# *In vitro* urease inhibition screening of some edible and medicinal herbs to combat *Helicobacter pylori* related gastric diseases

# Hina Zahid<sup>1</sup>, Bushra Hina<sup>2</sup> and Khizar Abbas<sup>3\*</sup>

<sup>1</sup>Department of Pharmacognosy, Dow College of Pharmacy, Dow University of Health Sciences, Karachi, Pakistan <sup>2</sup>Department of Pharmacognosy, Institute of Pharmaceutical Sciences, Jinnah Sindh Medical University, Karachi, Pakistan <sup>3</sup>Department of Pharmacognosy, Faculty of Pharmacy, Bahauddin Zakariya University, Multan, Pakistan

**Abstract**: Prevalence of gastric diseases caused by *Helicobacter Pylori* bacteria is very common especially in developing countries. *H. pylori* is not only responsible for initiating gastric complaints like gastritis and peptic ulcer but may also lead to gastric cancer. The aim of this research study is to explore natural flora that exhibit anti urease potential. For this purpose, fifteen edible and medicinal herbs (*Silybium marianum*, *Fagonia arabica*, *Nigella sativa*, *Curcuma longa*, *Ocimum sanctum*, *Salvia rosmainus*, *Hyssopus officinalis*, *Antrirrhinum majus*, *Salvia splendens*, *Tropaeolum majus*, *Dalbergia sisso*, *Aloe barbadensis*, *Abelmoschus esculentus*, *Cuscuta reflexa*, *Hibiscus schizopetalus*) were screened for anti-urease activity at three different concentration i.e,  $25\mu$ g/ml,  $50\mu$ g/ml and  $75\mu$ g/ml. The results indicated significant outcomes for urease inhibitory activity for all tested medicinal plants. However, *F. arabica* ( $87.2\pm1.47$ ), *N. sativa* ( $90.4\pm0.09$ ), *O. sanctum* ( $75.6\pm0.95$ ), *H. officinalis* ( $78.9\pm0.69$ ), *T. maju* ( $87.3\pm0.14$ ), *A. esculentus* ( $90.3\pm0.86$ ), *C. reflexa* ( $94.1\pm0.92$ ) showed significant results at  $75\mu$ g/ml when compared to Thiourea. Moreover IC <sub>50</sub> values were also calculated for urease inhibitory activity. It can be concluded that utilization of these valuable medicinal plants can not only decrease the prevalence of gastric diseases caused by *H. pylori* bacteria but a good candidate for therapeutic purposes.

Keywords: Herbs, GIT diseases, peptic ulcer, stomach cancer, urease inhibitory activity.

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# **INTRODUCTION**

Prevalence of gastric diseases caused by H. pylori bacteria is very common worldwide. Its manifestations start from gastritis, peptic ulcer and may lead to stomach According World Gastroenterology cancers. to Association (WGA) Helicobacter pylori has been a major disease-causing human pathogen for humanity since last four decades. H. pylori infection is a key reason of morbidity and mortality infecting about half of the world's population via spread of peptic ulcer diseases and gastric cancer. It is very challenging situation for health care professionals and researchers to prevent, treat and control the spread of diseases caused by H. pylori due to difference in spectrum of diseases as well as wide worldwide variation of suffering population (WGO, 2021; Chan and Lau, 2021). H. pylori infection is related to socioeconomic status and lifestyle pattern. Well-resourced population of developed countries is at lower risk, while prevalence of infection remains high in the regions of world that belong to developing countries where resources are limited (Cover and Blaser, 2020).

Survival of *H. pylori* in the highly acidic environment of stomach is mainly dependent on a nickel and enzyme Urease present in bacterial cytoplasm. Urease is responsible for metabolizing urea into ammonia and carbon dioxide adjusting the pH necessary for bacterial

\*Corresponding author: e-mail: Khizarabbas@bzu.edu.pk

survival (Amin et al., 2013). The schematic representation of urea hydrolysis by urease is depicted in scheme 1 (Yuri et al., 2018). The current regime of treatment of digestive diseases caused by H. pylori infection is based on triple therapy including two broadband antibiotics and proton pump inhibitor. However conventional therapy regimens of H. pylori are becoming complicated and ineffective due to antibiotic resistance as is counted in one of 16 antibiotic resistant dangerous bacterial strains (Suzuki et al., 2010; Cunha et al., 2021). Apart from bacterial resistance, high therapeutic cost and complex regimen of multiple daily doses makes the treatment unapproachable. As gastric infections caused by H. Pylori are highly prevalent around the globe as well as communicable through oral to oral and oral to fecal transmission, there is an urgent need to find a highly specific and targeted remedy (Sharaf et al., 2022).

