diagnose renal impairment. Gentamicin treatment greatly deteriorated kidney functions contrasted to the control group. Serum LDH levels increased significantly ($P < 0.05$). Prophylactic daily oral Mirazid (10 mg/kg) for 10 days resulted in a substantial ($P < 0.05$) decrease in serum LDH as compared to the Gentamicin group (Table 1).

Table 1. Effect of Mirazid (10mg/kg; p.o) on Gentamycin (100 mg/kg; i.p.) triggered a modification in serum creatinine, BUN, and LDH.

| Treatment group                  | Creatinine (mg/dl) | BUN (mmol/l) | LDH (U/L) |
|----------------------------------|--------------------|--------------|-----------|
| Control                          | 0.32 ± 0.02        | 36.75 ± 1.8  | 1350 ± 19.5|
| Gentamicin group                 | 0.74 ± 0.01 *      | 114.60 ± 3.1 * | 4409 ± 10.9 * |
| Mirazid prophylactic group       | 0.41 ± 0.01 #      | 41.33 ± 2.3 # | 1675 ± 11.94 # |

Results are expressed as mean± S.E.M, n = 8. Statistical analysis was performed using One-Way ANOVA followed by Tukey-Kramer multiple comparisons test at $P<0.05$. $^*P < 0.05$ concerning the control group, $^#P< 0.05$ concerning the Gentamicin group.

3.2. Effect of prophylactic Mirazid on Gentamicin triggered a modification in oxidants/antioxidant stress markers in the kidney; MDA content and decreased GSH

The administration of gentamicin elevated MDA levels while lowering GSH levels significantly. Concurrent administration of Mirazid (10 mg/kg, orally) significantly ($P<0.05$) ameliorated Gentamicin-induced kidney damage. As MDA content was reduced and GSH activity was restored, Mirazid significantly preserved kidney GSH activity when contrasted to the Gentamicin group and significantly decreased kidney MDA activity in rats in comparison with the disease group. On the other hand, Gentamicin administration showed a significant ($P<0.05$) up-regulation in MDA activity when contrasted to the control group (Table, 2).
Table 2. Effect of Mirazid (10 mg/kg; p.o) on Gentamycin (100 mg/kg; i.p.) triggered a modification in kidney homogenate GSH and MDA concentration.

| Groups                  | GSH (μmol/g tissue) | MDA (nmol/g tissue) |
|-------------------------|---------------------|---------------------|
| Control                 | 0.71 ± 0.02         | 33.75 ± 1.8         |
| Gentamicin group        | 0.32 ± 0.01 *       | 111.10 ± 3.3 *      |
| Mirazid prophylactic group | 0.69 ± 0.04 #      | 48.33 ± 2.3 #       |

Results are expressed as mean± S.E.M, n = 8. Statistical analysis was performed using One-Way ANOVA followed by Tukey-Kramer multiple comparisons test at P<0.05. *P< 0.05 concerning the control group, #P< 0.05 concerning the Gentamicin group.

3.3. Effect of prophylactic Mirazid on Gentamicin triggered a modification in kidney/body weight index

When compared to the control group, the kidney/body weight ratio of the Gentamicin group was considerably higher (P <0.05). The Mirazid prophylactic group, on the other hand, saw a significant (P <0.05) reduction in kidney weight when compared to the Gentamicin group. According to Mirazid prophylactic group, there was a significant improvement in modifying kidney/body weight index. However, the kidney/body weight index of Mirazid prophylactic group is still significantly increasing in comparison to the control group; it could not reach normal weight (Figure, 1).

