Molecular Properties, Bioactivity Scores, and Toxicity Predictions of the Phytoconstituents Present in Bauhinia Acuminata

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Abstract:
To develop the herbal drug with the least side effects, there are superior opportunities to discover the medicinal and other biological properties. Natural products serve as sources of beneficial chemical molecules. For this study, Bauhinia acuminata an important medicinal plant of the Indian subcontinent that belongs to the family Fabaceae was chosen. The plant is well known for its precautionary action in tuberculosis. It has been established to possess some pharmacological activities such as membranes Stabilizing activity1, antibacterial2, anti-nociceptive3, thrombolytic activity4, antioxidant5, anthelmintic6, anti-diarrheal7, Hepato-protective 8. Phytoconstituents present in Bauhinia acuminata obey Lipinski's rule (MiLog P <5) except Kaempferol-3-glucoside indicated their drug-likeness property. Rheoagenine, 9, 12, 15-octadecatrienoic acid, and 9, 12-octadecadienoic acid are the phytoconstituents showing all types of binding with all types of receptors binding except Kinase inhibitor activity. Rheoagenine, Alpha humulene, 9, 12, 15-octadecatrienoic acid, 9, 12-octadecadienoic acid, Alpha muurolol, Beta-sitosterol, Kaempferol-3-glucoside are the phytoconstituents that are free from any type of toxicity. The accurate prediction scores can be used as monographs by researchers and scientists for the development of potential Semisynthetic and synthetic drugs for multifarious usage.

Keywords: Bauhinia acuminata, Molinspiration software, Osiris software, Toxicity profile, Phytoconstituents, Bioactivity score.

I. Introduction:
The prehistoric people have great consciousness of the tradition of medicinal plants as herbal medicines. In the world, more than 80% of the living in minor developed countries reveals on customary medicine and humans are dependent on herbs for their basic requirements such as foodstuffs, clothing, flavor, shelter, fragrance, and medicines (Divya and Mini, 2011 & Manoj Kumar Mishra, 2016, Gurib-Fakim, 2006 and Brijesh & Madhusudan, 2015). The Discovery of drugs in medicinal plants affords better and vital leads, besides diverse pharmacological activities such as cytotoxic, anti-diarrheal, antimicrobial, anti-inflammatory, antioxidant, anthelmintic, anti-nociceptive, hemolytic activity. The plant is well known for its precautionary action in tuberculosis. As per the recommendations of Ayurveda Bauhinia acuminata is the one of important medicinal plants for the treatment of disorders. (Yi F et al, 2016)9. Bauhinia acuminata belonging to the family of Fabaceae is an evergreen large shrub grows in the areas of Southeast Asia such as Malaysia, Indonesia or Philippines. For conventional drugs, bark, leaves, stem, blooms, and Roots have been utilized. Chemical constituents present in Bauhinia acuminata leaves are palmitic acid, three phallic acid esters, gallic acid, and ursolic acid. The leaves and stems of B. acuminata have shown the presence of carbohydrate, saponins, phenolic compounds, flavonoids, oils, and fats, alkaloids, anthocyanoside, steroids, anthraquinone,
terpenoids, resins, amino acid, sugars, and cardiac glycosides. In phytochemical screening, leaf oil identified 13 compounds in B. acuminata through GC-MS analysis are Quercetin, Neophytadiene, Rhoeagenine, Alpha humulene, Isoaromadendrene epoxide (Vasudevan et al.,2013), Butanedioic acid diethyl ester, 9,12,15-octadecatrienoic acid, Beta-ionone, 9,12-octadecadienoic acid, Alpha muurolol, Bauhinione, Beta-sitosterol, Kaempferol-3-glucoside. Phytochemical plant extracts when screened have shown the presence of saponin, alkaloid, cardiac glycosides, flavonoids, and tannin and steroid compounds (Dongray et al, 2016) 10.

Molinspiration
i) Lipinski’s rule
Lipinski's rule of five also known as Pfizer's rule of five or simply the Rule of five (RO5) is a rule of thumb to evaluate drug-likeness or determine if a chemical compound with a certain pharmacological or biological activity has properties that would make it a likely orally active drug in humans. The rule was formulated by Christopher A. Lipinski in 1997. The rule describes molecular properties important for a drug's pharmacokinetics in the human body, including their absorption, distribution, metabolism, and excretion ("ADME") Components of Lipinski’s rule:

ii) Lipinski’s rule states
• A molecular mass of fewer than 500 daltons.
• An octanal-water partition coefficient log P not greater than 5.
• Not more than 10 hydrogen bond acceptors (nitrogen or oxygen atoms).
• Not more than 5 hydrogen bond donors (nitrogen or oxygen atoms with one or more hydrogen atoms).
• No more than one violation.

