GC–MS and Network Pharmacology Analysis of the Ayurvedic Fermented Medicine, Chandanasava, Against Chronic Kidney and Cardiovascular Diseases

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Abstract
Chandanasava is an Ayurvedic polyherbal fermented traditional medicine (FTM) used by traditional practitioners for millennia. Nevertheless, the mode of action and functional targets are still unknown. The current study includes a pharmacological network analysis to identify the Chandanasava compounds interacting with target proteins involved in chronic kidney disease (CKD) and cardiovascular disease (CVD). Sixty-one Chandanasava phytochemicals were obtained by GC–MS and screened using the Traditional Chinese Medicine Systems Pharmacology Database (TCMSP). The disease target genes were obtained from DisGeNET and GeneCards databases. Forty-five phytocompounds and 135 potential targets were screened for CKD and CVD target proteins and protein interaction networks were constructed. The pharmacological network was deciphered employing target proteins involved in the mechanical action of Chandanasava. The results indicated that 10 bioactive compounds exhibited higher binding affinity patterns with the screened 42 CKD and CVD target proteins. Gene Ontology and KEGG analysis revealed target pathways involved in CKD and CVD, which were further explored by detailed analysis and network-coupled drug profile screening. The molecular docking results showed piperine and melatonin as effective inhibitors/regulators of the hub genes of CKD and CVD. The current study establishing authentic bioactive compounds in FTM is based on deeper insights into recognized Ayurvedic medicines.

Keywords Network pharmacology · Phytoactive chemical screening · Protein–protein interaction network · Drug-protein interaction network · Molecular docking studies · GC–MS analysis

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Introduction

Chronic kidney diseases (CKD) and cardiovascular diseases (CVD) correspond to multifaceted, complicated disorders affecting a global human population, causing escalated mortality and morbidity rates owing to hospice conditions and survival challenges [1]. The negative correlations are aggravated in non-dialysis-dependent renal failure, myocardial ischemia, heart failure, and hypertension [2, 3]. End-stage renal disease patients are corroborated for an increased risk for cardiovascular complications and necessitate proper management [4]. Glomerular filtration rate and myocardial infarction have been associated with many population-based randomized trials [5, 6]. Life expectancy and increased prevalence of CVD and CKD among diabetic patients require impact assessment for cost-effective protocols, economic burden, and premature mortality management [7]. Ayurveda possesses sufficient potential to be used to prevent and treat various ailments [8]. The Ayurvedic pharmacopeia contains a variety of medications, including fermented traditional medicines (FTM), which are classified as Arishta (decoctions) and Asava (infusions), comprising polyherbal formulations invigorated by naturally occurring intrinsic microorganisms. Arishta and Asava are traditionally regarded as distinct and effective therapeutic formulations due to their improved storage efficiency, strengthened pharmacological activities, improved separation of bioactive compounds from herbal ingredients, and increased efficacy of dosage forms for enhanced health benefits [9].

Arishta and Asava formulations represent the distinct herbal fermentation products have been identified previously based on their composition and therapeutic uses [10, 11]. Furthermore, they are used to treat a plethora of disease conditions in pediatrics, systemic diseases affecting the central nervous system, circulatory system, respiratory system, digestive system, urinary system, reproductive systems, immune disorders, skin diseases, poisonous insect and snake bites, and alcoholism together with commonly encountered illnesses [11, 12]. Among the Arishta and Asava, Chandanasava is the most used medicine composed of 24 different polyherbal components and prescribed to treat gastrointestinal diseases, urinary disorders, spermatorrhea, gonorrhea, and autoimmune disorders as well as a diuretic appetizer, and has a cooling effect throughout the body [10, 11]. It is based on classical Ayurvedic text, namely, Bhaishajya Ratnavali in Sanskrit, with the known composition [11, 13] (Supplementary table S1). Singular composition rather than the combinatorial formulation of ArishtalAsava shows a wide variety of pharmacological effects rendering significant medicinal attributes for advantageous applications [14, 15]. The bioactivity is likely to be much more than the therapeutic effects described in Ayurvedic traditional knowledge due to the presence of a wide range of phytochemicals. Ayurvedic medicaments balance the tridoshas in the body and hence cure multiple diseases simultaneously. The enormity of traditional Ayurvedic formulations dates back to the ages for confronting the ancient and rich knowledge of Charaka Samhita in authenticity and authoritative medical applications of Indian origin [16]. Further ample literature is affirmative that traditional Ayurvedic medicines are included in alternative medicinal realms based on Ashtangahrudya and Bhaishajya Ratnavali [17, 18]. Moreover, therapeutic potentials of phytochemical Ayurvedic secondary metabolites have been included in the clinical trial literature and proved efficacious [19]. Nevertheless, natural product chemistry and drug discovery research are more inclined towards big data analytics for envisaging therapeutic roles in confirmed health augmentation and disease alleviation [20]. Hence, bioinformatics tools and computational biology platforms are largely resourced as effective tools for medicinal plant research in drug discovery, development, and deeper insights [21]. Databases for
screening dietary supplements and plant metabolites were emphasized for rational applications in phytomedicine [22]. Further health implications based on phytochemicals of plant origin were directed for advantageous applications in the arena of computational informatics [23]. In recent times, the network pharmacology approach has been used to examine the underlying mechanisms of TIM and TCM in order to effectively investigate the therapeutic efficacy using curated biological databases [24, 25]. In view of these, the current study is aimed to investigate the phytochemicals from an Indian fermented traditional medicine, Chandanasava, to treat chronic kidney and cardiovascular diseases by network pharmacology-based approach.