In current scenario a remarkable trend is noticed to explore nature as search engine for the discovery of phytochemicals having great therapeutical potentials. A lot of medicinal plants have been proved to have antibacterial properties. Keeping this fact in mind it can be believed that *H. pylori* gram negative bacteria responsible for digestive problems can also be eradicated with natural herbs. Main objective of this research study is to explore natural flora to identify and authenticate medicinal plants having anti-Urease activity. Blocking the Urease enzyme will result in eradication of bacteria hence protect from all complications like peptic ulcer and

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#### Scheme 1: Urea hydrolysis catalyzed by ureases.

cancers. In this regard we selected 15 medicinal plants Silybium marianum, Fagonia arabica, Nigella sativa, Curcuma longa, Ocimum sanctum, Salvia rosmainus, Hyssopus officinalis, Antrirrhinum majus, Salvia splendens, Tropaeolum majus, Dalbergia sisso (flowers), Dalbergia sisso (leaves), Aloe barbadensis, Abelmoschus esculentus (aerial parts), Abelmoschus esculentus (seeds), Cuscuta reflexa, Hibiscus schizopetalus) for anti-urease screening.

# MATERIALS AND METHODS

#### Material collection and identification

The selected 15 medicinal plants 100 gm each were purchased from the local herbal market of Karachi city and identified and authenticated by Prof. Dr. Ghazala H Rizwani, Department of Pharmacognosy, Faculty of Pharmacy Hamdard University. Botanical descriptions and Pharmacognostic profiles of all tested plants are mentioned in table 1.

#### Extract preparation

All plant materials were garbled, washed and shade dried individually. Extraction of each plant was performed separately by soaking in ethanol for 15 days at room temperature. It was then filtered through whatman filter paper No 1, followed by solvent evaporation under reduced pressure using rotary evaporator (Buchi, Switzerland), at a temperature of 46°C to give a residue (extract). After complete solvent evaporation, each of 15 extracts were weighed and stored in airtight bottles for further use. The % yield of extracts obtained were Silvbium marianum (7.2%), Fagonia arabica (5.9%), Nigella sativa (7.8%), Curcuma longa (6.6%), Ocimum sanctum (6.3%), Salvia rosmainus (4.3%), Hyssopus officinalis (9.8%), Antrirrhinum majus (5.6%), Salvia splendens (4.5%), Tropaeolum majus 3.9%, Dalbergia sisso (flowers) (6.0%), Dalbergia sisso (leaves) (10.0%), Aloe barbadensis (5.0%), Abelmoschus esculentus (aerial parts) (7.5%), Abelmoschus esculentus (seeds) (6.1%), *Cuscuta reflexa* (5.1%), *Hibiscus schizopetalus* (8.02%)

## Chemicals and reagents

All chemicals and reagents used were Analytical grade. Canavalia ensiformis (Avonchem Ltd, UK), Sodium Hypochlorite (HC Haq Chemicals, Pakistan), Sodium Nitroprusside (BDH Chemicals Ltd, England), Thiourea (Merck, Germany), Urea (Sigma-Aldrich).

## Anti-Urease activity

#### Sample preparation and incubation

Each of the extract weigh accurately (1 gm) and dissolve in solvent to prepare litre of solution. Then take 1 ml of it and dilute again to one litre to get  $\mu$ g/ml concentration. To evaluate the urease inhibitory activity of different medicinal plants the exact 20µl of enzyme (Canavalia ensiformis- Jack Bean) solution mixed with 55ul of phosphate buffer (0.2M) in each extract the mixture having pH of 7.4 (Weatherfourn, 1967). Then, the mixture was kept on incubation for 10 minutes at 30°C, later add 15µl urea and kept it again for incubation using the same time and temperature. After incubation, the known volume of each extract of medicinal plant (25µg/ml, 50µg/ml and 75µg/ml) respectively were added in 96 wells. Now add 25µg/ml of Thiourea (standard solution). Properly mixed it and kept it for 10 minutes at 37 °C. Then add 40µl Alkali reagent (0.5% w/v NaOH and 0.1% active chloride NaOCl) and 60µl phenol reagent (1% w/v phenol and 0.005% w/v sodium nitroprusside) in all wells and leave it at room temperature for 10 minutes (Khan et al., 2015). Experiments were performed in a triplicate fashion and thiourea was used as standard inhibitor.