iii) Drug likeness score
Molinspiration is web-based software which can be used to obtain parameters such as MiLog P, drug-likeness scores, TPSA. MiLog P is calculated by the methodology developed by Molinspiration which is a sum of fragment-based contributions and correction factors. Good permeability across the cell membrane can be checked with MiLog P value. Log P or Partition coefficient is an important parameter used to measure molecular hydrophobicity in rational drug design. The hydrophilic/lipophilic nature of drug molecules affects drug absorption, bioavailability, drug-receptor interactions, as well as their toxicity. Based on a sum of fragment contributions of O and N- centered polar fragments Molecular Polar Surface Area TPSA is calculated. Total polar surface area (TPSA) is closely related to the hydrogen bonding potential of a molecule and is a very good predictor of drug transport properties such as intestinal absorption, bioavailability, blood-brain barrier penetration, etc. Based on group contributors, the calculation of volume were developed at Molinspiration The number of rotatable bonds measures molecular flexibility. It is a very good descriptor of the absorption and bioavailability of drugs. Molecular properties and structure features with respect to known drugs can be checked through drug-likeness data of the molecule. MiLog P (partition coefficient), molecular weight, number of heavy atoms, number of hydrogen donors, number of hydrogen acceptors and number of violations, number of rotatable bonds, and volume 11 are the parameters considered for calculating the drug-likeness scores.

iv) Bioactivity score
Bioactivity of the drug can be checked by calculating the activity score of GPCR ligand, ion channel modulator, nuclear receptor legend, a kinase inhibitor, protease inhibitor, enzyme inhibitor. Calculated drug-likeness score of each compound and compared with the specific activity of each compound, and the results were compared with standard drug. All the parameters were checked with the help of the software Molinspiration drug-likeness score online (www.molinspiration.com). The probability for the organic molecules is if the bioactivity score is (>0), then it is active, if (-5.0-0.0) then moderately active, if (< -5.0) then inactive

II. Materials And Methods
i) Materials:
Phytochemical compounds present in Bauhinia acuminata like Quercetin, Neophytadiene, Rhoeagenine, Alpha humulene, Isoaromadendrene epoxide (Vasudevan et al.,2013), Butanedioic acid diethyl ester, 9,12,15-octadecatrienoic acid, Beta-ionone, 9,12-octadecadienoic acid, Alpha muurolol, Bauhinione, Beta-
sitosterol, Kaempferol-3-glucoside which are listed in **Table 1** were selected for Insilco prediction and the smile notations of used to generate the data.

**ii) Methods:**

Molinspiration Software: Structures of 13 phytochemical compounds selected for our work as given in Table 1 (reported in the literature resources) were drawn using online molinspiration for the calculation of molecular properties like MiLog P. Total polar surface area (TPSA), number of hydrogen bond donors and acceptors, molecular weight, number of atoms, number of rotatable bonds, etc., and bioactivity scores like GPCR ligands, kinase inhibitors, ion channel modulators, enzymes, and nuclear receptors. The molecular properties and bioactivity scores predicted by molinspiration were given in **Tables-2 and 3.**

For organic molecules, the probability is if the bioactivity scores (>0), then it is active, if (-5.0- 0.0) then moderately active, if (<-5.0) then inactive.

**Osiris software:** ([https://www.Osiris.com](https://www.Osiris.com)) This software predicts the toxicity risk assessment and calculates on-the-fly various drug-relevant properties like cLogP, solubility, Molecular Weight, Toxicity Risk Assessment, Overall Drug-Score, etc. When the structure is valid Prediction results are valued and color-coded like properties with a high risk of undesirable effect shown in red whereas green color indicates drug-conform behavior.

**III. Results**

**Table-1: General properties of Phyto-constituents of Bauhinia acuminata.**

| PHYTO-CONSTITUENT       | MOLECULAR FORMULA | SMILE NOTATION                                      | STRUCTURE |
|--------------------------|-------------------|-----------------------------------------------------|-----------|
| Quercetin                | C_{15}H_{10}O_{7} | C1=CC(=C(C=C1C2=C(C(=O)C3=C(C=C(C3O2)O)O)O)O)O       |           |
| Neophytadiene            | C_{20}H_{40}O_{0} | CC(C)CCCC(C)CCCC(C)CCCCC(C)CCCCC=C C=C            |           |
| Rhoeagenine              | C_{14}H_{17}NO_{6} | CN1CCC2=CC3=C(C=C24C1C5=C(C(O4)O)C6=C(C=C5)OCO6)O CO3 |           |
| Alpha humulene           | C_{15}H_{24}O_{4} | CC1=CCC(C=CCC(=CCC1C)C)C                                    |           |
| Isoaromadendrene epoxide | C_{14}H_{34}O_{2} | CCCC2C1C3C(C(C)C)CC4C2(O4)C                                   |           |
| Butanedioic acid diethyl ester | C_{8}H_{14}O_{4} | CCOC(=O)CCC(=O)OCC                                          |           |
| 9,12,15-octadecatrienoic acid | C_{18}H_{28}O_{2} | CCCCCC=CCC=CCCC=CCCCC(=O)O                                   |           |
| Beta-ionone              | C_{13}H_{20}O_{4} | C1=C(C(CCC1)C)C=CC(=O)C                                    |           |
| 9,12-octadecadienoic acid | C_{18}H_{28}O_{2} | CCCCCC=CCC=CCCCCCCCCC(=O)O                                   |           |
Table 2: Computed molecular properties of the phytoconstituents present in Bauhinia acuminata.

