Original article

Single-step green synthesis of gold conjugated polyphenol nanoparticle using extracts of Saudi’s myrrh: Their characterization, molecular docking and essential biological applications

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A B S T R A C T

The progress in the innovative nanocrystal synthesis process by using environmentally benign and low-priced nontoxic chemicals, solvents, and renewable sources remains a challenging task for researchers worldwide. The majority of the existing synthesis techniques engage in the potentially dangerous, for either human health or the environment. Current investigation has been centered on green synthesis processes to create novel nanomaterials, which are eco-friendly as well as safer for sustainable marketable feasibility. The current work provides the green synthesis method for gold nanoparticle (GNPs) synthesis using Commiphora myrrh (C.myrrh) extract. This simple method includes 6 ml of HAuCl₄·3H₂O treated with 4 ml C.myrrh extract having pH 4.5 after 80 min at 25°C temperature. In this novel method, green synthesized GNPs characterized by UV–Vis, X-ray diffraction spectroscopy (XRD), zeta potential, fourier transform infrared (FT-IR), high-resolution transmission electron microscopy (HR-TEM), energy dispersive X-ray spectroscopy (EDXA), and dynamic light scattering (DLS). During the development successful antioxidant assay, the DPPH assay was applied. The cell toxicity of green synthesized GNPs was evaluated following an MTT assay against HCT-116 (colon cancer) and MCF-7 (breast cancer).

Besides molecular docking in the δ-elemene for inhibitor to VEGFR-2 domain revealed more negative docking score (−3.976) which is an excellent binding affinity to the C.myrrh@GNP. The synthesized GNPs showed anti-diabetic, antibiotic, and antibacterial properties and anti-inflammatory inhibition against COX-1, and COX-2 enzymes.

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1. Introduction

Nanotechnology is a scientific study of properties and the extent of response capacity in addition to the directing materials on the molecular and atomic scale ranging from 1 to 100 nm. Despite the small scale or size of these materials (Kim et al., 2016), they can help in solving some complicated humanity problems (Abdelbasir and Shalan, 2019; Li et al., 2022; Liao et al., 2021; Qiu et al., 2021; Yang et al., 2021; Yue et al., 2021). Currently, nanotechnology is renowned as a reputable progressive field having an extensive choice of applications including pharmaceutical and further industries. Metal nanoparticle production has gained growing concentration owing to its broad array of exercises in a variety of areas of medicine, electronics, and energy, (Doroudian et al., 2021; Kamat et al., 2021; Khalaf et al., 2021; Khalaf et al., 2021; Liang and Liang, 2021; Neha et al., 2021). Additionally, last decade, metal oxide nanoparticles also proved to be the subject of wide research, owing to their different application in many scientific areas (Aziz et al., 2015; Gan et al., 2019; Lee et al., 2020; Liu et al., 2019; Xue et al., 2017; Yang et al., 2018; Zhou et al., 2019), metal oxide (Faisal et al., 2021; Dizaj et al., 2014; Falcaro et al., 2016; Fernandez-Garcia and Rodriguez, 2007; Stoimenov et al., 2002; Franke et al., 2006). Nano-sized materials have been broadly used mainly considering their countless applications in different fields including chemical, physical
and biological (Al Jahdaly et al., 2021; Ranke et al., 2006; Nikam et al., 2018; Richter et al., 2010; De et al., 2008; Llop et al., 2014). Liquid metal nanoparticles are created from noble metals like Pt (Al-Radadi, 2019; Al-Radadi and Adam, 2020), Au (Dreaden et al., 2012; Sperling et al., 2008), and Ag (Al-Radadi and Abu-Dief, 2022; Abdullah et al., 2021; Al-Radadi, 2018; Al-Radadi and Al-Youbi, 2018a; Kaviya et al., 2011). Nanoparticles (denoted to NPs) showed excellent potential seeing that antimicrobial (Al-Radadi, 2022a; Behzad et al., 2021; Siege1 et al., 2013), antioxidant (Desai et al., 2020; Zhang et al., 2018), anti-inflammatory agents (Sengupta et al., 2018), antidiabetic (Nirmala Grace and Pandian, 2007), anti-Alzheimer (Al-Radadi, 2022b; Al-Radadi et al., 2022) and anti-cancer (Al-Radadi, 2022c; Jain et al., 2007) having a strong capability of retarding the growth of unnecessary and injurious bacteria causing threatening disease. Gold nanoparticles (further termed as GNPs) have attracted lots of interest among noble metals due to their outstanding physicochemical and optical features, which can be broadly exploited in severe domain names of human activity (Al-Radadi, 2021a; Al-Radadi and Al-Youbi, 2018b). GNPs can in addition exhibit antimicrobial (Zhang et al., 2015), antioxidant potential (Al-Radadi, 2021b), and catalysis (Aswathy Aromal and Philip, 2012; Ramakrishna et al., 2016). GNPs are highly sensitive (Chen et al., 2008), stable (Balasubramanian et al., 2010), highly reliable, and less toxic (Lasagna-Reeves et al., 2010) than additional metal NPs.

However, a crucial asset of GNPs is their biocompatibility with living creatures building them broadly utilized within biomedical applications, drug delivery, sensor and diagnosis, and bio-imaging. (Kumar et al., 2011; Malathi et al., 2013). GNPs derived from natural sources have shown to be effective against a variety of uncontrolled growing cells i.e. cancer cell types (Khandalou et al., 2018; Parida et al., 2014; Singh et al., 2019). As a result of widespread toxic concerns connected with chemical and physical processes, the eco-friendly ‘green’ synthesis of nano-sized materials has been a crucial and popular topic in support of material research in recent years (Albrecht et al., 2006; Khan, 2020). Plants are favored over some other organic procedures due to the fact they take away the requirement of retaining cell culture (David et al., 2014; Eccles, 1999).