![Figure 1](image)

**Figure (1):** Effect of Mirazid (10mg/kg) orally for 7 days on Gentamycin (100 mg/kg; I.P)-induced changes in kidney/body weight index. Results are expressed as mean± S.E.M, n = 8. Statistical analysis was performed using One-Way ANOVA followed by Tukey-Kramer multiple comparisons test at P < 0.05. *P < 0.05 concerning the control group, #P < 0.05 concerning the Gentamicin group.
3.4. Effect of prophylactic Mirazid on hematoxylin and eosin-stained kidney specimens, gentamicin caused histopathological changes

As illustrated in Figure 2, kidney tissue photomicrographs of A, control animal showing normal glomeruli with an intact bowman's capsule (arrow); B, sections of animals treated with gentamycin showing glomerular congestion (arrow), inflammatory cell infiltration (filled arrowhead), and necrosis (open arrowhead); C, sections of animals exposed to gentamycin and treated with mirazid showed isolated mild tubular damage in the form of tubular dilatation with an irregular contour and tubular vacuolization (filled arrowhead), but no glomerular abnormalities or neutrophil infiltration were observed (arrow). Scale bar = 20 µm.

Figure (2): Representative photomicrographs for sections from renal tissue of rats stained with H&E stain. Scale bar = 20 µm.

Table 3. Effect of Mirazid on Gentamycin (100 mg/kg; I.P)- triggered a modification in inflammatory cells count for H&E.

| Groups                        | Inflammatory cells (cells / µm²) |
|-------------------------------|----------------------------------|
| Control group                 | 20.50 ± 1.11                     |
| Gentamicin group              | 177.3 ± 6.33 *                   |
| Mirazid prophylactic group    | 120.7± 5.01 *#                  |

Results are expressed as mean± S.E.M, n = 8. Statistical analysis was performed using One-Way ANOVA followed by Tukey-Kramer multiple comparisons test at P<0.05. P< 0.05 concerning the control group, *P< 0.05 concerning the Gentamicin group.

3.5. Effect of prophylactic Mirazid on Gentamicin triggered activation of renal apoptosis; immunohistochemical analysis of caspease-3 expression

As an apoptotic marker, the expression of caspase 3 stained cells in tissue was evaluated. Caspase-3 immunostaining was negative in the control group. On the contrary, the Gentamicin group
showed strong immunostaining for caspase-3 indicating the existence of apoptotic activity in kidney tissue contrasted to the control group. Caspase-3 expression was downregulated in the prophylactic daily oral Mirazid (10 mg/kg) contrasted to the Gentamicin group (Figure, 3).

**Figure (3):** A) Microscopic pictures of immune-stained renal sections against Caspase3. IHC counterstained with Mayer's hematoxylin. Black arrows point to positive tubules. Low magnification X:100 bar 100 and high magnification X:400 bar 50 B). Effect of Mirazid (10 mg/kg) orally for 7days on Gentamycin (100 mg/kg; I.P)-induced change in inflammatory cells to count for H&E. Results are expressed as means S.E.M, n = 8. Statistical analysis was performed using One-Way ANOVA followed by Tukey-Kramer multiple comparisons test at P<0.05. *P< 0.05 concerning the control group, #P< 0.05 concerning the Gentamicin group
3.6. Effect of prophylactic Mirazid on Gentamicin triggered activation of renal inflammation; immunohistochemical analysis of i-NOS and JNK1 expressions

Expressions of i-NOS and JNK1 stained cells were evaluated as a marker of inflammation. The Control group revealed negative immunostaining for i-NOS (Figure 4) and JNK1 (Figure 5). The Gentamicin group showed strong immunostaining for i-NOS and JNK1 expressions contrasted to the control group. On the other hand, Prophylactic daily oral Mirazid (10 mg/kg) dose downregulated i-NOS and JNK1 expressions contrasted to Gentamicin group, yet this decrease failed to reach a normal level.

![Microscopic pictures of immune-stained renal sections against iNOS](image)

**Figure (4):** A) Microscopic pictures of immune-stained renal sections against iNOS. IHC counterstained with Mayer's hematoxylin. Black arrows point to positive tubules. Low magnification
X:100 bar 100 and high magnification X:400 bar 50  B) Effect of Mirazid (10 mg/kg) orally for 7 days on Gentamycin (100 mg/kg; I.P)-induced change in inflammatory cells to count for H&E. Results are expressed as mean± S.E.M, n = 8. Statistical analysis was performed using One-Way ANOVA followed by Tukey-Kramer multiple comparisons test at P<0.05. *P< 0.05 concerning the control group, *P< 0.05 concerning the Gentamicin group.