#### Absorbance using UV spectrophotometer

Measure the absorbance spectrophotometric at 625 nm. Urease inhibition percentage was calculated by using the mentioned formula:

% inhibition = [1- (A<sub>625</sub> of sample/A<sub>625</sub> of control)  $\times$  100]

#### IC 50 value calculations

 $IC_{50}$  values of all medicinal plants and standard Thiourea were calculated by plotting graph between % urease inhibitory activity verses doses/concentration of extracts, then fitting the data with a straight trendline. Linear regression Equation thus obtained was used to  $IC_{50}$ determination.  $Y = M^* X + C$ 

# STATISTICAL ANALYSIS

Anti-urease activity of all 15 medicinal plants along standard thiourea is represented as mean  $\pm$  S.E.M. In order to find the significant differences in activity among them One way ANOVA was employed followed by Tukey's posthoc test at *p*<0.05. SPSS version 20 was used for statistical calculations.

## RESULTS

Anti-urease activity of ethanolic extracts of all 15 medicinal plants (three different doses 25, 50 and 75µg/ml) has been analyzed using UV spectrophotometer keeping thiourea as standard. Fig. 1a, 2a, 3a showed the graphical representation of % urease inhibition of extracts of medicinal plants and standard thiourea at the concentration of 25, 50 and 75µg/ml respectively. However, *F. arabica* (87.2±1.47), *N. sativa* (90.4±0.09), *O. sanctum* (75.6±0.95), *H. officinalis* (78.9±0.69), *T. maju* (87.3±0.14), *A. esculentus* (90.3±0.86), *C. reflexa* (94.1±0.92) showed significant results at 75µg/ml when compared to Thiourea.

After estimation of % urease inhibitory activity of all plant extracts,  $IC_{50}$  was calculated that is shown in table 2.  $IC_{50}$  value of standard thiourea was found 35.6µl. Extracts having excellent  $IC_{50}$  values were *Cuscuta reflexa* (35.4µg/ml), *Aloe barbadensis* (36.5µg/ml), *Nigella sativa* (37.1µg/ml), *Fagonia arabica* (39µg/ml), *Tropaeolum maju* (40.17µg/ml), *Abelmoschus esculentus* (seed 40.4µg/ml), *Ocimum sanctum* (41µg/ml), *Abelmoschus esculentus* (Aerial parts 41.2µg/ml) and *Hyssopus officinalis* (42µg/ml).  $IC_{50}$  values of medicinal plant extracts and standard Thiourea for Anti-urease activity were showed in table 2. Comparison of three different concentrations (25ug/ml, 50ug/ml and 75ug/ml) was depicted in fig. 2.

# DISCUSSION

*H. pylori* is a causative bacteria responsible for widespread spectrum of gastric diseases including multifocal atopic gastritis, stomach cancers and peptic ulcers. Stomach cancer is listed as the  $3^{rd}$  main cause of morbidity among population due to cancer and behind 90% of such cases the causative organism was detected as *H. pylori* (Cunha *et al.*, 2021). It is the need of time to explore natural flora to find medicinal and edible plants that are capable of destroying the colonies of this gramnegative bacterial pathogen in the stomach. Our current research study is designed to investigate some common edible as well as medicinal herbs for their urease inhibitory effect so that increasing morbidity and mortality due to *H. pylori* can be minimized and controlled.