| Phyto-constituent     | Mi log p | TPSA     | natoms | MW    | nON | nOHNH | n violations | nrotb | volume  |
|-----------------------|----------|----------|--------|-------|-----|--------|--------------|-------|---------|
| Quercetin             | 1.68     | 131.35   | 22     | 302.24| 7   | 5      | 0            | 1     | 240.08  |
| Neophytadiene         | 7.55     | 0        | 20     | 278.52| 0   | 0      | 1            | 13    | 336.04  |
| Rheoeagenine          | 2.34     | 69.64    | 27     | 369.37| 7   | 1      | 0            | 0     | 312.06  |
| Alpha humulene        | 5.3      | 0        | 15     | 204.36| 0   | 0      | 1            | 0     | 234     |
| Isoaromadendrene epoxide | 4.27   | 12.53    | 16     | 220.36| 1   | 0      | 0            | 0     | 228.88  |
| Butanedioic acid diethyl ester | 0.71 | 52.61   | 12     | 174.22| 4   | 0      | 0            | 7     | 168.9   |
| 9,12,15-octadecatrienoic acid | 6.37  | 37.3     | 20     | 278.44| 2   | 1      | 1            | 13    | 306.47  |
| Beta-ionone           | 3.45     | 17.07    | 14     | 192.32| 1   | 0      | 0            | 2     | 208.76  |
| 9,12-octadecadienoic acid | 6.86    | 37.3     | 20     | 280.45| 2   | 1      | 1            | 14    | 312.65  |
| Alpha muurolol        | 4.97     | 20.23    | 16     | 222.37| 1   | 1      | 0            | 1     | 243.65  |
| Bauhinione            | 2.77     | 52.61    | 21     | 284.31| 4   | 0      | 0            | 2     | 256.42  |
| Beta-sitosterol       | 8.62     | 20.23    | 30     | 414.72| 1   | 1      | 1            | 6     | 456.52  |
| Kaempferol-3-glucoside | 0.12    | 190.28   | 32     | 448.38| 11  | 7      | 2            | 4     | 364.19  |

Table 3: Computed Bioactivity scores of the Phyto-constituents of Bauhinia acuminata by Molinspiration

| PHYTO-CONSTITUENT             | GPCR LIGAND | ION CHANNEL MODULATOR | KINASE INHIBITOR | NUCLEAR RECEPTOR LIGAND | PROTEASE INHIBITOR | ENZYME INHIBITOR |
|-------------------------------|------------|-----------------------|------------------|-------------------------|--------------------|------------------|
| Quercetin                     | -0.06      | -0.19                 | 0.28             | 0.36                    | -0.25              | 0.28             |
| Neophytadiene                 | -0.12      | -0.02                 | -0.35            | 0.2                     | -0.11              | 0.14             |
| Rheoeagenine                  | 0.21       | 0.13                  | -0.14            | -0.09                   | 0.02               | 0.17             |
| Alpha humulene                | -0.14      | 0.02                  | -0.93            | 0.34                    | -0.67              | 0.31             |
| Isoaromadendrene epoxide      | -0.39      | -0.37                 | -0.69            | -0.01                   | 0.02               | -0.05            |
| Butanedioic acid diethyl ester| -0.93      | -0.35                 | -1.19            | -0.91                   | -0.92              | -0.46            |
| 9,12,15-octadecatrienoic acid | 0.29       | 0.17                  | -0.16            | 0.31                    | 0.12               | 0.38             |
| Beta-ionone                   | -0.9       | -0.26                 | -1.34            | 0.25                    | -0.79              | 0.28             |
| 9,12-octadecadienoic acid     | 0.29       | 0.17                  | -0.16            | 0.31                    | 0.12               | 0.38             |
| Alpha muurolol                | -0.09      | 0.05                  | -0.87            | 0.39                    | -0.63              | 0.4              |
| Bauhinione                    | -0.13      | -0.18                 | 0.1              | 0.06                    | -0.29              | 0.34             |
| Beta-sitosterol               | 0.14       | 0.04                  | -0.51            | 0.73                    | 0.07               | 0.51             |
| Kaempferol-3-glucoside        | 0.06       | -0.05                 | 0.1              | 0.2                     | -0.05              | 0.41             |