Materials and Methods

Sample Collection

Ayurvedic polyherbal fermented traditional medicine (FTM), Chandanasava, was procured from Ashtanga Ayurvedics PVT, Limited, Tiruchirapalli, Tamil Nadu, India.

Solvent Extraction

Hexane and ethylacetate solvents were employed for Chandanasava in a liquid–liquid separation technique with minor modifications. A sample of 5 mL of Chandanasava and 100 mL of solvents were mixed in a 500-mL separating funnel and shaken well steadily for 15 min. After 5 min, the solvent phase was detached. The solvent phase was then harvested, and these steps were repeated two times using the same sample. Sodium thiosulphate (anhydrous) was added to the mixture extracts to eliminate moisture levels and condensed with a rotary concentrator (BUCHI R-210, Buchi India Pvt Ltd), then purged with nitrogen gas till 1 mL was obtained.

Gas Chromatography‑Mass Spectrometry (GC–MS) Analysis of Ayurvedic FTM

Perkin Elmer Clarus 500 (CT, USA) instrument with a flame ionization detector with a capillary column (30 m length × 0.25 mm ID) coated with 5% phenyl 95% dimethylpolysiloxane with a film thickness of 0.25 μm was used for GC–MS analysis. The carrier gas was helium with a flow rate of 1 mL/min and the injection port was maintained at 280 °C and 1 μL of the sample was injected. The initial temperature of the column oven was 60 °C, ramped to 300 °C at a rate of 10°C/min⁻¹. In mass spectrometry, the full scan mode and 40–450 Daltons scan range were optimized for spectral data collection. The National Institute of Standards and Technology library was utilized to cross-match the obtained peaks with the standard.

Screening of Chemical Components from Biological Databases

Traditional Chinese Medicine Systems Pharmacology and Analysis Platform (TCMSP) were used to profile the phytoconstituents in Chandanasava. The respective constituents were identified using the GC–MS analysis and compounds were screened using the database [26]. Chandanasava’s phytoactive constituents were screened for oral bioavailability
(OB) (> 30%) and drug-likeness (DL) (0:18) for assimilatory profiles. Absorption, distribution, metabolism, and excretion (ADME) properties of medicines and their oral administration in the bloodstream correspond to oral bioavailability [27]. The Pub Chem (https://pubchem.ncbi.nlm.nih.gov/) and Drug Bank (https://www.drugbank.ca/drugs) were used to obtain chemical information on the constituents, including their molecular formula and gene codes.

**Screening of Phytochemicals and Diseases-Related Target Genes**

DisGeNET and GeneCards database systems were used to acquire target genes information correlated to CKD with cardiovascular complications [28]. These databases were searched using the keywords “chronic kidney diseases” and “cardiovascular diseases” to identify the targeted disease and related genes.

**Protein–Protein Interaction (PPI) Network Construction Based on Drug-Protein Interaction**

A rough and basic interaction involving target genes for relevant bioactive metabolites in Chandanasava was built using the STRING version 11.0 to gather knowledge about the PPI network [29]. To validate the accuracy of our study, the species chosen is “Homo sapiens,” and the protein interaction is computed to a 70% similarity (0.700) high confidence level. The identified genes associated with the target diseases were cross-verified with the STITCH v5.0 server [30]. Cytoscape 3.8.0 software was used to build and validate the final PPI network based on the draft network data [31]. The network analyzer module in the Cytoscape is utilized for the network validation [32]. Cytohubba module in Cytoscape was used for profiling the disease hub genes associated with the disease targets [33].