It is very important to understand the mechanism of *H. pylori* survival to eradicate the gastric diseases caused by this bacterium. The bacterial mechanism behind survival in highly acidic environment of stomach has been explained by various researchers. It can be concluded from literature that Urease is the key bacterial enzyme composed of two genes *Ure* A and *Ure* B, responsible for increasing the pH of stomach by liberating ammonia. Moreover, gene transcription and reduced mucin viscoelasticity also help of *H. pylori* in adherence to the

surface epithilum cells of stomach promoting vacuolating cytotoxic activity and growth- inhibitory factor for stomach cell proliferation (Smoot, 1997; Cellia et al., 2009; Graham and Miftahussurur, 2018; Khan et al., 2017). This available scientific data on mechanism of  $H_{\rm c}$ pylori survival suggests inhibiting the urease enzyme in the stomach hence maintaining the lower acidic pH resulting in bacterial death. In our research study we have explored fifteen medicinal and edible plants from Pakistan for their anti-urease potential to combat gastric diseases caused by H. pylori. Utilization of these plants will inhibit the bacterial urease enzyme resulting in bactericidal effect. Analytical results indicate that all plants showed positive anti-urease activity in vitro. Among the tested samples some plants such as Nigella sativa (Black cumin), Ocimum sanctum (basil), Abelmoschus esculentus (lady finger), Salvia rosmainus (rosemary) and Curcuma longa (turmeric) are edible.

These plants are easily available and their incorporation in daily cuisine of general population may not only help in spread of gastric diseases but also fighting against *H. pylori* via inhibition of bacterial urease. On the other hand, medicinal plants like *Silybum marianum*, *Fagonia arabica*, *Hyssopus officinalis*, *Antrirrhinum majus*, *Salvia splendens*, *Tropaeolum majus*, *Dalbergia sisso*, *Aloe barbadensis*, *and Cuscuta reflexa* has been proved as good candidates to be employed for therapeutic use among patients of gastric diseases.

Upon comparison of extracts with standard thiourea, it was concluded that most of the tested medicinal plants (*Fagonia arabica, Nigella sativa, Ocimum sanctum, Hyssopus officinalis, Tropaeolum maju, Abelmoschus esculentus* (Aerial parts), *Abelmoschus esculentus* (seed), *Cuscuta reflexa*) showed significant urease inhibitory activity at all doses of 25µg/ml, 50µg/ml and 75µg/ml. On the other hand, *Silybum marianum, Aloe barbadensis, Hibiscus schizopetalus* showed moderate activity and found statistically significant at doses of 50µg/ml and 75µg/ml, while flowers of *Dalbergia sisso* was found significant at dose of 75µg/ml when compared to thiourea.

However. Curcuma longa. Salvia rosmainus. Antrirrhinum majus, Salvia splendens although showed anti-urease activity at all doses but not as significant as Thiourea. Regarding mechanism it is evident from results that urease enzyme inhibitory activity of all medicinal plants is dependent on concentration, and it is increasing in dose dependent manner (Figs. 1a, 1b and 1c). Our research findings suggest that the use of edible herbs having anti urease potential in daily routine can minimize the spread of H. pylori infections. Moreover, natural flora can serve as search engine for the discovery of new anti H. pylori drugs due to the presence of active phytochemicals having therapeutic potential. Extraction, isolation and structure elucidation of such active constituents from these herbs can serve as model for developing their synthetic analogue.