Control group  Gentamicin group  Mirazid prophylactic group

**A**

Control group  Gentamicin group  Mirazid prophylactic group

**B**

IHC staining intensity scores of JNK-1 (0-3)
Figure (5): A) Microscopic pictures of immune-stained renal sections against JNK. IHC counterstained with Mayer's hematoxylin. Black arrows point to positive tubules. Low magnification X:100 bar 100 and high magnification X:400 bar 50 B) Effect of Mirazid (10 mg/kg) orally for 7 days on Gentamycin (100 mg/kg; I.P)-induced change in inflammatory cells to count for H&E. Results are expressed as mean± S.E.M, n = 8. Statistical analysis was performed using One-Way ANOVA followed by Tukey-Kramer multiple comparisons test at P<0.05. *P< 0.05 concerning the control group, #P< 0.05 concerning the Gentamicin group

3.7. Docking studies

Through the creation of five H-bonds with Met108, Glu109, and Met111 amino acids, the x-ray structure of JNK revealed the fitting of its co-crystallized inhibitor (TCU) inside its binding pocket. Also, it formed two extra H-bonds with Gln117 amino acid through an intermediate bridging H2O515 molecule. Molecular docking of the previously isolated compounds from myrrh resin compared to the docked co-crystallized inhibitor of JNK revealed the following descending binding order: oxalic acid, hexyl 2-methyl phenyl ester (12) > docked co-crystallized TCU inhibitor (18) > germacrene B (4) > (-)-e Ima-1,3,11(13)-trien-12-ol (2) > isosericenine (5) > 3-[(E)-2-phenyl-1-propenyl]cyclohexanone (6) > 2,5,8-trimethyl-1-nonen-3-yn-5-ol (7) ≥ 2-(2-hydroxy-2-methyl-2-phenylethyl)-3-methyl (14) > curzerene (3) > beta selinene (8) > spathulenol (9) ≥ 1-deoxycapsidiol (10) > myrcenol (15) > (-)-caryophyllene oxide (11) > (-)-(R)-ipsdienol (13) ≥ 2,8-decadiene (16) > R(+)-limonene (1) > bicyclo[3.1.1]hept-2-ene-2-carboxaldehyde,6,6-dimethyl-,(1S)- (17). Moreover, their binding scores and interactions with the amino acids of the JNK pocket are depicted in (Table 4) and supplementary data.

The findings of docking simulation showed that the docked co-crystallized TCU inhibitor (18) showed nearly the same binding mode of its native co-crystallized form, where it formed four H-bonds with Met108, Glu109, and Met111 amino acids. At the same time, it formed the two extra H-bonds with Gln117 amino acid through the intermediate bridging H2O515 molecule. Its binding score was found to be -6.95 kcal/mol and the RMSD value was 1.18. On the other hand, surprisingly, oxalic acid, hexyl 2-methyl phenyl ester (12) achieved a superior binding score (-7.19 kcal/mol) than the docked co-crystallized TCU inhibitor (18) with an RMSD value of 1.25. It got stabilized through the formation of only one H-bond with Asn114 amino acid which indicates a highly stabilized fitting of the molecule regardless of the formed amino acid interactions compared to the co-crystallized TCU inhibitor (docked, 18) as represented in (Tables 4 and 5). At the same time, germacrene B (4), (-)-e Ima-1,3,11(13)-trien-12-ol (2), and isosericenine (5) compounds showed very good binding scores which were very close to that of the docked co-crystallized TCU inhibitor (18) with score values of -6.34, -6.30, and -6.25, respectively (Table 4).
Table 4: Binding scores and interactions of the seventeen isolated and identified compounds from myrrh resin (1-17) compared to TCU (docked, 18) inside the binding pocket of JNK-1.