**Gene Ontology (GO) and KEGG Pathway Enrichment Analysis**

Polyherbal Ayurvedic FTM Chandanasava components for target genes of CKD and CVD genes and appropriate for gene function and signaling cascades were obtained from the DAVID database system GO functional annotation, and KEGG pathway enrichment for deriving the biological process (BP), cellular components (CC), and molecular function (MF) of proteins used was analyzed [34].

**Target Protein Pathway Network Construction**

Enriched pathway associated with target diseased genes was further imported into the Cytoscape to derive an “Ingredient-Target-Pathway.” The network interconnections of Chandanasava constituents in the target protein pathway were built using nodes corresponding to diseases, targets, and associated pathways. The edges were elaborated on the connections between the nodes. A network model map for Chandanasava components and target genes for chronic renal and cardiovascular disease has subsequently arrived for a preliminary theoretical assessment of targeted medications.
Molecular Docking Analysis

Data Bank (https://www.rcsb.org/) and Drug Bank (a familiarity foot of pharmaceutical library) were used to obtain the tertiary structures of the top-hit hub proteins and generated secondary metabolites. The protein structures were prepared by removing hydrogen from the receptor and subtracting water molecules. Flexible docking with AUTODOCK VINA was used to study the molecular interaction between the top-hit hub proteins and ligands and binding sites were automatically detected [35]. The LigPlot+ software was used to identify the molecular interaction in a 2-dimensional view [36]. Finally, hydrogen bonds, hydrophobic interactions, and binding scores of the docked complex were noted.

Results

GC–MS Profiling of Chandanasava

The hexane and ethyl acetate fractions of the Chandanasava formulation were analyzed using gas chromatography-mass spectrometry. As illustrated in the overlay plot (Fig. 1A and B), the GC–MS spectra indicated the presence of a broad set of secondary metabolites that differ in retention times, molecular formulae, and peak areas. The total ion chromatogram of the Chandanasava formulation revealed that most of the compounds had a similar type of metabolic profile, while others exhibited variance (Fig. 1A and B). The peaks were integrated and correlated to the spectra databases available in the GC-MS libraries (NIST and Wiley library). In Chandanasava formulations, a total of 62 secondary metabolites were identified using GC–MS. Heterocyclic compounds, hydrocarbons, esters, acids, amines, halides, ethers, and aldehydes were present in the Chandanasava formulation.

Screening of Targets of Phytoconstituents from Chandanasava

The identified 61 compounds were searched by chemical name strategy using the TCMSP database to screen bioactive components in Chandanasava. The potential and pharmacologically effective phytochemicals were screened with the value of ≥ 30% OB and ≥ 0.18 DL (Table 1). The phytochemical compounds of the canonical SMILES are represented in Table 2. In a total of 61 compounds, 16 (MOL004067, MOL003353, MOL005028, MOL008670, MOL001573, MOL008712, MOL001230, MOL012364, MOL005117, MOL002139, MOL001212, MOL002092, MOL007525, MOL004791, MOL001212, and MOL001815) were eliminated due to the lack of related diseases and target proteins involved in CKD and CVD. After eliminating overlaps from 418 target genes, a total of 135 target genes for CKD and CVD were identified (Supplementary Table S2). Cytoscape 3.8.0 software view of the interaction of the 45 bioactive Chandanasava metabolites with their respective target genes was represented (Fig. 2). The analysis of component-target network consisted of 180 clusters and 418 edges. The red diamond clusters corresponded to the 45 phytocompounds, while the 7 yellow and 15 blue circle clusters corresponded to chronic kidney disease and cardiovascular disease, respectively. Moreover, 20 pink circle nodes represented both common genes of chronic kidney and CVD. The remaining 93 green circle nodes are attributed to the other target-related genes of the bioactive compounds. Eugenol, stigmasterol, isonipecotic acid,
1-methyl-4-phenyl-, ethyl ester, piperine, elemicin, 1,2-benzenedicarboxylic acid, dibutyl ester, and 5-hydroxymethylfurfural were found to have a strong interaction with multiple target genes among the 45 metabolites screened from Chandanasava. The results indicated that these active metabolites were the key components in the network which were attributed to the treatment of CKD and CVD.