The plant-intervened engineered approach (Al-Radadi, 2022c), is a significant procedure that can be helpfully made and engineered (Rajeshkumar and Bharath, 2017), the utilization of extracellular extract was viewed as more powerful and proficient in controlling the size, shape, and dispersity of the NPs (David et al., 2014; Narayanan and Sakthivel, 2011; Udayasoorian et al., 2011), therefore it is observed a typical plant referred to in old stories medication as “myrrh” is one of the most widely recognized spices in Saudi Arabia. Commiphora myrrh (C. myrrh) has a place with the class Commiphora and the family Burseraceae (Sheir et al., 2001; Su et al., 2011). C. molmol Engler (Burseraceae) is local to Madagascar, the Arabian islands, and India (Ben-Yehoshua and Borowitz, 2012). It is one of the best homegrown medications on the planet for sore throats, ulcers, and gum disease (Khalil et al., 2020). C. molmol Engler is one of the types of myrrh which has been utilized in the treatment of twisted and as a fast-acting medicine (Alwhibi et al., 2020), likewise it is a powerful antimicrobial specialist as it is valuable for the treatment of skin breakout, bubbles, and joint pain. It shows solid antithrombotic activity (Bhattacharjee and Alenezi, 2020). Myrrh extract has been utilized as reducing (Elia et al., 2014) self-assembly (Zhao et al., 2018) and covering specialists (Atta et al., 2014) for the union of NPs which could be the benefit over microbial combination as there is no need of the explained course of refined and keeping up with the phones (Wu et al., 2010). Myrrh has neighborhood energizer and insect mending, and disinfector properties for wounds and scraped spots. It is utilized as a mouth wash (Eid et al., 2021), and as a uterine energizer and emmenagogue, it is utilized in the treatment of diseases in the mouth such as mouth ulcers, pyorrhea as well as catarrhal issues of pharyngitis and sinusitis. It is incredible in irritated mouth and outrageous ulceration of inconsistent ptyalism. The concentration of myrrh (gum) diminishes the outright augmentation of blood glucose over the fasting focus consistently of the oral glucose resistance test in both ordinary and diabetic rats (Al-Awadi and Gumaa, 1987), and may end up being a valuable helpful specialist in the treatment of non-insulin subordinate diabetes mellitus. A color of myrrh is utilized for the treatment of aphthous ulcers (stomatitis aphthosa). It shows solid antithrombotic movement. Extract of myrrh is utilized as stomach-related help medication, and it is endorsed by the FDA for the utilization in food and oral medical care drug items. It was given GRAS status as a flavor fixing by FEMA (Satpathy et al., 2020). Myrrh is utilized in conventional Chinese medicina to diminish torment and enlarging because of awful injury (Lee and Lam, 1993). It is utilized as a hypolipidemic specialist. Myrrh is helpful in constant gastritis and nuclear dyspepsia with full pale tongue and layer, as well as successive mucous stouts joined by a fart. It helps in the treatment of laryngitis and respiratory whines. As of late the cytotoxic and antitumor movement of myrrh has ended up being identical to those of the standard cytotoxic medication cyclophosphamide. It likewise offered insurance against mucosal harm brought about by indomethacin (Al-Harbi et al., 1997). Commiphora molmol showed mitigating action, it showed an undeniable hindrance of ADP, adrenaline, and serotonin-instigated platelet collection. It additionally showed a solid thyroid stimulatory activity when directed to pale-skinned person rodents (Wijesekera, 2016). The constituents of myrrh, incorporate unpredictable oil (2–8%), tar (23–40%), gum (40–60%), and harsh standards (10–25%) (Chen et al., 2013). In Fig. 1 compound of myrrh incorporate phenols, furan sesquiterpenes, β-sitosterol, and liquor dissolvable tars with powerful sterile, cancer prevention agent and calming exercises which can be utilized actually in the mending of different kinds of wounds and ulcers (Cao et al., 2019; Negahdari et al., 2017; Walsh et al., 2010). The antimicrobial activity of phenolic compounds is portrayed by restricting nucleophilic amino acids in bacterial proteins, which prompts inactivation of the protein and loss of its capacity, in this way avoidance of the bacterial life cycle (Wink and Mitchell, 1998). Besides, phenolic intensifies focus on a superfluous level of uncovered adhesins in the microbial cell, which will forestall bacterial adherence with different cells and arrangement of biofilms. In this concentration, it is zeroed in on the amalgamation of GNPs to raise the effectiveness of myrrh, and it is involved in the treatment of wound sterilization (Haffor, 2010), and come to an incredible outcome. In this methodology, the regular non-poisonous fixing removed from the myrrh plant was utilized, rather than normal unsafe synthetic compounds, in a blend with UV light illumination to synthesize GNPs. The Cmyrrh@GNPs are homogenous, round, and three-sided shape, accomplish 1.68–10 nm particular sizes, and are all around scattered without collection, the portrayal of combined GNPs under various circumstances was dissected utilizing UV-apparent retention spectra, high-resolution transmission electron microscope (HR-TEM) examination, transmission electron microscope (TEM), fourier-transform infrared spectroscopy (FT-IR), powder XRD, dynamic light dissipating (DLS), energy dispersive X-ray spectroscopy (EDS), X-beam photoelectron spectroscopy (XPS) analysis and zeta potential.
This study investigated the property of *C.myrrh* extract in the development of GNPs and cell reinforcement, cytotoxicity, anti-cancer, mitigating, hostile to diabetic, and anti-microbial specialists. These amino acids and nutrients are likewise present in *C.myrrh* separate in the Kingdom of Saudi Arabia (KSA) locale. The current article investigates the natural potential of the antioxidants, amino acids and some secondary metabolites of a particular *C.myrrh* plant for the biosynthesis of GNPs. The biosynthesized *C.myrrh*@GNP applied as a multi pharmaceutical agent including anti-diabetic, antibiotic, anti-oxidant, anti-bacterial, anti-inflammatory and Lastly anti-tumorigenic towards two malignant growth cell lines, colon carcinoma (HCT-116) and breast adenocarcinoma (MCF-7).

2. Materials and methods

2.1. *C.myrrh* extract preparation

The oleo-gum-resin of *C.myrrh* was bought from a local market in Madinah Munawara, KSA, and washed with double distilled water (ddH2O) followed by ethanol. The oleo-gum-resin of *C.myrrh* was grounded into coarse powder by a grinder. The oleo-gum-resin of *C.myrrh* powder (500 mg) was extracted using ethanol or ddH2O by maceration for 24 h (h) at room temperature (RT) with occasional shaking (250 g × 2L). The extract was then filtered and dried by rotatory evaporator. The oleo-gum-resin of *C.myrrh* extract was utilized for further experiments.
2.2. Synthesis of GNPs

(HAuCl₄·3 H₂O) derived from Sigma Aldrich with high purity (9.99%), C.myrrh stock solution prepared and mixed with 4 mL volume of (1 × 10⁻³) M of HAuCl₄·3 H₂O. The reduction reaction of Au³⁺ to Au⁰ was confirmed when the yellow color changed to ruby-red.

2.2.1. Optimization for C. myrrh@GNPs synthesis

A few elements influence the combination of GNPs, for example, the amount of myrrh extract, the amount of (HAuCl₄·3 H₂O), response time, pH, and the temperature was contemplated to gauge the improvement boundaries for the union of C.myrrh@GNPs from an extract of myrrh.

2.2.2. Optimization for C. Myrrh extract

Various volumes of C. myrrh extract from 1 to 6 mL were treated with 4 mL of 1 mM HAUCl₄·3 H₂O independently and the temperature was retained at RT intended for around 80 min for the production of C.myrrh@GNPs.

2.2.3. Optimization for the volume of HAU Cl₄·3 H₂ O

The impact of volume of HAU Cl₄·3 H₂ O to 6 mL of Cmyrrh extract was examined by treating 1 to 4 mL volumes of HAU Cl₄·3 H₂ O independently for about 80 min at RT.

2.2.4. Optimization for the reduction of time

The reaction response time impact was examined by checking the UV–Vis spectra of C.myrrh@GNPs synthesized in the solution having maintained reaction time from 20 to 120 min, and different volumes of C.myrrh extract and gold salt.

2.2.5. Optimization of pH

A pH optimization was studied at various levels from 1 to 9 after 80 min of reaction time with optimized C.myrrh extract and salt concentration at RT. The optimization study of pH was done by 0.1 M HCl and 0.1 M NaOH.

2.2.6. Optimization for reaction temperature

The reaction temperature was examined by setting different reaction temperatures from 15 °C to 35 °C for around 80 min with the above-streamlined conditions to acquire C.myrrh@GNPs.

2.3. Characterization of GNPs

To guarantee the physio-substance nature of incorporated C.myrrh@GNPs, an assortment of insightful methods were utilized for the investigation. UV–Vis absorbance spectra were done with quartz cuvettes utilizing Cary 100, UV–Vis spectrophotometer within the frequency range 200–800 nm wavelength. FTIR was examined by 0.1 M HCl and 0.1 M NaOH. C.myrrh after 80 min of reaction time with optimized volume of (1/2) (9.99%), from an extract of myrrh.

2.3.1. Characterization of C. myrrh extract

The impact of volume of HAU Cl₄·3 H₂ O to 6 mL of Cmyrrh extract was examined by treating 1 to 4 mL volumes of HAU Cl₄·3 H₂ O independently for about 80 min at RT.

2.3.2. Characterization of the size of tiny particle that exists within a solution.

is a significant method moderately used to understand the distribution profile of C.myrrh@GNPs utilizing a Physical Electronics PHI 5000 Versa test II instrument was utilized.