| No. | Compound                                                                 | $S^a$ | RMSD $^b$ | Amino acid bond                        | Length (Å) |
|-----|--------------------------------------------------------------------------|-------|-----------|----------------------------------------|------------|
| 1   | R(+) - Limonene                                                          | -4.96 | 1.51      | -                                      | -          |
| 2   | (-)-Elema-1,3,11(13)-trien-12-ol                                        | -6.30 | 0.98      | Met108/H-donor                          | 3.88       |
| 3   | Curzerene                                                               | -5.76 | 1.12      | Val40/pi-H                             | 4.20       |
| 4   | Germacrene B                                                            | -6.34 | 0.86      | -                                      | -          |
| 5   | Isosericenine                                                           | -6.25 | 1.28      | Met108/H-donor                          | 3.74       |
|     |                                                                          |       |           | Ser155/pi-H                             | 3.84       |
| 6   | 3-[(E)-2-phenyl-1-propenyl]cyclohexanone                                | -6.04 | 1.63      | -                                      | -          |
| 7   | 2,5,8-Trimethyl-1-nonen-3-YN-5-ol                                       | -6.00 | 1.43      | Asn114/H-donor                          | 2.98       |
|     |                                                                          |       |           | Ser155/H-acceptor                       | 3.29       |
| 8   | Beta selinene                                                           | -5.65 | 1.38      | -                                      | -          |
| 9   | Spathulenol                                                             | -5.63 | 1.08      | Ser155/H-acceptor                       | 2.80       |
| 10  | 1-Deoxycapsidiol                                                        | -5.63 | 1.13      | -                                      | -          |
| 11  | (-)-Caryophyllene oxide                                                 | -5.43 | 0.91      | Asn114/H-acceptor                       | 2.87       |
| 12  | Oxalic acid, hexyl 2-methylphenyl ester                                 | -7.19 | 1.25      | Asn114/H-acceptor                       | 2.90       |
| 13  | (-)-(R)-Ipsdienol                                                       | -5.30 | 2.08      | Ser155/H-acceptor                       | 2.99       |
|     |                                                                          |       |           | Asn114/H-acceptor                       | 3.15       |
| 14  | 2-(2-Hydroxy-2-methyl-2-phenylethyl)-3-methyl                           | -6.00 | 1.53      | Val40/pi-H                             | 4.15       |
| 15  | Myrcenol                                                               | -5.57 | 1.03      | Ser155/H-acceptor                       | 2.96       |
|     |                                                                          |       |           | Asn114/H-acceptor                       | 3.32       |
| 16  | 2,8-Decadiene                                                           | -5.26 | 0.55      | -                                      | -          |
| 17  | Bicyclo[3.1.1]hept-2-ene-2-carboxaldehyde, 6,6-dimethyl-,(1S)-         | -4.91 | 1.03      | Gln37/H-acceptor                       | 3.24       |
| 18  | Docked co-crystallized inhibitor                                        | -6.95 | 1.18      | Met111/H-donor                          | 2.95       |
|     |                                                                          |       |           | Gln117(H2O515)/H-acceptor               | 2.99       |
|     |                                                                          |       |           | Met111/H-acceptor                       | 3.03       |
|     |                                                                          |       |           | Gln117(H2O515)/H-donor                  | 3.04       |
|     |                                                                          |       |           | Glu109/H-donor                          | 3.19       |
|     |                                                                          |       |           | Met108/H-donor                          | 3.43       |

$^a$ $S$: the score of a compound inside the protein binding pocket (Kcal/mol),

$^b$ RMSD: The Root Mean Squared Deviation between the predicted pose and the crystal structure.
Table 5: 3D binding interactions and positioning between the most promising tested compound of myrrh resin (Oxalic acid, hexyl 2-methyl phenyl ester, 12) at the JNK-binding pocket compared to TCU (docked, 18).

The red dash represents H-bonds and the black dash represents H-pi interactions.

| Compound | 3 D interactions | 3 D positioning |
|----------|------------------|-----------------|
| 12       | ![Image of compound 12] | ![Image of positioning 12] |
| 18       | ![Image of compound 18] | ![Image of positioning 18] |

Regarding the docking results of the isolated tested compounds of myrrh resin compared to TCU, represented a good idea concerning their binding affinities towards JNK-1. Many isolates of the resin showed ideal and promising binding, which indicates high affinities and predicted intrinsic activities as JNK inhibitors as well.