![Fig. 1](image)

Fig. 1 Identification of Chandanasava ingredients by GC–MS using solvent extracts (A) ethyl acetate fraction and (B) hexane fraction
Table 1 Representation of molecular properties from traditional Chinese medicine systems pharmacology database and analysis platform for the chemical components of Chandanasava

| MOL ID   | Molecule name                  | Molecular formula | CAS No. | MW (g/mol) | OB%  | BB%  | DL%  | HL%  | Structural formula |
|----------|--------------------------------|-------------------|---------|------------|------|------|------|------|-------------------|
| MOL000449| Stigmasterol                   | C_{29}H_{48}O_{8}  | 3-48-7  | 412.77     | 43.83| 1.00 | 0.76 | 5.57 |                   |
| MOL003064| Quinoline                      | C_{9}H_{7}N_{9}    | 1-22-5  | 129.17     | 35.25| 1.71 | 0.03 | 11.89|                   |
| MOL004067| Nootkatone                     | C_{6}H_{12}O_{6}   | 4674-50-4| 218.37     | 33.04| 1.51 | 0.10 | 4.39 |                   |
| MOL001431| Melatonin                      | C_{23}H_{36}N_{2} | 73-31-4 | 232.31     | 52.96| 0.49 | 0.11 | -1.27|                   |
| MOL00353 | 9(10H)-Anthracenone, 1,3,8-trihydroxy-6-methyl- | C_{16}H_{12}O_{6} | 491-60-1| 256.27     | 24.72| 0.24 | 0.21 | N/A  |                   |
| MOL009594| Isonipecotic acid, 1-methyl-4-phenyl-, ethyl ester | C_{15}H_{22}N_{2}O_{4} | 57-42-1 | 247.37     | 77.48| 1.36 | 0.10 | 3.78 |                   |
| MOL008475| Formosan-16-carboxylic acid, 19-methyl-2-oxo-, (19alpha)- | C_{20}H_{22}N_{2}O_{4} | 10126-00-8 | 354.44     | 31.70|-0.17 | 0.70 | 10.40 |                   |
| MOL009475| Anethole &lt(E)&gt;            | C_{10}H_{12}O       | 104-46-1| 148.22     | 32.49| 1.81 | 0.03 | 1.68 |                   |
| MOL00528 | Cannabinol                     | C_{21}H_{26}O_{2}  | 521-35-7| 310.42     | 22.04| 1.05 | 0.32 | N/A  |                   |
| MOL010749| 1H-Pyrazole, 1,3-dimethyl-      | C_{5}H_{8}N_{2}    | 694-38-4| 96.15      | 50.33| 1.77 | 0.01 | 4.17 |                   |
| MOL00670 | Fluoroacetamide                | C_{5}H_{10}O_{2}   | 640-19-7| 77.07      | 49.07| 0.27 | 0.00 | 12.04|                   |
| MOL007580| Picoline &lt;meta&gt;          | C_{6}H_{10}N       | 108-99-6| 93.14      | 73.75| 1.88 | 0.01 | 11.95|                   |
| MOL007197| Benzenamine, N-phenyl-         | C_{12}H_{14}N       | 122-39-4| 169.24     | 31.13| 1.87 | 0.05 | 7.72 |                   |
| MOL006767| 2,6-Pyridinecarboxylic acid, 4-(4-amino-1-carboxy-4-oxobutyl)jimino)ethyliden)-1,2,3,4-tetrahydro- | C_{10}H_{10}N_{2}O_{7} | 904-62-1 | 339.34     | 56.14|-2.19 | 0.26 | 37.55|                   |
| MOL00032 | Eudesmol &lt;beta&gt;          | C_{15}H_{20}O_{6}  | 473-15-4| 222.41     | 26.09| 1.38 | 0.10 | N/A  |                   |
| MOL000991| Cinnamaldehyde                 | C_{9}H_{10}O       | 104-55-2| 132.17     | 31.99| 1.48 | 0.02 | 4.73 |                   |
| MOL006469| 9,10-Anthracenedione           | C_{16}H_{14}O_{5}  | 84-65-1 | 208.22     | 56.10| 0.29 | 0.14 | 33.53|                   |
| MOL001580| Piperonyl aldehyde             | C_{12}H_{18}O_{2}  | 120-25-7| 150.14     | 32.74| 0.64 | 0.04 | 3.73 |                   |
| MOL003050| Pelargonic acid                | C_{9}H_{18}O_{2}   | 112-05-0| 158.27     | 40.51| 1.08 | 0.02 | 4.15 |                   |