2.4. Antioxidant study

Antioxidant study of HAU Cl₄·3 H₂ O, C.myrrh and C.myrrh@GNPs entirely set in stone by (Al-Radadi, 2021b; Hosseini-Mehr et al., 2011), where 50 μL of various convergences of HAU Cl₄·3 H₂ O, C.myrrh and C.myrrh@GNPs were occupied in various test tubes, then, 2.5 mL of 0.1 mM DPPH methanolic arrangement was mixed with these test tubes and thoroughly shaken. The test tubes were permitted to remain at 25 °C for 20 min. The separate control test tube was made ready as above utilizing methanol dissolvable rather than HAU Cl₄·3 H₂ O, C.myrrh and C.myrrh@GNPs. The DPPH free radical scavenging test was completed for the assessment of the antioxidant test. This examines the free radical scavenging limit of the researched extricate. DPPH is a particle including a steady free extremist. Within the sight of a cell reinforcement, which can give an electron to DPPH, the purple tone commonplace for nothing DPPH radical rots, and the alteration in absorbance is estimated at λ = 517 nm. The anti-radical action of the GNPs, C.myrrh, and C.myrrh@GNPs was estimated as the free extremist rummaging action DPPH. Shortly, within 3 mL of every GNPs, C.myrrh, and C.myrrh@GNPs 1 ml of methanol arrangement of 0.1 mM/L DPPH solution was mixed. The combination was set aside in dark RT intended for 30 min and the absorbance was estimated at λ = 517 nm against a blank. The percentage activity of free radical scavenging was determined utilizing the following formula:

% potential of DPPH free radical scavenging = (control-test)/control × 100.

The IC₅₀ upsets of the HAU Cl₄·3 H₂ O, C.myrrh and C.myrrh@GNPs, for example, the convergence of HAU Cl₄·3 H₂ O, C.myrrh separate and C.myrrh@GNPs. Important to diminish the underlying union of DPPH by half were determined. The IC₅₀ esteem μg/mL is the viable focus at which DPPH radical was looking through by half and the worth was acquired by addition from the straight declined examination.

2.5. Examination of cell viability against of C.myrrh extract and C. myrrh@GNPs

For cell viability study, Human umbilical vein endothelial cells (HUVECs) were cultivated in M199, enhanced with heat-inactivated 20% FBS (WELGEN Inc.), 20 ng/ml of bFGF, 100 units/ml of penicillin, and 100 μg/l of streptomycin within 5% CO₂ incubator at 37 °C.

The impact of HAU Cl₄·3 H₂ O, C.myrrh, and C.myrrh@GNPs on the suitability of HUVECs were resolved to utilize the MITT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide) test, which depends on the change of MTT to insoluble MTT-formazan by cleavage of the tetrazolium ring by mitochondrial dehydrogenase compounds in living cells (Al-Radadi, 2022a; Arulmozhi et al., 2013). In brief, HUVECs were developed in M199 with 20% FBS at a thickness of 2 × 104 cells on 24-well culture plates. Following one night, the media was supplanted by M199 including 1% FBS, and HAU Cl₄·3 H₂ O, C.myrrh, and C.myrrh@GNPs and the cells were then brooded for 24 h at 37 °C under a humidified air that included 5% CO₂. The cells were allowed to react with different concentrations of HAU Cl₄·3 H₂ O, C.myrrh, and C.myrrh@GNPs. Then, 5 ng/mL in H₂O of MTT solution was added to each well, trailed by the addition of 0.3 ml of dimethyl sulfoxide to break down the MTT-formazan. MTT formazan was estimated by recording the optical density at 570 nm wavelength as indicated by the
following equation. The experiment was performed in triplicate and repeated thrice.

\[
\text{Percentage of cell viability} (\%) = \left( \frac{\text{Sample Absorbance}}{\text{Control absorbance}} \right) \times 100
\]

2.6. Cytotoxicity activity

MTT compound was used to assess cell toxicity (Al-Radadi, 2022c, 2021a; Yas et al., 2021). Cell lines of the colon (HCT-116) and breast carcinoma (MCF-7) were grown in Eagle’s minimum essential medium with 2 mM L-glutamine concentration and Earle’s salts. In a nutshell, the cell lines were allowed to grow at \(1 \times 10^4\) cells/well in a 96 well plate, then incubated at 37 °C in a 5 percent CO\(_2\) incubator for 24 h. \(C\).\text{myrrh} and \(C\).\text{myrrh}@GNPs were used in concentrations of 15 g, 25 g, 50 g, and 100 g and incubated for 24 h. Following a 4 h incubation period, MTT (5 mg/ml in PBS) was added to each well. After discarding the supernatant, 100 µl of DMSO was mixed in each well of the plate, which was then smoothly shaken for dissolving the formazan formed and recorded the absorbance utilizing a plate reader at 530 nm. The inhibition of cell growth in percentage was measured utilizing the following formula and the concentration required for inhibiting 50% of the cell growth was also assessed for \(C\).\text{myrrh} and \(C\).\text{myrrh}@GNPs. DMSO was selected as negative control while doxorubicin was added as a standard drug and acted as a positive control.

The IC\(_{50}\) standards were estimated by mean values of three experiments with four concentrations.

\[
\% \text{ Inhibition} = 100 - \left( \frac{\text{OD of Sample}}{\text{OD of Control}} \right) \times 100
\]

2.7. Anti-inflammatory assay

Cyclooxygenase (COX) is an enzyme that produces prostaglandins that cause fever, pain, and inflammation. Herein, the biosynthetic GNP was examined for inhibition potential to COX-1 and COX-2 enzymes to decrease inflammation. COX-1 and COX-2 from Ovine Kit 701050, France were used to measure the anti-inflammatory capacity of \(C\).\text{myrrh}@GNP. For positive control ibuprofen (10 mM) was utilized. NPs of various concentrations, including 25, 50, 100, 200, and 400 µg/ml were utilized for test inhibition and calculated according to the manufacturer’s instructions. 96-well microplates were scanned at 530 nm wavelength absorbance for the presence of N, N, N', N' tetramethyl-phenylenediamine.

2.8. Antidiabetic assay

An \(\alpha\)-amylase and \(\alpha\)-glucosidase assays were performed to evaluate the anti-diabetic potential.

2.8.1. \(\alpha\)-amylase

The capacity of concentrates to repress \(\alpha\)-amylase was examined utilizing a chromogenic technique recently depicted by (Sigma Aldrich, USA). In 0.1 M phosphate buffer having pH 6.8, the 1 m/ml protein was ready and completely blended in with 5 mM, 4-nitrophenyl-\(\alpha\)-D-maltopentaoside solution. A little part of the examination sample was acquainted with the reaction mixture and incubated for 30 min at 37 °C. The percent inhibition activity of the sample was determined by subtracting the absorbance of two samples when the reaction mixture include extract and while it was excluded by utilizing a microplate reader at wavelength 530 nm.

2.8.2. \(\alpha\)-glucosidase

To further evaluate the anti-diabetic potential, the \(\alpha\)-glycosidase inhibition test was carried out using a color-producing method having a 0.45 m polyethylene filter end-capped column. A little portion of the experimental sample was reacted with 5 mM of 4-nitrophenyl-\(D\)-glucopyranoside (4NP; Sigma, country name?) 1 ml of intestinal fluid and allowed to incubate at 37 °C for 30 min. The potential activity was evaluated using percent inhibition potential via calculating a diversity between absorbance standards while containing the extracts and the absorbance of the examined sample on a 530 nm wavelength.

Fig. 2. Schematic diagram for the reduction process and stabilization of GNPs by \(C\).\text{myrrh} aquas extract, the work plane for synthesis of \(C\).\text{myrrh}@GNPs.
2.9. Bioassay of antibacterial activities

The antibacterial effectiveness of every extract was tested utilizing a total of four bacterial strains, two of which were *Staphylococcus aureus* and *Bacillus subtilis* Gram-positive and two of which were *Escherichia coli* and *Pseudomonas aeruginosa*, Gram-negative that contaminate food. The strains of bacteria were derived from King Fahad Medical Research Centre (KFMRCC) at King Abdulaziz University, Jeddah, Saudi Arabia.

2.9.1. Agar well diffusion method

The anti-bacterial potential of C.myrrh@GNPs was analyzed using the conventional agar well diffusion technique (Kunchandy and Rao, 1990) utilizing a 5 mm borer about six wells were created in an aseptic environment. The bacterial cultures were poured throughout the Petri plates by spreader and various C.myrrh extract concentrations and C.myrrh@GNPs were added to 15 μg/ml, 25 μg/ml, 50 μg/ml, and 100 μg/ml, into the 5 mm well. Streptomycin acted as a positive control and lacking samples were retained as a negative control. The Petri plates were kept at 37 °C for incubation for 24 h. The inhibition area was calculated by a MIC ruler along with mentioned in mm.