S. No	Scientific names	Local names	Family	Part	Phytochemicals	Therapeutic uses
1.	Silybium marianum	Milk thistle	Asteraceae	Flower	Silybin, silidianin, Silicristin	Liver disorder, hepatitis, cirrhosis, jaundice. diabetes. Indigestion (Wang <i>et al.</i> , 2020)
2.	Fagonia arabica	Dhamasa/ sachi booti	Zygophyllaceae	Aerial part	Glycoside, flavonoids, terpenoids, Saponin, Alkaloids	Antidiabetic, anticancer, antipyretic, laxative antioxidant, anti inflammatory (Iftikhar <i>et</i> <i>al.</i> , 2022)
3.	Nigella sativa	Kalonji	Ranunculaceae	Seed	Fatty acid, terpene, alcohol, volatile oil	Antioxidant, inflammation, Asthma, hypertension, Cancer (Sharma <i>et al.</i> , 2009)
4.	Curcuma longa	Turmeric	Zinigiberaceae	Root and Rhizome	Flavonoids, curcumin, volatile oil, sugar, protein, Resin	Inflammation, Digestion, Antioxidant, Depression, Cancer (Omosa <i>et al.</i> , 2017)
5.	Ocimum sanctum	Holy Basil	Labiateae	Leaf	Flavonoids, tannin, saponin, phenolic, essential oil	Bronchitis, bronchial asthma, diarrhea, skin disease, Malaria (Pattanayak <i>et al.</i> , 2010)
6.	Salvia rosmainus	Rosemary	Lamiaceae	Aerial part	Phenolic acid, flavonoids, carnosic acid	Headache, dysmenhorrea, epilepsy, rheumatic pain, spasm (Choukairi <i>et al.</i> , 2019)
7.	Hyssopus officinalis	Hyssop/ Zofa	Lamiaceae	Aerial part	Beta pinene, limonene, caryphyllene	Asthma, sore throat, antioxidant, anorexia, Brochitis (Said <i>et al.</i> , 2015)
8.	Antrirrhinum majus	Snap dragon/ gul e meymoon	Scrophulariaceae	Flower	Amino acid, pigments, flavonoids, oil	Diuretic, sucrvy, liver disorder, tumor (Al sanafi, 2015)
9.	Salvia splendens	Scarlet sage	Lamiaceae	Leaves	Flavonoids, triterpenes, saponin, tannin, alkaloids, phenolic content	Wound dressing, dysentery, colic hemorrhides (Moharram <i>et al.</i> , 2012)
10	Tropaeolum majus	Nasturtium	Tropaeolaceae	Flower	Ascorbic acid, flaconoid, phenoiliv content	expectorant, anti cancer (Garzón and Wrolstad, 2009)
11	Dalbergia sisso	Sheesham	Fabaceae	Leaves	Carbohydrate, protein, amino acid, flavonoids, phenolic content	Skin disorder, peptic ulcer, Bleeding, Leprosy (Bhattacharya <i>et al.</i> , 2014)
12	Dalbergia sisso	Sheesham	Fabaceae	Flower	Carbohydrate, protein, amino acid, flavonoids, phenolic content	Skin disorder, peptic ulcer, Bleeding, Leprosy (Bhattacharya <i>et al.</i> , 2014)
13	Alo barbadensis	Alo vera	Liliaceae	Aerial parts	Anthraquinone, carbohydrate, saponin, steroids, tannin	Wound healing, Diabetes, constipation (Manvitha and Bidya, 2014)
14	Abelmoschus esculentus	Lady finger	Malvaceae	Aerial parts	Cellulose, lignin, Hemicellulose	Diabetes, weight loss, constipation (Jain <i>et al.,</i> 2012)
15	Abelmoschus esculentus	Lady finger	Malvaceae	Seed	Cellulose, lignin, Hemicellulose	Diabetes, weight loss, constipation (Jain <i>et al.,</i> 2012)
16	Cuscuta reflexa	Giant dodder	Cuscutaceae	Sap	Alkaloid, flavonoid, terpenoids, saponin, tannin, steroids	Purgative, cough, jaundice, muscle pain (Saini <i>et al.</i> , 2015)
17	Hibiscus schizopetalus	Shoe- flower	Malvaceae	Leaves	Terpenoids, steroids, Phenolics	Arthritis, diabetes, cough, malaria (Wong <i>et al.</i> , 2016)

Table 1: Detail Description of Medicinal plants selected for anti- Urease screening.



Fig. 1: Effect of extracts at 25ug/ml on urease inhibition activity. Data are shown as the mean  $\pm$  SD (n = 3). Asterisk represents statistically significant.



Fig. 2: Effect of extracts at 50ug/ml on urease inhibition activity. Data are shown as the mean  $\pm$  SD (n = 3). Asterisk represents statistically significant.

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Fig. 3: Effect of extracts at 75ug/ml on urease inhibition activity. Data are shown as the mean  $\pm$  SD (n = 3). Asterisk represents statistically significant.