Screening of Common Targets in Chandanasava for CKD and CVD

The common and corresponding targets for 27 CKD- and 35 CVD-related genes were obtained from the DisGeNET and GeneCards databases and were represented in the Venn diagram (Fig. 3A and B). The data output about these disease target genes obtained from DisGeNET and GeneCards was given (Supplementary Tables S3 and S4). The CKD- and CVD-associated genes interaction with 42 potential target genes obtained from Chandanasava components are represented in Table 3.
Table 1 (continued)

| MOL005549 | 2H-pyran-2-one, 5,6-dihydro-4-methyl- | C₂H₄O₂ | 2381-87-5 | 112.14 | 53.46 | 1.39 | 0.02 | 11.55 |
| MOL000953 | Cholest-5-en-3-ol, (3.beta.)- | C₂H₄O | 57-88-5 | 386.73 | 37.87 | 1.13 | 0.68 | 4.52 |
| MOL001192 | Caryophyllene alcohol | C₁₅H₂₀ | 472-97-9 | 220.44 | 38.06 | 2.08 | 0.13 | 10.07 |
| MOL009238 | Eugenol <methoxy-> | C₁₇H₁₆O₃ | 8627-88-9 | 194.25 | 65.28 | 1.35 | 0.05 | 3.10 |
| MOL001787 | Beta-D-Ribofuranose, 1-{(6-amino-9H-purin-9-yl)-1-deoxy-} | C₁₀H₁₃N₅O₄ | 58-61-7 | 267.28 | 15.98 | -2.22 | 0.18 | N/A |
| MOL001573 | Hydrocinnamate <methyl-> | C₂₅H₂₂O₄ | 103-25-3 | 164.22 | 29.05 | 1.31 | 0.03 | N/A |
| MOL004650 | Vanillin acetate | C₁₃H₁₀O₂ | 881-68-5 | 194.20 | 28.17 | 0.38 | 0.06 | N/A |
| MOL010419 | Delta-Tocopherol | C₂₀H₃₂O₂ | 119-13-1 | 402.73 | 16.36 | 1.35 | 0.48 | N/A |
| MOL010904 | Chamazulene | C₃₀H₄₂O | 520-05-5 | 184.30 | 25.35 | 1.84 | 0.06 | N/A |
| MOL008712 | Heptanoate <ethyl-> | C₁₅H₂₄O₂ | 106-30-9 | 158.27 | 45.19 | 1.28 | 0.02 | 4.20 |
| MOL003790 | Androstanone | C₁₉H₂₆ | 438-22-2 | 260.51 | 32.50 | 2.05 | 0.26 | 4.37 |
| MOL009677 | Lanost-8-en-3beta-ol | C₁₅H₂₀O | 79-62-9 | 428.82 | 34.23 | 1.25 | 0.74 | 5.48 |
| MOL006290 | N-amidinosarcosine | C₁₁H₁₆N₄O₂ | 57-00-1 | 131.16 | 60.47 | 0.13 | 0.02 | 11.53 |
| MOL003795 | 2-Piperidine carboxylic acid | C₁₉H₁₈N₂O₂ | 535-75-1 | 129.18 | 66.14 | 0.38 | 0.02 | 11.08 |
| MOL004366 | Formate <octyl-> | C₄H₉O₂ | 112-32-3 | 158.27 | 53.32 | 1.24 | 0.02 | 6.99 |
| MOL001684 | Isoquinoline, 6-amino- | C₁₇H₁₄N₂ | 23687-26-5 | 144.19 | 30.46 | 0.11 | 0.04 | -2.16 |
| MOL001592 | Piperine | C₁₀H₁₀NO₃ | 94-62-2 | 285.37 | 42.52 | 0.62 | 0.23 | 10.25 |
| MOL001230 | Undecyl alcohol | C₁₁H₂₂O | 112-42-5 | 172.35 | 21.94 | 1.23 | 0.03 | N/A |
| MOL012364 | Elemol | C₁₅H₂₆O | 639-99-6 | 222.41 | 31.91 | 1.32 | 0.07 | 8.14 |
| MOL003483 | Caprylic acid ethyl ester | C₁₀H₂₂O₂ | 106-32-1 | 172.30 | 33.05 | 1.30 | 0.03 | 4.60 |
| MOL009811 | 2-Ethyl-5,5-dimethylpyrazine | C₁₁H₁₂N₂ | 13925-07-0 | 136.22 | 36.17 | 1.08 | 0.02 | 11.44 |