2.10. Molecular docking

2.10.1. Preparation of ligand

The main compounds polyphenols in C.myrrh planned to be used for docking were prepared using the default protocol of the Ligprep program (2021) in the Schrödinger’s suite. All compounds were docked to the target protein using the glide dock XP protocol without using perform post-docking minimization. XP G-score (Kcal mol⁻¹) was used as ranking criteria for the best-docked ligands.

2.10.1.1. Preparation of protein. VEGFR-2 was found to be crucial for cell survival, which regulates endothelial differentiation in both the breast cancer cells (MCF-7) and human colorectal carcinoma (HCT-116) (El-Adl et al., 2021). Thus each experiment used VEGFR-2 downloaded from the Protein Data Bank (PDB ID: 1YWN) (Miyazaki et al., 2005). Furthermore, the three-dimensional structures of ovine COX-1 (PDB ID: 3KK6) (Rimon et al., 2010) and mouse COX-2 (PDB ID: 3LN1) (Wang et al., 2010) were selected for anti-inflammatory docking (Karim et al., 2019). Finally, for the antidiabetic docking (Missioui et al., 2021), the crystal structure of α-amylase (PDB ID: 4GQR) (Williams et al., 2012) and α-glucosidase (PDB ID: 5NN5) (Roig-Zamboni et al., 2017) were selected.

All three-dimensional complex structures were downloaded from the PDB database (https://www.rcsb.org/). The protein structures were prepared using the protein preparation wizard program from the Schrödinger suite (Schrödinger Suite, 2021-2) in which water molecules (>5Å radius) and small molecules present were removed from the structure part, disulfide bonds were created and hydrogens were added to the PDB structures. Restrained impreff minimization with default settings was performed on the structure with optimized potentials for liquid simulations (OPLS-2005) force field. The resulting structures were used for receptor grid generation for docking.

3. Result

3.1. Analytical study of C.myrrh

GNPs were synthesized in a green, environmental technique with an eco-friendly approach without using any organic solvent. Several devices were used to characterize GNPs synthesized with myrrh extract, under several conditions of reaction, the production of GNPs was performed within the different environments including at various temperatures, pH, varying time intervals varying volumes of C.myrrh extract, and gold salt. From experience, it is found that the best result to get the best images taken by electron microscopy of GNPs were observed to be small, spherical, homogeneous.
and with no aggregation, when *C.myrrh* extract (6 ml) was added to 1 mM HAuCl₃·3H₂O (4 ml) and after 80 min at 25 °C and pH 4.5. The solution color transformed to ruby-red from yellow. Studying the UV–VIS spectroscopy of *C.myrrh*@GNPs showed various absorbance peaks, it is found that the surface plasma resonance (SPR) for GNPs strong, and the best peak for the highest absorption λₑант = 532 nm. It is noticed that in Fig. 3 small amount of volume, the absorption is too low and broad, but when the increase the volume, the absorption increases as it becomes sharper and narrower, the best volume was (6 ml) absorbed at 1.4 and λₑант = 532 nm in Fig. 4. It is noticed that when the volume of HAuCl₃·3H₂O increased the absorption increased as well, the best curve observed for 4 ml at wavelength 532 nm as shown in Fig. 4. It can be seen that (4 ml) of (HAuCl₃·3H₂O) is narrower and sharper than (3 ml) which is broad which indicates the formation of GNPs. In Fig. 5, the time effect on *C.myrrh*@GNPs can be seen with the best absorption at 80 min, when GNPs form at a wavelength λₑант = 532 nm and absorption more than 2. Fig. 6, strength variations of UV–Vis peak of GNP as a purpose of (4 ml) (HAuCl₃·3H₂O) and (6 ml) of *C.myrrh* extract following 80 min after mixture at 25 °C. The absorbance spectra present at 532 nm corresponds to Au⁰ that showed continuous increase till 80 min. and stabilized at a 532 nm peak that is equivalent to the synthesis of Au⁰ nanostructures. There was a difference in UV Vis absorbance spectra and time is taken to achieve stabilized λₑант demonstrated by absorbance to time spectra. These findings recommended that the first 80 min is sufficient to accomplish the uppermost concentration of stabilized *C.myrrh*@GNPs. Fig. 7 showed the best effect of temperature on GNPs at 25 °C that have absorption at 1.9 and wavelength λₑант = 532 nm, which was the narrower and sharper at pH 4.5. It is noticed that the best absorption with wavelength λₑант = 532 nm then the absorption decreases every time increased the pH which is shown in Fig. 8.

3.2. TEM analysis

With the aim of the morphological and size-dependent study of the *C.myrrh*@GNPs electron microscopy was performed in transmission mode. The size of *C.myrrh*@GNPs found in TEM images ranges from 1.68 – 7.86 nm. The TEM images of *C.myrrh*@GNP in Fig. 9-B presented nano size with spherical shape and did not exhibit aggregation. Further approach devoted to morphological study shown in TEM images (Fig. 9-A) given a clear idea of the size of *C.myrrh*@GNPs range from 1.68 nm, 7.86 nm, and 10.8 nm at 20 nm resolution. The HAADF-STEM images of the *C.myrrh*@GNPs in Fig. 9-C demonstrate the gold in bright ring structures. Additionally, the intensity of 4 nm diameter-sized nanoparticles was confirmed in the Fig. 9-D.

3.3. EDX study

Further characterization of the elemental gold was performed by the EDX technique. The elemental analysis profile of *C.myrrh*@GNPs exposed a sturdy signal for gold at 2.8 KeV alongside carbon and oxygen peaks. The consequential other signals may arise due to the secondary metabolite capped around the surface of the *C.myrrh*@GNPs (Fig. 10).
Fig. 5. Intensity variant within UV–Vis. spectra of C.myrrh@GNPs as function of time at room temperature 25 °C with (6 ml) of C.myrrh extract and (4 ml) (HAuCl₄·3H₂O) (1 × 10⁻³) M.

Fig. 6. Absorption Intensity variant within UV–Vis spectra of C.myrrh@GNPs as a function of (4 ml) (HAuCl₄·3H₂O) (110⁻³) M stock solution and (6 ml) of C.myrrh extract after 80 min of addition at 25 °C.
Fig. 7. UV–vis spectra for C.myrrh@GNPs at different temperatures (15–35), after 80 min with (6 ml) of C. myrrh extract and (4 ml) (HAuCl₄·3H₂O) (1x10⁻³) M.

Fig. 8. UV–vis spectra for C.myrrh@GNPs at different pH values (1–9) at room temperature 25 °C, after 80 min with (6 ml) of C.myrrh extract and (4 ml) (HAuCl₄·3H₂O) (1x10⁻³) M.
Fig. 9. (A) HR-TEM image of spherical C.myrrh@GNPs, (B) TEM image of C.myrrh@GNPs, (C) HAADF-STEM images of the C.myrrh@GNPs and the elemental mapping images, (D) Size distributions graph of C.myrrh@GNPs.

Fig. 10. EDX analysis for C.myrrh@GNPs.
3.4. DLS analysis

The analysis of the zeta potential study found (-10.8) mV that indicated the stability of C.myrrh@GNPs at 25°C (Fig. 11). It was in collaboration with the earlier images shown in TEM mean triplicate (Fig. 12).

3.5. FTIR analysis

FTIR analysis was completed to distinguish the bound bioparticles responsible for covering and producing GNPs utilizing C.myrrh extract (Fig. 13). The FTIR range of GNPs showed an exceptionally solid top at 3400 cm\(^{-1}\) was allocated as (–OH) bunches extending, the 3010 cm\(^{-1}\) peak acquired because of aromatic (C-H) bonds, where at 2900 cm\(^{-1}\) the top for aliphatic (C-H) gatherings, at 1700 cm\(^{-1}\), wave-numbers tops happened because of the presence of [C = O] (carbonyl groups) and 1030 cm\(^{-1}\) (C-O) extending vibrations of alcohols, carboxylic acids, ester, or ethers of bioparticles present in the C.myrrh.