Table 4: Computed Toxicity profile of the Phyto-constituents of Bauhinia acuminata by Osiris software

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IV. Discussion

Table 1: General properties and structures of the phytoconstituents present in Bauhinia acuminata

Table 2 and 3: Molinspiration (molecular properties and Bioactivity scores)

a) Molecular property of the phytochemical compounds
Among the all the 13 phytochemicals screened for Insilco drug activity prediction which are done by molinspiration showed that all the phytochemical obeyed the Lipinski’s rule of five and reflected good drug-likeness scores. The values of MiLog P for these compounds were found to be < 5 indicating their good permeability across the cell membrane.

All the phytochemicals found to have molecular weight <500, TPSA well below 160Å2, Number of hydrogen bond donors < 5, hydrogen bond acceptors < 4, n-violations 0, and number of rotatable flexible bonds <5. The observed data indicated that except Kaempferol-3-glucoside all the phytoconstituents were found to have drug-likeness property.

b) Bioactivity scores of the compounds
For organic molecules, the probability is if the bioactivity scores (>0) then it is active, if (-5.0 - 0.0) then moderately active, if (<-5.0) then inactive.

The bioactivity scores of the 13 phytoconstituents observed were as follows:

i) GPCR Ligand
Rhoeagenine, 9, 12-octadecadienoic acid, Beta-sitosterol, Kaempferol-3-glucoside are showing good GPCR binding activity

ii) Ion channel modulator
Rhoeagenine, Alpha humulene, 9, 12, 15-octadecatrienoic acid, 9, 12-octadecadienoic acid, Alpha muurolol, Beta-sitosterol are showing good ion channel modulator activity.

iii) Kinase inhibitor
Quercetin, Bauhinione, Kaempferol-3-glucoside are having good Kinase Inhibitor activity

iv) **Nuclear receptor Ligand**
Rhoeagenine, Isoaromadendrene epoxide, Butanedioic acid diethyl ester all the other Phytoconstituents are showing good Nuclear receptor binding activity

v) **Protease inhibitor**
Rhoeagenine, Isoaromadendrene epoxide, 9, 12, 15-octadecatrenic acid, 9, 12-octadecadienoic acid, Beta-sitosterol are showing good Protease inhibitor activity.

vi) **Enzyme inhibitor**
Except for Isoaromadendrene epoxide, Butanedioic acid diethyl ester all the phytoconstituents are showing good enzyme binding activity.

Rhoeagenine, 9, 12, 15-octadecatrienoic acid, and 9, 12-octadecadienoic acid are the phytoconstituents showing all types of binding with all types of receptors except Kinase inhibitor activity.

**Table 4: Osiris (Toxicity prediction)**

Zero and positive values indicate good drug score and drug likeliness. Drug score and drug likeliness are good for Rhoeagenine

**Mutagenicity:** Quercetin showing Mutagenicity

**Tumorogenicity:** Quercetin, Isoaromadendrene epoxide, Beta-ionone are showing tumorogenicity

**Skin and eye Irritation:** Isoaromadendrene epoxide, Beta-ionone, Butanedioic acid diethyl ester are showing skin and Eye Irritation

**Reproductive effects:** Beta-sitosterol is showing Reproductive effects.

Rhoeagenine, Alpha humulene, 9, 12, 15-octadecatrienoic acid, 9,12-octadecadienoic acid, Alpha muurolol, Beta-sitosterol, Kaempferol-3-glucoside are the phytoconstituents that are free from any type of toxicity.

**Conclusion**
In the present study, we used Molinspiration, Osiris online software tool which is available free for the users to evaluate the molecular properties, bioactivity scores, toxicity predictions of the phytoconstituents present in Bauhinia acuminata. The phytoconstituents of the plants were enlisted through the software includes Quercetin, Neophytadiene, Rhoeagenine, Alpha humulene, Butanedioic acid diethyl ester, 9,12,15-octadecatrienoic acid, Beta-ionone, 9,12-octadecadienoic acid, Alpha muurolol, Bauhinione, Kaempferol-3-glucoside. Accordingly, the respective software data for the phytoconstituents are computed and depicted in respective tables. Further, these values can be used as monographs by researchers and scientists for the development of potential Semisynthetic and synthetic drugs for multifarious usage.

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“Not applicable”.

**Conflicts of Interest**

There are no conflicts of interest.

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