Collectively, this study proposed the promising affinity of myrrh isolates against JNK-1, especially for oxalic acid, hexyl 2-methyl phenyl ester (12) which showed a superior binding affinity compared to the docked co-crystallized TCU inhibitor (18). Accordingly, we propose such a compound for further in vitro and in vivo studies to gain an effective anti-inflammatory and subsequently an apoptotic therapeutic against nephrotoxicity. Moreover, the previously studied isolates may be examined either alone or in combinations with each other's against nephrotoxicity.
4- Discussion

Mirazid's renal protective effect against experimentally generated nephrotoxicity was investigated in this study. Effects on inflammation, antioxidants, and apoptosis were discovered to be responsible for the renal protective effect, which was mainly due to a modulatory effect on JNK1/iNos pathway. For nephrotoxicity induction, a well-standardized experimental model was used.

Although the benefits of Gentamicin in reducing a wide range of bacterial infections, mostly Gram-negative bacteria, have been demonstrated\textsuperscript{73}. Gentamicin-induced renal toxicity which is a major clinical challenge to its wide therapeutic application\textsuperscript{1, 74, 75} Gentamicin administration raised serum creatinine and urea levels, as well as kidney/body weight, in the current study. In agreement with this result, through drug-induced free radical generation, gentamicin has been linked to the radical formation and oxidant injury in experimental models\textsuperscript{76}. Studies showed that serum urea and creatinine elevation is considered to be an important marker of renal dysfunction (glomerular damage marker)\textsuperscript{77}. The most sensitive markers for kidney disease detection in experimental trials were serum creatinine, blood urea nitrogen, and kidney weight/body index, which were all raised by gentamicin\textsuperscript{77, 78}.

A sensitive indicator of tubular injury is the LDH enzyme found in the proximal renal tubules\textsuperscript{79}. LDH activity was considerably higher in the Gentamicin group than in the control group. This increase can be explained by the fact that Gentamicin administration caused a change in redox status, which was demonstrated by a decrease in the concentration of glutathione and an increase in lipid peroxidation\textsuperscript{80}.

Inflammation causes a wide spectrum of inflammatory mediators to be released. It is characterized by tissue destruction and secretion of many inflammatory cytokines\textsuperscript{81, 82}. When these cells are activated, more cytokines such as nuclear factor kappa B (NF-κB) and iNOS are generated. iNOS causes inflammatory cells to migrate to the wounded area and generates cytokines such as NF-κB\textsuperscript{83}. Gentamicin caused peroxynitrite production by inducing the expression of iNOS in glomeruli and mesangial cells\textsuperscript{84}. Gentamicin-induced nephrotoxicity is also suggested as a result of nitric oxide(NO) overproduction\textsuperscript{85}. Lee et al. showed that NO is produced during inflammation\textsuperscript{86}. These findings corroborated the results of the current investigation, which indicated Gentamicin's nephrotoxicity.

The decline in antioxidant enzyme activity could be indicative of the negative impacts of Gentamicin. This is in alignment with Abdel-Zaher et al.\textsuperscript{87} when it comes to Probucol's effect in
preventing nephrotoxicity triggered by Gentamicin was studied in rats, and also in line with Pai, P.G., et al.\textsuperscript{88} where the protective action of ursolic acid is activated against Gentamicin nephrotoxicity was studied\textsuperscript{88}. The main mechanism by which Gentamicin mediates kidney injury is oxidative stress.

The level of MDA in the Gentamicin group was significantly greater than in the control group, whereas the content of GSH was significantly lower in the Gentamicin group than in the control group. These findings were in line with earlier research which reported that ROS has a significant impact on renal disease pathophysiology\textsuperscript{89}. Gentamicin increases ROS production \textit{in vivo} and \textit{in vitro} by modifying mitochondrial respiration. According to Khan et al., free radicals and ROS mediate polyunsaturated fatty acid peroxidation (PUFAs). An overabundance of PUFAs increases the kidneys' vulnerability to ROS\textsuperscript{90}. Biological membranes contain significant amounts of PUFAs that are especially vulnerable to lipid peroxide-producing peroxidative attacks\textsuperscript{91}.