3.6. XRD study

XRD is an effective technique for analyzing crystalline materials, where it gives information about crystal structure and crystal size for Nanomaterial. X-ray diffraction of C.myrrh@GNPs showed 5 peaks at the range which is attributed to the sides of the cube are centered on the side of the crystal that applies to the Joint Committee for Powder Diffraction Set (JCPDS) with the known information of X-Ray diffraction for the gold metal. The crystalline character of C.myrrh@GNPs was examined using the X-Ray diffraction method and found four major peaks at 38.1, 44.5, 65, 78, and 82 coupled theta, allocated to (111), (200), (220), (311) and (222) crystalline planes respectively of gold. These found indices indicate the FCC structure of GNPs. In Fig. 14 the additional peak shown at 82 coupled theta may arise due to the extract's amorphous nature. The average crystallite size is determined by using the Debye–Sheerrer formula: 

\[
D = \frac{K \lambda}{b \cos \theta}
\]

Where D represents the average thickness of crystalline grains, K represents the Scherrer constant (0.89), \(b\) represents the FWHM (Full-Width Half Maximum), \(\theta\) represents the angle of diffraction, and \(\lambda\) represents the X-ray wavelength.

3.7. XPS analysis

To additionally affirm the existence of polyphenol covering the surface of the NPs, an XPS investigation was carried out. Fig. 15-A, demonstrate the settled spectra of the center degrees of Au4f1/2 and Au4f3/2 at 84 eV and 88 eV, separately, relating admirably with the earlier reports of metallic Au. Fig. 15-B-C-D addresses the acquired spectra for N1s, C1s, and O1s, individually. The C 1s top at 284.5 eV and O1s top at 532.3 eV N1s top at 399.5 eV may be because of the extract of C.myrrh.

3.8. Antioxidant activity of GNPs by DPPH

DPPH (1,1-diphenyl -2-picryl-hydrazine) is a well-acknowledged approach to determining antioxidant characteristics as a result of its capacity in reducing the free radicals. DPPH radicals can
**Fig. 13.** FTIR spectrum of *C. myrrh* extracts and *C. myrrh*@GNPs.

**Fig. 14.** XRD spectrum of synthesized GNFs with *C. myrrh* extract.
immediately react to antioxidants. When antioxidants attract this radical, DPPH is revitalized from an antioxidant with the aid of using receiving a hydrogen atom. The free radicals of this compound produce a purple color and might absorb on the wavelength of 517 nm. Its absorption depth is reduced with the aid of using revitalization and it adjustments to yellow. The antioxidant energy relies upon the discount percent of preliminary darkish red color to the yellow color. The extra is the wide variety of hydroxyl corporations of antioxidant phenyl loop; the better is the wide variety of hydrogen atoms for response with DPPH and its stabilization. DPPH free radical scavenging impact C.myrrh and C.myrrh@GNPs in numerous concentrations like 10, 20, 30, 40, 50, 60, 70, 80, 90, 100 µg/ml. The observed IC<sub>50</sub> of C.myrrh, DPPH, and C.myrrh@GNPs were 100, 93, and 84 respectively (Fig. 16).

3.9. Cell viability against of C.myrrh and C.myrrh@GNPs

The selected cell lines (HUVEC) were allowed to react with different concentrations of HAuCl<sub>4</sub>·3H<sub>2</sub>O, C.myrrh, and C.myrrh@GNPs in an MTT test used for 48 h regarding the cytotoxic effect on HUVEC cells. The UV–Vis optical density absorbance intensity was identified at 570 nm wavelength, which specifies unexpected absorbance for HUVEC cell line till equal to 1 mg/ml for HAuCl<sub>4</sub>·3H<sub>2</sub>O, C.myrrh, and C.myrrh@GNPs as presented in Fig. 17. On the
basis of data shown here in figure it clearly indicates that when cell concentration increases, percent viable cell count decreases.

3.10. Cell toxicity study

The C.myrrh@GNPs were estimated for their anticancer property towards HCT-116 and MCF-7, by applying an MTT assay. Table 1 disclose that C.myrrh@GNPs were efficient towards HCT-116 and MCF-7 cell lines. Different concentrations of C.myrrh@GNP like 15 μg, 25 μg, 50 μg, and 100 μg were prepared and treated with the above-mentioned cell lines in the MTT assay. Cell toxicity study was executed to explore the cytotoxic activities of C.myrrh and C.myrrh@GNPs against cancer cell lines HCT-116 (colon cancer) and MCF-7 (breast cancer). Amazingly, in this study C.myrrh did not show considerable cell-toxicity towards selected cancer cell lines (HCT-116 and MCF-7). Cell toxicity against HCT-116 cell line in C.myrrh@GNPs was 59.3% for 15 μg/mL, 70.41%, for 25 μg/mL, 85.1 % for 50 μg/mL and highest percent inhibition of 96.5 for 100 μg/mL concentration (Fig. 18(A)). Particularly the cell toxicity against MCF-7 in C.myrrh@GNPs was observed highest in 100 μg/mL concentration i.e. 79.92%, 67.43% for 50 μg/mL, 55.20% for 25 μg/mL, 42.37% 15 μg/mL concentrations. Whereas the standard showed 98.18 percent activity as shown in (Fig. 18(B)). The results of the cell viability and toxicity studies indicated that when concentration increases cell viability and cell toxicity increased. The findings clearly showed the C.myrrh@GNPs presents more effective proliferation inhibition percentwise than C.myrrh extract shown in Fig. 19 for both the cell lines.

3.11. Anti-inflammatory assay

In this study biosynthesized GNP were used to inhibit COX-1, and COX-2 by making the different concentrations of C.myrrh@GNPs like 25 μg/ml, 50 μg/ml, 100 μg/ml, 200 μg/ml, and 400 μg/ml. The results show that as the C.myrrh@GNPs concentration increased the percent inhibition also increased accordingly (Fig. 20).

3.12. Antidiabetic

Diabetes mellitus is a metabolic problem caused by abnormalities in insulin release, insulin activity, or both. Therefore, insulin deficiency causes chronic hyperglycemia that adversely affects the digestion of starch, fat, and protein. It is one of the chronic non-communicable diseases (CNCD) that has appeared as a predominant disease in the world. The basis is the investigation of new sources of regular mixtures with potential anti-diabetic effects from tropical greens and nanodevices. An α-amylase is a compound that can break down starch into monomers of glucose. These factors, which are primarily involved in the inhibition of the well-known enzymes α-amylase and α-glucosidase, are of huge significance to diabetes research.

In this study, C.myrrh@GNPs were used to inhibit at concentration dependant way to α-amylase and α-glucosidase enzymes. The antidiabetic property of C.myrrh@GNPs was found to be increased when a high concentration (400 μg/ml) was used in a dose-dependent manner. The percent inhibition was observed at 50 ± 0.58 and 44 ± 0.49 for α-amylase and α-glucosidase enzymes respectively (Fig. 21).

3.13. Bioassay of antibacterial activities

Medicinal plants contain composite phytochemical components including proteins, saponins, terpenes, alcohols, alkaloids, and phenols, whereas microbes possess key enzymes that can behave as a reducing and stabilizing agent in nanomaterial production. This study reported the antibacterial activity of C.myrrh@GNP. The NPs exhibited the highest zone of inhibition towards all four (Bacil-
Fig. 17. Percent viability measured on human umbilical vein endothelial cells after treatment with present C. myrrh, HAuCl₃·3H₂O and C. myrrh@GNPs.