Fig. 4: Representation of comparison of three different concentrations (25ug/ml, 50ug/ml and 75ug/ml)

S No.	Medicinal Plants (Extracts)	IC <sub>50</sub> (µg/ml)
1	Silybum marianum	54.16
2	Fagonia arabica	39
3	Nigella sativa	37.1
4	Curcuma longa	58
5	Ocimum sanctum	41
6	Salvia rosmainus	64
7	Hyssopus officinalis	42
8	Antrirrhinum majus	71.6
9	Salvia splendens	68
10	Tropaeolum majus	40.17
11	Dalbergia sisso (Leaves)	63
12	Dalbergia sisso (Flowers)	60
13	Aloe vera	36.5
14	Abelmoschus esculentus (Aerial parts)	41.2
15	Abelmoschus esculentus (seed)	40.4
16.	Cuscutare flexa	35.4
17.	Hibiscus schizopetalus	44.7
Standa	rd Thiourea	35.6

**Table 2**:  $IC_{50}$  values of medicinal plant extracts and standard Thiourea for Anti-urease activity.

### CONCLUSION

Gastric problems caused by *H. pylori* are prevalent among at least 50% of the population. It is evident from this research study that utilization of medicinal plants will provide highly effective, cheap and safe treatment to people suffering from such gastric diseases. Incorporation of food, vegetables and herbs having anti-urease potential in our daily routine will prevent disastrous effects of *H. pylori* on stomach health and upgrade the quality of life. However, to achieve optimum therapeutic benefits from these herbs it is mandatory to utilize best quality genuine herbs free from adulteration and all kinds of environmental contamination. Our research stipulate prospects for developing reliable and precise anti-*H. pylori* drugs from natural sources.

## REFERENCES

- Al-Snafi AE (2015). The pharmacological importance of Antirrhinum majus-A review. *Int. J. Pharm. Sci. Techno.*, **5**(4): 313-320.
- Amin M, Anwar F, Naz F, Mehmood T and Saari N (2013). Anti *Helicobacter pylori* and urease inhibition activities of some traditional medicinal plants. *Molecules*, 18: 2135-2149.
- Bhattacharya M, Singh A and Ramrakhyani C (2014). Dalbergia sissoo - An important medical plant. J. Med. Plant Studies, **2**(2): 76-82.
- Cellia JP, Bradley S Turnerb, Nezam H Afdhalb, Sarah Keatesb, Ionita Ghiranb and Ciaran P Kelly and Ewoldt RH McKinley GH, So P, Erramilli S and Bansil R

(2009). *Helicobacter pylori* moves through mucus by reducing mucin viscoelasticity. *PNAS*, **106**(34): 14321-14326.

- Chan FKL and Lau JYW (2021). Peptic ulcer disease. *In*: Feldman M, Friedman LS, Brandt LJ, eds. Sleisenger and Fordtran's Gastrointestinal and Liver Disease. 11<sup>th</sup> ed. Philadelphia, PA: Elsevier: Chap 53.
- Choukairi Z, Hazzaz T, Lkhider M, Ferrandez JM and Fechtali T (2019). Effect of *Salvia officinalis* L. and *Rosmarinus officinalis* L. leaves extracts on anxiety and neural activity. *Bioinformation*, **15**(3): 172-178.
- Cover TL and Blaser MJ (2020). *Helicobacter pylori* and other gastric *Helicobacter* species. *In*: Bennett JE, Dolin R, Blaser MJ, eds. Mandell, Douglas and Bennett's Principles and Practice of Infectious Diseases. 9<sup>th</sup> ed. Philadelphia, PA: Elsevier; Chap 217.
- Cunha ES, Chen X, Sanz-Gaitero M, Mills DJ and Luecke H (2023). Cryo-EM structure of *Helicobacter pylori* urease with an inhibitor in the active site at 2.0 Å resolution. *Nat. Commun.*, **14**(1): 648.
- Cunha ES, Chen X, Sanz-Gaitero M, Mills DJ and Luecke H (2021). Cryo-EM structure of *Helicobacter pylori* urease with an inhibitor in the active site at 2.0 Å resolution. *Nat. Commun.*, **12**: 230.
- Garzón GA and Wrolstad REJFC (2009). Major anthocyanins and antioxidant activity of Nasturtium flowers (*Tropaeolum majus*). *Food Chem.*, **114**(1): 44-49.
- Graham DY and Miftahussurur M (2018). *Helicobacter pylori* urease for diagnosis of *Helicobacter pylori* infection: A mini review. J. Adv Res., **13**: 51-57.
- Hanif M, Shoaib K, Saleem M, Rama HN, Zaib S and Iqbal J (2012). Synthesis, urease inhibition, antioxidant, antibacterial and molecular docking studies of 1,3,4-oxadiazole derivatives. *ISRN Pharmacology*, pp.1-9.
- Iftikhar N, Chatha SAS, Ahmad T, Ali Q, Hussain AI and Rathore HA (2022). *Fagonia arabica* L.: A review of its phytochemistry, pharmacology and traditional uses. *Comb. Chem. High Throughput Screen.*, **25**(7): 1187-1199.
- Jain N, Jain R, Jain V and Jain S (2012). A review on *Abelmoschus esculentus. Pharmacia*, 1(3): 84-89.
- Khan, BA, Akhtar, N, Khan H, Mustafa G, Niazi ZR and Menaa F (2017). Short communication - Urease inhibitory activity of *Hippophae rhamnoids* and *Cassia fistula*. *Pak. J. Pharm. Sci.*, **30**(5): 1779-1781.
- Khan H, Saeed M, Muhammad N, Gaffar R, Gul F and Raziq N (2015). Lipoxygenase and urease inhibition of the aerial parts of the *Polygonatum verticillatum*. *Toxico. Indus. Health*, **31**(8): 758-763.
- Manvitha K and Bidya B (2014). Phytochemistry. Aloe vera: A wonder plant its history, cultivation and medicinal uses. J. Pharmacogn. Phytochem., 2(5): 85-88.
- Moharram FA, Marzouk MS, El-Shenawy SM, Gaara AH and El Kady WM (2012). Polyphenolic profile and