The Gentamicin group had tubular degradation and necrosis, as well as mononuclear cell infiltration, as demonstrated in the histological image of the kidney tissues. The discovered histopathological renal changes as a result of nephrotoxicity induced by Gentamicin are consistent with findings from Kuatsienu et al.\textsuperscript{92} Alarifi et al. reported, necrosis, degeneration, and vacuolization were early signs of tubular changes caused by gentamicin treatment. By the conclusion of the gentamicin treatment, tubular abnormalities in the kidney had emerged, and their severity had grown. The majority of the proximal convoluted tubules and, to a lesser extent, the distal tubules were affected by degeneration up to severe necrosis\textsuperscript{93}.

The activation of the apoptotic pathway was linked to Gentamicin-induced kidney damage, implying that the apoptotic pathway was linked to renal damage caused by Gentamicin. According to the study, NF-κB activation enhanced Gentamicin-induced apoptosis in rat tubular cells \textsuperscript{94}. The considerable increases in caspase3 expression in renal cortical tissue revealed that Gentamicin produced endoplasmic reticulum (ER) stress and activation of ER-mediated cell death indicators in this investigation\textsuperscript{95}. One of the fundamental processes that provide protection and repair in stress-induced cellular dysfunction by inducing cell death is the activation of ER stress\textsuperscript{96}.

In the current study, Mirazid prophylactic showed a reno-protective effect against Gentamicin triggered nephrotoxicity, Gentamicin-induced nephrotoxicity was significantly improved, as seen by considerable reductions in (creatinine, urea, and LDH) levels and kidney weight. These findings support Hanan's findings that Mirazid therapy improved renal function in rats\textsuperscript{97}. Numerous experimental animal models have shown a relationship between oxidative stress and
nephrotoxicity. Furthermore, it has been demonstrated that treating rats with hydroxyl radical scavengers protects them against acute renal failure caused by Gentamicin.

Prophylactic treatment with Mirazid showed inhibition of kidney oxidative stress. It reduced MDA levels, meaning that it inhibited lipid peroxidation and the production of ROS, which was accompanied by increased GSH content, implying a significant boost in antioxidant defenses. Significant reductions in serum LDH activity accompanied these improvements in oxidative/antioxidant balance. Mirazid's capacity to prevent lipid peroxidation and dramatically improve the activity of antioxidant enzymes has been established in previous research to have a protective impact. Various studies have reported findings that are consistent with the ones presented here, Polyphenolic groups in myrrh extract induce a protective action against ROS. Due to its free radical-scavenging properties, it exhibited a preventive effect against stomach ulcers. It was recently discovered that myrrh, a powerful antioxidant, works by enhancing the total antioxidant activity of the serum and tissues. Meanwhile, Mirazid inhibited the production of MDA in Gentamicin induced renal cells. Improved antioxidant defense and reduced ROS generation can help maintain cellular integrity and provide structural, biochemical, and physiological benefits. Mirazid's antioxidant properties were associated with a significant decrease in Pathological alterations caused by gentamicin, as well as a return to normal metabolic equilibrium and cellular hemostasis. Mirazid has been proven to protect the nephrons, which is assumed to be related to its antioxidant effects and the antioxidants attenuate lipid peroxidation induced by Gentamicin.

In this study, there was an elevation in the expression of iNOS and JNK1. This can be explained by; Gentamicin administration has led to stimulation of inflammatory pathways through the upregulation of iNOS expressions. Furthermore, tumor necrosis factor-alpha (TNF-α) is a pro-inflammatory cytokine produced by glomerular and tubular cells, as well as extrinsic inflammatory cells, in response to Gentamicin treatment. TNF-α operates through the mitogen-activated protein kinase (MAPK) and NF-κB signaling pathways. Tissue damage and inflammation are important nephrotoxicity triggers. Furthermore, Gentamicin activates the signaling pathways of MAPK. JNK1 is one of three well-known MAPK pathways; it is regarded as a pro-inflammatory pathway. Cell proliferation, differentiation, migration, and apoptosis are also controlled. The effect of Gentamicin obtained in the current study can be explained by these findings. (Figure 6).
**Figure (6):** The protective role of Mirazid in Gentamicin-induced nephrotoxicity.