Table 1
Inhibitory activity % for HCT-116 and MCF-7 cell line at different concentrations of C.myrrh@GNPs.

| Sample           | Concentration | Standard | C.myrrh | GNPs  |
|------------------|---------------|----------|---------|-------|
| HCT-116 Inhibitory activity% | 100 µg/mL     | 99.41    | 47.50   | 96.50 |
|                  | 50 µg/mL      | 95.54    | 39.40   | 85.10 |
|                  | 25 µg/mL      | 92.89    | 26.53   | 70.41 |
|                  | 15 µg/mL      | 90.96    | 16.43   | 59.30 |
| MCF-7 Inhibitory activity%   | 100 µg/mL     | 98.18    | 38.32   | 79.92 |
|                  | 50 µg/mL      | 93.90    | 29.22   | 67.43 |
|                  | 25 µg/mL      | 91.59    | 18.69   | 55.20 |
|                  | 15 µg/mL      | 90.01    | 9.89    | 42.37 |
*lus subtilis*, *Staphylococcus aureus*, *Pseudomonas aeruginosa*, and *Escherichia coli*) microorganisms utilized. The maximum inhibition varies from $18 \pm 0.9$ mm to $22 \pm 0.5$ mm having the uppermost concentration of C. myrrh@GNPs next to every pathogen.

The highest inhibition zone was observed to be $22 \pm 0.5$ mm versus *B. subtilis*, $21 \pm 0.1$ mm versus *E. coli*, $20 \pm 0.7$ mm versus *S. aureus*, and $18 \pm 0.9$ mm versus *P. aeruginosa*. Whereas C. myrrh extract showed $17 \pm 0.1$ mm in *B. subtilis*, $16 \pm 0.2$ mm in *E. coli*, $15 \pm 0.5$ nm in *S. aureus*, and $14 \pm 0.9$ nm in *P. aeruginosa* at a maximum concentration as displayed in Table 2 and Fig. 22.

### 3.14. Molecular docking

The molecular interaction between drugs and DNA-proteins has been an exciting branch of research in chemical pharmacology and biology (Tian et al., 2008). The important role of peptides as hormones, enzyme inhibitors, neurotransmitters, and immunomodulatory in living systems plays a significant role in the treatment of many diseases (Cruciani et al., 1991; Mai et al., 2001).

The molecular interaction of *C. myrrh* compounds was studied against the three-dimensional complex structure of VEGFR-2.

![Fig. 18. Inhibitory activity % at different concentrations of C.myrrh@GNPs for (A) HCT-116 and (B) MCF-7 cell line.](image-url)
(PDB ID: 1YWN), ovine COX-1 (PDB ID: 3KK6), mouse COX-2 (PDB ID: 3LN1), α-amylase (PDB ID: 4GQR) and α-glucosidase (PDB ID: 5NN5). The obtained results and dock XP G-score and RMSD values (Å) are in Tables 3-7 and shown graphically in Figs. 23-27 and Fig. 1S-36S, supplementary materials.

The interaction of compounds under study can be classified as follow:

i. Hydrogen bonds to the protein, broken down into backbone and side chain, donor and acceptor.

ii. Hydrophobic interactions are broken down into π-π stacking (two aromatic groups stacked face-to-face or face-to-edge).

iii. Water bridge interactions that hydrogen bonding via a water bridge molecule, divided into protein donor and protein acceptor.

iv. The solvent exposure effect can only decrease binding affinity and it occurs when polar or charged groups of either the compound or protein were exposed to solvent leading to desolvation when being placed in contact with groups to which they cannot hydrogen bond effectively.

v. No interaction between molecules and the protein.

A cursory a glance at data reported in Tables 3-7 and shown graphically in Figs. 23-27 and Fig. 1S-36S, supplementary materials, one can conclude the following remarks:

**Fig. 19.** Inhibition of (HCT-116) and (MCF-7) cell lines by (A) C.myrrh and (B) C.myrrh@GNPs.
a. All C.myrrh compounds have interaction with α-amylase while about 50% of C.myrrh compounds have interaction with α-glucosidase which agrees with experimental inhibition of antidiabetics showing that α-amylase is greater than that of α-glucosidase.

b. δ-elemene has the highest dock XP G-score with −3.976 kcal/mol and RMSD = 0.987 Å towards VEGFR-2. Moreover, Furanoeudesma-1,3-dien has the highest dock XP G-score with −8.179 kcal/mol and RMSD = 1.128 Å towards COX-1.

Fig. 20. Anti-inflammatory inhibition of GNPs with C.myrrh extract.

Fig. 21. Anti-diabetic potential of C.myrrh@GNPs.
The aqueous extract of *C. myrrh* indicated the potential for reduction and stabilization of GNPs. When the extract concentration-dependent study was done it is found that the 6 ml extract has the potential to reduce so that the produced NPs possessed a spherical shape.

The absorption maxima were found to be 532 nm this is in the known range of GNPs in scientific research ([ElMitwalli et al., 2020; Kumari and Meena, 2020](#)). Here the extract volume also interestingly provides varying results at a small amount of volume, the absorption is very low with broadness, but increasing volume showed increased absorption i.e. 1.3 and the peak also got sharper and narrower so the best volume was at 6 ml and $\lambda_{\text{max}} = 532$ nm, that made the $\text{C. myrrh}$ @GNPs found out that 1.68 nm to 7.86 nm. The Surface Plasmon Resonance (SPR) for *C. myrrh*@GNPs exhibits two contents: first scattering and second absorption. The scattering content is identified as designated for fluorescence improvement while the absorption content is identified as fluorescence quencher ([Slocik et al., 2008](#)). The SPR spectrum is dependent on the size, and shape of the NPs, other than additionally being contingent on several outside possessions of the NPs environment like temperature, dielectric constant, medium, and refractive index of the solvent ([Unamaheswari et al., 2018](#)). The TEM images showed a circular shape below 10 nm size, this is in collaboration with earlier biosynthesis of GNPs. The NPs are homogenously bounded through Plasmon Resonance (SPR) for *C. myrrh*GNPs which was also supported by earlier reports on other plant extract-mediated GNPs ([Al-Radadi and Al-Youbi, 2018b](#); [Okitsu et al., 2009](#); [Singh and Srivastava, 2015](#); [Sun et al., 2009](#)). The size of the *C. myrrh*GNPs found out that 1.68 nm to 7.86 nm. The Surface Plasmon Resonance (SPR) for *C. myrrh*@GNPs exhibits two contents: first scattering and second absorption. The scattering content is identified as designated for fluorescence improvement while the absorption content is identified as fluorescence quencher ([Slocik et al., 2008](#)). The SPR spectrum is dependent on the size, and shape of the NPs, other than additionally being contingent on several outside possessions of the NPs environment like temperature, dielectric constant, medium, and refractive index of the solvent ([Unamaheswari et al., 2018](#)). The TEM images showed a circular shape below 10 nm size, this is in collaboration with earlier biosynthesis of GNPs. The NPs are homogenously bounded through Plasmon Resonance (SPR) for *C. myrrh*GNPs which was also supported by earlier reports on other plant extract-mediated GNPs ([Al-Radadi and Al-Youbi, 2018b](#); [Okitsu et al., 2009](#); [Singh and Srivastava, 2015](#); [Sun et al., 2009](#)). The size of the *C. myrrh*GNPs found out that 1.68 nm to 7.86 nm.