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biological activity of *Salvia splendens* leaves. J. Pharm. Pharmacol., **64**(11): 1678-1687.

- Omosa L, Midiwo J and Kuete V (2017). *Curcuma longa*. *In*: Medicinal spices and vegetables from Africa. *Elsevier*, pp.425-435.
- Pattanayak P, Behera P, Das D and Panda SK (2010). Ocimum sanctum Linn. A reservoir plant for therapeutic applications: An overview. Pharmacog. Rev., 4(7): 95-105.
- Said-Al Ahl HA, Abbas ZK and Sabra AS (2015). Tkachenko KG. Essential oil composition of *Hyssopus* officinalis L. cultivated in Egypt. Int. J. Plant Res., 1(2): 49-53.
- Saini P, Mithal R and Menghani E (2015). A parasitic medicinal plant *Cuscuta reflexa*: An overview. *Int. J. Sci. Eng. Res.*, 6: 951-959.
- Sharaf M, Arif M, Hamouda HI, Khan S, Abdalla M and Shabana S, Hussein ER, Tehsin UK, Chi Z and Liu C (2022). Preparation, urease inhibition mechanisms, and anti-*Helicobacter pylori* activities of hesperetin-7rhamnoglucoside. *Curr. Res. Micro. Sci.*, **3**: 100103.
- Sharma NK, Ahirwar D, Jhade D and Gupta S (2009). Medicinal and phamacological potential of *Nigella* sativa: A review. *Ethnobotanical Leaflets*, 13: 946-955.
- Smoot DT (1997). How does *Helicobacter pylori* cause mucosal damage? Direct mechanisms. *Gastroenterology*, **113**(6): S31-S50.
- Suzuki H, Nishizawa T and Hibi T (2010). *Helicobacter pylori* eradication therapy. *Future Microbiol.*, **5**(4): 639-648.
- Wang X, Zhang Z and Wu SC (2020). Health benefits of Silybum marianum: Phytochemistry, pharmacology, and applications. J. Agri. Food Chem., 68(42): 11644-11664.
- Weatherfourn MW (1967). PhenoI-hypochlorite reaction for determination of ammonia. *Anal. Chem.*, **39**(8): 971–974.
- Wong S, Chan E and Chan HT (2016). A review on the phytochemistry and pharmacology of two lesser-known Hibiscus species: *H. taiwanensis* and *H. schizopetalus*. *Int. J. Pharmacog. Phytochem. Res.*, 8: 1341-1346.
- World Gastroenterology Organization. Global Guidelines Helicobacter pylori. pp.4-10.
- Yuri FR, Marcelo PQ, Tiago OB, Priscila GC, Vagner T and Fernando MJ (2018). Review on the development of urease inhibitors as antimicrobial agents against pathogenic bacteria. J. Adv. Res., 13: 69-100.