Gentamicin caused nephrotoxicity by increasing membrane lipid peroxidation and the formation of lipid aldehyde by stimulating ROS. JNK1 is one of three well-known MAPK pathways that activate protein kinase.; it is regarded as a pro-apoptotic and pro-inflammatory pathway. JNK activates the apoptosis pathway and the production of inflammatory markers. Gentamicin induced inducible nitric oxide synthase expression in glomeruli and mesangial cells and caused peroxynitrite production. This results in insult of inflammation and damage of the glomerulus (Nephrotoxicity). Mirazid possesses nephroprotective benefits due to its antioxidant, anti-inflammatory, and anti-apoptotic qualities.

Mirazid's anti-inflammatory and immune-modulating properties are confirmed by a reduction in renal iNOS content. In agreement with these results, the transcription factor NF-κB and MAPK that regulate the expression of many immune and inflammatory genes were inhibited by Mirazid, and several inflammatory mediators needed for initiation, maintenance of an inflammatory process, and reduction of oxidative stress were modulated by Mirazid. Because it regulates genes and coordinates the expression of pro-inflammatory enzymes and cytokines including iNOS and TNF-α, as well as interleukin-6, NF-κB is generally understood to be critical for cell viability (IL-6). In inflammatory and immune reactions, the NO radical is recognized to have a central role. iNOS is inactive in resting cells under normal physiological conditions, but it produces a large amount of NO under pathological conditions, resulting in interferon (IFN) and lipopolysaccharide (LPS).
increased endothelial nitric oxide synthase (eNOS) levels 10 times, which has a dual function in chronic infection, inflammation, and carcinogenesis\textsuperscript{114}.

Mitigation of Gentamicin-induced histological alterations confirm the anti-inflammatory impact of mirazid; of worth, choosing is the significant downregulation of apoptotic and inflammatory responses. Mirazid caused caspase-3 expression to decrease, indicating that apoptosis was retracted\textsuperscript{94}. In light of the situation, by inhibiting the activation of the ER stress and NF-\kappa B pathways, Mirazid protected cells from apoptosis. These mechanisms that have been proposed are backed up by prior research that found Mirazid is an effective free radical scavenger in the kidney inhibiting inflammatory signaling pathways and preventing gentamicin-induced renal toxicity\textsuperscript{50}.

5- Conclusion

Finally, by blocking the JNK1/iNOS pathway, Mirazid generates anti-inflammatory, anti-oxidant, and anti-apoptotic effects, as demonstrated in this study. Molecular docking was used to investigate seventeen identified chemicals from myrrh resin using GC–MS and ICP–MS separation techniques against JNK-1 which is responsible for stopping both the apoptotic and iNOS pathways and therefore proposed to be the main mechanism of action for its anti-inflammatory effects. Among the tested isolates oxalic acid, hexyl 2-methyl phenyl ester (12) was found to be superior to the docked co-crystallized inhibitor (18) with a binding score of -7.19 kcal/mol compared to -6.95, respectively. Our findings could help in the exact identification of the main constituents of myrrh responsible for its anti-inflammatory effects by targeting JNK-1. Especially oxalic acid, hexyl 2-methyl phenyl ester (12), germacrene B (4), (-)-elema-1,3,11(13)-trien-12-ol (2), and isosericenine (5) isolates represent the most promising compounds for further preclinical and clinical studies for the treatment of inflammation.
Author Contributions

Conceptualization: SAA, AEK; Data curation: AAA, MM, MAE, and AEK; Formal Analysis: AAA, MM, MAE, SS, and AEK; Methodology: AAA, AM, MM, MAE, SS, and AEK; Project administration: AAA and AEK; Resources: AAA and AEK; Supervision: AAA and AEK; Validation: SAA, AAA, and AEK; Visualization: AAA, AM, SS, and AEK; Writing – original draft: SAA, AAA, AM, SS, and AEK; Writing– review and editing: SAA, AAA, and AEK. All authors approved the final version of the manuscript.

Conflict of interest

The authors declare no conflict of interest.

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