Construction and Evaluation of Target Genes PPI Network

The STRING database was used to analyze selected CKD-associated target genes [28] and CVD-associated target genes [37], which were shown as an interaction network with a high confidence score of 0.700. For CKD, the network of PPI interactions between target genes was constructed using 26 clusters, 180 edges, and an average node degree of 13.8. PPI of CVD target genes, a total of 33 clusters and 191 edges, and
**Table 1** (continued)

| Compound ID | Name                          | Formula | Molecular Weight | MW | OB | BBB | DL | HL |
|-------------|-------------------------------|---------|------------------|----|----|-----|----|----|
| MOL004466  | (S)-1-Phenyl-2-propylamine    | C9H13N  | 135.23           | 44.85 | 1.14 | 0.02 | -2.48 |
| MOL004783  | Nonane                        | C9H20   | 128.29           | 29.23 | 1.95 | 0.01 | N/A |
| MOL010529  | Jasmone <(Z)>                | C9H16O  | 164.27           | 32.07 | 1.50 | 0.03 | 2.71 |
| MOL05117   | Anisyl alcohol <para->        | C10H10O | 105-13-5         | 138.18 | 5.64 | 0.79 | 0.02 | N/A |
| MOL012192  | Labdanol                      | C14H18O2| 204.29           | 26.47 | 1.33 | 0.06 | N/A |
| MOL000269  | Elemicin                      | C10H10O | 208.28           | 21.94 | 1.28 | 0.06 | N/A |
| MOL005117  | Anisyl alcohol               | C8H10O  | 120-57-0         | 32.74 | 0.64 | 0.04 | 3.73 |
| MOL012192  | Labdanol                      | C14H18O2| 140-10-3         | 19.68 | 0.96 | 0.03 | N/A |
| MOL007180  | Beta-Tocopherol               | C24H34O4S| 940.69          | 32.29 | -0.58 | 0.70 | 0.04 |
| MOL002295  | Cinnamic acid                | C9H8O2  | 330.36           | 8.46  | 0.08 | 0.33 | N/A |
| MOL002139  | Acetylphenylhydrazine         | C8H10N2O| 114-83-0         | 24.76 | 1.15 | 0.03 | N/A |
| MOL001300  | Phenylethyl alcohol           | C8H10O  | 150.20           | 24.76 | 1.15 | 0.03 | N/A |
| MOL00748   | 5-Hydroxymethylfurfural      | C4H5O   | 126.12           | 45.07 | -0.27 | 0.02 | 11.73 |
| MOL01212   | 1-Tetradecanol               | C14H30O | 214.44           | 14.19 | 0.98 | 0.05 | N/A |
| MOL02092   | 2,4-Di-tert-butylphenol      | C14H18O | 206.36           | 26.74 | 1.81 | 0.06 | N/A |
| MOL07525   | 3-Eicosene, (E)-             | C20H38   | 280.60           | 18.19 | 1.92 | 0.13 | N/A |
| MOL03509   | 1-Nonanol                    | C13H26O | 144.29           | 33.19 | 1.15 | 0.01 | 6.07 |
| MOL04791   | 1-Hexadecanol                | C18H36O | 242.50           | 13.32 | 1.07 | 0.08 | N/A |
| MOL00676   | 1,2-benzenedicarboxylic acid, dibutyl ester | C22H20O4| 278.38           | 64.54 | 0.56 | 0.13 | 5.41 |
| MOL01212   | Myristic alcohol             | C14H26O | 214.44           | 14.19 | 0.98 | 0.05 | N/A |
| MOL01815   | 1-Tricosene                  | C30H48   | 322.69           | 16.25 | 1.92 | 0.21 | N/A |