Table 2

| Microorganisms | Concentration (µg/mL) | C. myrrh | GNPs |
|----------------|----------------------|---------|------|
|                | 15                    | 3 ± 0.3 | 7 ± 0.2 |
|                | 25                    | 9 ± 0.1 | 12 ± 0.9 |
|                | 50                    | 13 ± 0.1 | 16 ± 0.3 |
|                | 100                   | 15 ± 0.5 | 20 ± 0.7 |
| Positive control | 24 ± 0.6 | 25 ± 0.6 |
| negative control | 0.0 ± 0.0 | 0.0 ± 0.0 |
| 15 µg/mL | 5 ± 0.1 | 8 ± 0.3 |
| 25 µg/mL | 11 ± 0.4 | 15 ± 0.6 |
| 50 µg/mL | 14 ± 0.8 | 19 ± 0.2 |
| 100 µg/mL | 17 ± 0.1 | 22 ± 0.5 |
| Positive control | 25 ± 0.3 | 24 ± 0.2 |
| negative control | 0.0 ± 0.0 | 0.0 ± 0.0 |
| 15 µg/mL | 4 ± 0.3 | 7 ± 0.6 |
| 25 µg/mL | 10 ± 0.1 | 12 ± 0.5 |
| 50 µg/mL | 13 ± 0.8 | 18 ± 0.7 |
| 100 µg/mL | 16 ± 0.2 | 21 ± 0.1 |
| Positive control | 23 ± 0.6 | 25 ± 0.6 |
| negative control | 0.0 ± 0.0 | 0.0 ± 0.0 |
| 15 µg/mL | 5 ± 0.6 | 5 ± 0.9 |
| 25 µg/mL | 8 ± 0.4 | 11 ± 0.1 |
| 50 µg/mL | 11 ± 0.5 | 15 ± 0.8 |
| 100 µg/mL | 14 ± 0.9 | 18 ± 0.9 |
| Positive control | 25 ± 0.6 | 24 ± 0.6 |
| negative control | 0.0 ± 0.0 | 0.0 ± 0.0 |

c. From the interaction with COX-2 and α-amylase, Lindestrene shows the highest dock XP G-score with −8.200 and −5.581 kcal/mol, respectively resulting from interaction through hydrogen bonds TYR341→(OH) for COX-2 and (OH)→(H2O) For α-amylase with distances 2.11 and 1.59 Å, respectively.

d. The highest dock XP G-score (−4.293 kcal/mol) with RMSD 1.500 Å for Furanodiene with TRP373 of COX-2 and Furanodiene and benzenmethanol-3-methoxy-phenyl with TRP59 and TRP481, respectively.

e. π - π stacking appeared in 2-tert-butyl-1,4-naphthoquinone and curzerene with TYR355 of COX-1. In addition, this interaction appears in Furanodiene with TRP373 of COX-2.

f. π - π stacking appeared for α-amylase and α-glucosidase in Furanodiene and benzenemethanol-3-methoxy-α-phenyl with TRP59 and TRP481, respectively.

g. T-Cadinol shows no interaction with VEGFR-2. While benzenemethanol-3-methoxy-α-phenyl has no interaction with both COX-1 and COX-2. Finally, β-elemene, Furanouedsesma-1,3-dien, β-Bourbonene, Curzerene, alloaromadendrene, β-Ylangene, α-Selinene, Elemol, Bicyclogermacrene, β-Caryophyllene, and Furanodiene have no interaction with α-glucosidase.

4. Discussion

The aqueous extract of *C. myrrh* indicated the potential for reduction and stabilization of GNPs. When the extract concentration-dependent study was done it is found that the 6 ml extract has the potential to reduce so that the produced NPs possessed a spherical shape.
outcomes got from the FTIR examination (Al-Radadi, 2021a; Ahmad et al., 2018; Vijayakumar et al., 2020). In FTIR the C = O (carbonyl gatherings) and hydroxide may be liable for the arrangement of the NPs and may assume a significant part in the adjustment of the shape of GNPs. The flavonoids that existed in the C. myrrh seed extract were a strong reducing potential which might be responsible for the reduction of gold salt. Hence it is observed that C.myrrh extract can fill double roles of reducing and stabilizing agent for C.myrrh@GNPs earlier reports of other plant extract showed if these molecules are present then its extract works as a dual function (Clarance et al., 2020; Shejawal et al., 2021; Yas et al., 2021). The C.myrrh@GNP was examined for crystallography using X-ray diffraction and found four major peaks of 2θ at 38.1, 44.5, 65, 78, and fifth at 82 which were allocated to (111), (200), (220), (311) and (222) crystalline planes respectively of gold. The peak ensuing to (111) was sufficiently tough as compared to other present planes suggesting that the C.myrrh extract produced crystalline GNP has crystalline nature and (111) was observed principal orientation (Al-Radadi, 2022b; Parida et al., 2011). The stability of the NPs was examined through DLS which was found to be (−10.8) mV. There was a discrepancy in TEM size and DLS size analysis of C.myrrh@GNP it is well known that DLS determines radii of gold and might the organic compounds surrounded by the GNPs synthesized by C.myrrh (Al-Radadi, 2021b; Khan et al., 2020; Perveen et al., 2021; Vijayakumar et al., 2020). The free radical scavenging activity using DPPH showed good improvement as compared to C.myrrh extract and its GNPs. Different concentrations such as from 10 to 100 μg/ml. The determined IC50 values for C.myrrh, DPPH, and C.myrrh@GNPs were 100, 93, and 84 correspondingly (Al-Radadi, 2021a; Khan et al., 02020). These findings are consistence with the earlier reports for different plants (Al-Radadi, 2022c; Kiran et al., 2021; Satpathy et al., 2020; Zhang et al., 2020) which presented that the plant extract while reacting with metal salts it loses its cytotoxicity (Al-Radadi, 2022b; Al-Radadi, 2021a; Mai et al., 2021; Parida et al., 2014). Anticancer activity against breast and colon cell lines showed a dose-dependent response. Amongst the different concentrations utilized 100 μg/ml concentration was inhibitory the concentration of
C.myrrh@GNPs for which it is found excellent inhibition as compared to C.myrrh extract. These outcomes recommended that the antioxidant activity designate concentration and dose-dependent (Al-Radadi, 2022c; Perveen et al., 2021; Al-Radadi, 2021a; Shejawal et al., 2021; Siegel et al., 2013; Anand et al., 2015; Wang et al., 2019). The inflammatory study was also carried out which showed that the GNP’s were used to inhibit COX-1, and COX-2, it is found that the response was having direct contact with the concentration of C.myrrh@GNPs here it is observed that 400 µg/ml concentration showed excellent inhibition of COX-1 and COX-2. These results exhibited that the phenolics and flavonoids in C. myrrh extract impart excellent anti-inflammatory stress via numerous mechanisms (Al-Radadi, 2022b; Hwang et al., 2015; Singh et al., 2018). The C.myrrh@GNPs used to treat α-amylase and α-glucosidase enzymes to detect the anti-diabetic property of C.myrrh@GNPs was found to be increased when concentration was increased. The 400 µg/ml was used in a dose-dependent manner. The percent inhibition was observed at 50 ± 0.58 and 44 ± 0.49 for α-amylase and α-glucosidase enzymes respectively (Al-Radadi, 2022b; Kiran et al., 2021; Senthilkumar et al., 2019; Vijayakumar et al., 2020). The findings of the antibacterial study exhibited that the C.myrrh@GNPs at 100 µg/ml concentrations may retard the bacterial growth of pathogens liable for food toxicity (Al-Radadi, 2021b; Siegel et al., 2013). The medicinal value also reported the

### Table 3
Molecular interactions of C.myrrh compounds predicted for inhibitor binding to VEGFR-2.

| Compound                                      | Docking score | RMSD (Å) | Interaction Bond Type distance (Å) |
|-----------------------------------------------|---------------|----------|-----------------------------------|
| i-elemene                                     | -3.976        | 0.987    | -                                 |
| Furanoeudesma-1,3-dien                        | -3.912        | 0.967    | -                                 |
| β-Bourbonene                                  | -3.806        | 1.199    | H-Bond 2.21                       |
| menthofuran                                   | -3.568        | 1.188    | (H₂O)–(C=O) hetero-ring H-Bond 2.22 |
| 3,4,4-trimethyl-3-(3-oxobut-1-enyl)bicyclo(4.1.0)heptan-2one | -3.537 | 1.480 | ASP1044–(C = O) H-Bond 2.22 |
| 1-Elemene                                     | -3.507        | 0.970    | -                                 |
| 2-tert-butyl-1,4-naphthoquinone               | -3.449        | 1.083    | -                                 |
| α-Humulene                                    | -3.433        | 0.958    | -                                 |
| Curzerene                                     | -3.371        | 0.921    | -                                 |
| allo-Aromadendrene                            | -3.325        | 1.141    | Solvent exposure –                |
| β-Ylangene                                    | -3.222        | 0.764    | Solvent exposure –                |
| benzene methanol-3-methoxy-α-phenyl           | -3.205        | 0.887    | (OH)–(H₂O) H-Bond 1.69           |
| β-Selinene                                    | -3.165        | 1.259    | Solvent exposure –                |
| β-elemene                                     | -3.160        | 0.790    | Solvent exposure –                |
| Elemol                                        | -3.160        | 0.757    | (OH) → ASP1044 H-Bond 1.93       |
| γ-Murolene                                    | -2.951        | 1.502    | -                                 |
| Bicyclogermacrene                             | -2.805        | 1.228    | -                                 |
| β-Caryophyllene                               | -2.709        | 1.869    | -                                 |
| α-selinene                                    | -2.642        | 1.375    | Solvent exposure –                |
| α-Copaene                                     | -2.488        | 1.353    | Solvent exposure –                |
| Lindestrene                                   | -2.425        | 1.788    | Solvent exposure –                |
| 2-methoxyFuranodiene                          | -2.287        | 0.764    | Solvent exposure –                |
| Furanodiene                                   | -1.655        | 1.390    | Solvent exposure –                |
| T-Cadinol                                     | No interaction|         | -                                 |