*MW,* molecular weight, *OB,* oral bioavailability, *BBB,* blood–brain barrier, *DL,* drug-likeness, *HL,* half-life
| Molecule name                                                        | Canonical SMILES                                                                 |
|--------------------------------------------------------------------|----------------------------------------------------------------------------------|
| Stigmasterol                                                       | CCC(C=CC(C) C1CCC2C1(CCC3C2CC=C4C3(CCC(C4)O)C)C)C)C                               |
| Quinoline                                                         | C1=CC=C2C(=C1)C=C=CC=N2                                                        |
| Nootkatone                                                        | CC1CC(=O)C=C2C1(CC(CC2)C(=C)C)                                                 |
| Melatonin                                                        | CC(=O)NCC1C1CNC2=C1C=C=C2OC                                                    |
| 9(10H)-Anthracenone, 1,3,8-trihydroxy-6-methyl-                   | CC1=CC2=C(C(=C1)O)C(=O)C3=C=C2                                               |
| Isonipeotic acid, 1-methyl-4-phenyl-, ethyl ester                  | CCO(=O)C1CCN(CC1)C2=CC=CC=CC=CC=C2                                                |
| Formosanan-16-carboxylic acid, 19-methyl-2-oxo-, (19alpha)-       | CC1C2CN3CCC4(C3C2C(=CO1)C(=O)O)C(=O)O)C(=O)O                                    |
| Anethole <(E)->                                                   | CC=CC1=CC=C(C=C1)OC                                                         |
| Cannabinol                                                       | CCCCCC1CC1C=C2C=C1(=C1)OC                                                   |
| 1H-Pyrazole, 1,3-dimethyl-                                        | CC1=NN(C=C1)C                                                               |
| Fluoroacetamide                                                  | C(C(=O)N)F                                                                   |
| Picoline <meta->                                                  | CC1=CN=CC=C1                                                                |
| Benzenamine, N-phenyl-                                            | C1=CC=C(C=C1)NC2=CC=CC=C2                                                   |
| 2,6-Pyridinedicarboxylic acid, 4-(((4-amino-1-carboxy-4-oxobutyl)imino)ethylidene)-1,2,3,4-tetrahydro- | C1C(N=C(C=C1C=NC(CCC(=O)N)C(=O)O)C(=O)O)C(=O)O                                    |
| Eudesmol <beta->                                                 | CC12CCCC(=C)C1CC(CC2)C(C)C)O                                                |
| Cinnamaldehyde                                                   | C1=CC=C(C=C1)C=CC=O                                                          |
| 9,10-Anthracenedione                                              | C1=CC=C2C=C1C(=O)C                                                          |
| Piperonyl aldehyde                                                | C1OC2=C(O1)C=C(C=C2)C=O                                                      |
| Pelargonic acid                                                  | CCCCCCCCCC(=O)O                                                              |
| 2H-pyran-2-one, 5,6-dihydro-4-methyl-                            | CC1=CC(C=O)OCC1                                                             |
| Cholest-5-en-3-ol, (3beta)-                                      | CC(C)CCCC(C) C1CCCC2C1(CCC3C2CC=C4C3(CCC(C4)O)C)C                                |
| Caryophyllene alcohol                                            | CC1CC2C1CCCC2CC=C4C3(CCC(C4)O)C                                              |
| Eugenol <methoxy->                                                | COC1=CC=C(C=O)OCC=CC                                                          |
| Beta-D-Ribofuranose, 1-(6-amino-9H-purin-9-yl)-1-deoxy-           | C1=NC(=C2C(=N1)N(C=N2)C3=C3(C=C(O3)CO)O)O)N                                   |
| Hydrocinnamate <methyl->                                         | COC(=O)CCC1CC=CC=CC=C1                                                      |
| Vanillin acetate                                                 | CC(=O)OC1=C(C=C(C=C1)C=O)OC                                                  |
| Delta-Tocopherol                                                 | CC1=CC=CC2C1OC(CC2)C(CCC(C))C(CCC1CCCC(C))C                                  |
| Chamazulene                                                      | CCC1=CC2=C(C=CC2=C(C=C1)C)                                                 |
| Heptanoate <ethyl->                                              | CCCCCCCCC(=O)OCC                                                             |
| Androstane                                                       | CC12CCCC1C3CCCC4CCCC4C3CC2C                                                 |
| Lanost-8-en-3beta-ol                                              | CC(C)CCCC(C)C1CCCC2C1CCCC3=C2CCCC4CC3(CC(C4)(C)O)O)C(C                                 |
| N-amidinosarcosine                                               | CN(CC(=O)O)OJC(=N)[NH3+]                                                   |
| 2-Piperidine carboxylic acid                                     | C1CC[NH]2+[C=C1]C(=O)[O-]                                                   |
| Formate <octyl->                                                 | CCCCCCCCCOC=O                                                                |
| Isoquinoline, 6-amino-                                            | C1=CC2=C(C=CN=C2)C=C1N                                                     |
an intermediate node degree of 11.6 were acquired, representing the target genes. Target genes involved in CKD: DRD1, DRD2, IL2, F10, CASP3, CREB1, and AHSA1 are shown (Fig. 3C), whereas CVD have a gene cluster as ADH1C, ADRA2A, SLC6A2, LTA4H, MAOA, ADRB1, SCN5A, ADR1A, MMP8, PDE3A, DPEP1, AR, NFkBA, MMP13, and KCNMA1 (Fig. 3D). The target gene functional network of 42 potential target genes in Chandanasava ingredients is depicted (Fig. 4). The evaluation of the protein–ligand interaction with 82 nodes and 149 edges resulted in the appropriate network. The red diamond nodes indicate 45 Chandanasava bioactive components, the yellow circle nodes denote 7 target proteins of CKD, and 15 blue circles show CVD. Moreover, the 20 pink circle nodes exhibit common genes of CKD and CVD.