### Table 4
Molecular interactions of C.myrrh compounds predicted for inhibitor binding to COX-1.

| Compound                                      | Docking score | RMSD (Å) | Interaction Bond Type distance (Å) |
|-----------------------------------------------|---------------|----------|-----------------------------------|
| Furanoeudesma-1,3-dien                        | -8.179        | 1.218    | -                                 |
| Furanodiene                                   | -7.834        | 1.664    | -                                 |
| δ-elemene                                     | -7.436        | 1.029    | -                                 |
| β-Caryophyllene                               | -7.411        | 1.685    | -                                 |
| 2-tert-butyl-1,4-naphthoquinone               | -7.206        | 0.927    | Aromatic ring—TYR355 π-π stacking 5.33 |
| Lindestrene                                   | -6.866        | 0.905    | -                                 |
| β-Ylangene                                    | -6.854        | 1.372    | -                                 |
| γ-Murolene                                    | -6.750        | 0.904    | -                                 |
| β-elemene                                     | -6.666        | 1.074    | -                                 |
| α-selinene                                    | -6.523        | 1.039    | -                                 |
| β-Selinene                                    | -6.516        | 0.894    | -                                 |
| α-Copaene                                     | -6.506        | 0.462    | -                                 |
| α-Humulene                                    | -6.384        | 0.694    | -                                 |
| β-Bourbonene                                  | -6.345        | 0.767    | -                                 |
| Bicyclogermacrene                             | -6.338        | 1.292    | -                                 |
| Menthofuran                                   | -6.185        | 0.981    | -                                 |
| γ-Elemene                                     | -6.044        | 0.655    | -                                 |
| allo-Aromadendrene                            | -5.986        | 0.870    | -                                 |
| Curzerene                                     | -5.720        | 1.693    | ARG120–(C=O) ring             H-Bond 1.83 |
| 3,4,4-trimethyl-3-(3-oxobut-1-enyl)bicyclo(4.1.0)heptan-2one | -5.532 | 0.861 | Furan ring—TYR355 π-π stacking 4.61 |
| T-cadinol                                     | -5.411        | 0.871    | -                                 |
| Elemol                                        | -5.359        | 1.693    | -                                 |
| 2-methoxyFuranodiene                          | -3.802        | 0.931    | Solvent exposure –                |
| benzene methanol-3-methoxy-α-phenyl           | No interaction|         | -                                 |
effect of C.myrrh extract against the diabetics and antibacterial but here GNPs capped with the C.myrrh extract have shown the excellent report on diabetic and antibacterial activity (Khalil et al, 2020; Al-Radadi, 2022b). The reason for this antibacterial activity is that the C.myrrh extract contains many phytochemicals that were capped around the GNPs (Al-Radadi, 2022c).

5. Future perspective

Some antibacterial, anti-inflammatory and anti-diabetic medications are being repurposed in clinical trials globally and several potential findings been lately been stated. In this case, NPs attract the center and are thought to be a viable substitute. Additionally,
C.myrrh@GNP found potential findings that these NPs can be applied against pathogenic bacteria, as an anti-inflammatory and anti-diabetic, as well as anticancer agent. The C.myrrh@GNP is currently exhibiting the potential revolutionary agent which will create a constructive and encouraging impact on biomedical fields in the future.

6. Conclusions

The C.myrrh is of high medicinal importance. This method of synthesis of GNPs will obviously increase the total medicinal value of the GNPs capped by C.myrrh biomolecules. The overall finding of the work enlightens the medicinal importance of C.myrrh@GNPs. The method is the most effective method, regarding price, and energy and it also exhibits simplicity. GNPs characterized by spectroscopic and microscopic techniques revealed that there is a direct correlation between volume and SPR peak of the synthesized GNP. The size structure and morphology of the nanomaterials are spherical, crystalline, and with an average size of 4 nm, made up of metallic gold. The functional molecules surrounding the surface of the NPs are coming from the C.myrrh extract and contributing to the cell toxicity, anti-biotic (antibacterial), anti-inflammatory, antidiabetic and antioxidant properties. C.myrrh@GNPs demonstrated a remarkable cell line proliferation inhibition activity against MCF-7 and HCT-116 cell lines. The molecular docking of Polyphenol exhibits the high affinity of Polyphenol against HCT-116, inflammatory and diabetic.

Table 7

| Compound | Docking score | RMSD (Å) | Interaction | Bond Type | distance (Å) |
|----------|---------------|----------|-------------|-----------|--------------|
| 3,4,4-trimethyl-3-(3-oxobut-1-enyl)bicyclo(4,1,0)heptan-2-on | -4.293 | 1.500 | ARG600 → (C = O) | H-Bond | 2.15 |
| T-cadinol | -4.245 | 1.715 | (OH) → ASP616 | H-Bond | 1.49 |
| α-Humulene | -3.952 | 1.697 | – | – | – |
| α-Copaene | -3.243 | 1.826 | Solvent exposure | – | – |
| β-Selinene | -3.195 | 1.897 | Solvent exposure | – | – |
| benzene methanol-3-methoxy-α-phenyl | -2.684 | 1.083 | (OH) → ASP282 | H-Bond | 1.95 |
| δ-elemene | -2.515 | 1.187 | Solvent exposure | – | – |
| γ-Murolene | -1.857 | 1.148 | – | – | – |
| Lindestrene | -1.757 | 0.968 | TRP618 → (H₂O) → (C = O) | H-Bond via water molecule | 1.98, 2.11 |
| 2-methoxyFurane | -1.477 | 1.126 | TRP618 → (H₂O) → (C = O) | H-Bond via water molecule | 1.98, 2.11 |
| 2-tert-butyl-1,4-naphthoquinone | -1.373 | 1.126 | ALA284 → (C = O) | H-Bond | 2.33 |
| menthofuran | -0.909 | 0.964 | TRP618 → (H₂O) → (C = O) | H-Bond via water molecule | 1.98, 2.18 |
| γ-Elemene | 1.254 | 1.000 | Solvent exposure | – | – |
| δ-elemene | No interaction | – | – | – | – |
| Furaneodesma-1,3-dien | No interaction | – | – | – | – |
| β-Bourbonene | No interaction | – | – | – | – |
| Curzerene | No interaction | – | – | – | – |
| allo-Aromadendrene | No interaction | – | – | – | – |
| β-Ylangene | No interaction | – | – | – | – |
| α-Selinene | No interaction | – | – | – | – |
| Elemol | No interaction | – | – | – | – |
| Bicyclogermacrene | No interaction | – | – | – | – |
| β-Caryophyllene | No interaction | – | – | – | – |
| Furanodiene | No interaction | – | – | – | – |

Fig. 23. 3D and 2D molecular interaction of δ-elemene for inhibitor to VEGFR-2.
Fig. 24. 3D and 2D molecular interaction of Furanoeudesma-1,3-dien for inhibitor to COX-1.

Fig. 25. 3D and 2D molecular interaction of Lindestrene for inhibitor to COX-2.

Fig. 26. 3D and 2D molecular interaction of Lindestrene for inhibitor to α-amylase.
Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jsps.2022.06.028.

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