| Molecule name                                    | Canonical SMILES                        |
|-------------------------------------------------|----------------------------------------|
| Piperine                                        | C1CCN(CC1)C(=O)                         |
| Piperine                                        | C = CC = CC2 = CC3 = C(C = C2)OCO3      |
| Undecyl alcohol                                 | CCCCCCCCC                          |
| Elemol                                          | CC(=C)C1C(CCC1(C)C = C)(C)(C)O         |
| Caprylic acid ethyl ester                       | CCCCCCCCC(=O)OCC                    |
| 2-Ethyl-3,5-dimethylpyrazine                    | CCC1 = NC = C(N = C1)C               |
| (S)-1-Phenyl-2-propylamine                      | CC(CC1 = CC = CC = C1)N              |
| Nonane                                          | CCCCCCCCC                          |
| Jasmonene < (Z) >                               | CCC = CCC1 = C(CCC1 = O)C            |
| Anisyl alcohol < para- >                       | COC1 = CC = C(C = C1)CO              |
| Labdanol                                        | CC(C)OC(=O)C = CC1 = CC = CC = C1      |
| Elemicin                                        | COC1 = CC(=CC(=CC1OC)OC)CC = C       |
| Piperonal                                       | C1OC2 = C(O1)C = C(C = C2)C = O       |
| Beta-Tocopherol                                 | CCC1 = C(CN(C1 = O)C(=O)NCCC2 = CC = C(C = C2)S(=O)(=O)NC(=O)NC3CCC(C3)C |
| Cinnamic acid                                   | C1 = CC = C(C = C1)C = CC(=O)O       |
| 5-Hydroxy-7- 3 ‘,4’-trimethoxyflavanone          | COC1 = CC2 = C(C = C1) (=O)CC(=O2)    |
| Acetylphenylhydrazine                           | C3 = CC(=C(C = C3)OC)OC)O            |
| Phenylethyl alcohol                             | C1 = CC = C(C = C1)CO                |
| 5-Hydroxymethylfurfural                        | C1 = C(OC(=C1)C = O)CO              |
| 1-Tetradecanol                                  |CCCCCCCCCCCCCCCCO                  |
| 2,4-Di-tert-butylphenol                         | CC(C)(C1 = CC(=C(C = C1)O)(C)(C)C |
| 3-Eicosene, (E)-                               | CCCCCCCCCCCCCCCCC = CCC            |
| 1-Nonanol                                       | CCCCCCCCCCO                        |
| 1-Hexadecanol                                   | CCCCCCCCCCCCCCCCCCO                |
| 1,2-benzenedicarboxylic acid, dibutyl ester     | CCCCCO(=O)C1 = CC = CC = C1C(=O)OCCC |
| Myristic alcohol                                | CCCCCCCCCCCCCCO                  |
| 1-Tricosene                                     | CCCCCCCCCCCCCCCCCCCCCC = C        |
GO Enrichment Analysis

DAVID server was utilized to perform the GO enrichment analysis against the disease-associated target genes. As illustrated in Fig. 5, a $p$-value ($p < 0.05$) was used to demarcate the best 20 significantly enriched terms in the BP and MF criteria. These findings revealed that these CKD and CVD target genes are involved in the molecular regulation of various biological processes. Totally, 189 BP primarily were attributed to significant functions for positive regulation of cellular transcription in the formation of RNA polymerase II promoter that negatively regulates the drug cell proliferation, inflammatory response, ERK1 and ERK2 cascade positive regulation, transcription factor positive regulatory activity, nitric oxide biosynthetic process positive regulation, lipopolysaccharide-mediated signaling pathway, and response to hypoxia along with the many other genes involved in biological processes. Among the 33 MF, the respective protein targets were binding with enzyme, norepinephrine, epinephrine interleukin-1 receptor, dopamine, protease, cytokine activity, protein homodimerization activity, binding, and serine-type endopeptidase activity that are compiled for molecular functioning genes.

Fig. 2 Interaction network of Chandanasava ingredients with target diseases. Note: The red diamond nodes represent the 45 compounds, 7 yellow circle nodes represent the genes of chronic kidney diseases, 15 blue circle nodes represent the genes of cardiovascular diseases, 20 pink circle nodes represent the genes involved in both chronic kidney and cardiovascular diseases, and the remaining 93 green circle nodes represent the other related target corresponding genes of the compounds.
KEGG Pathway Enrichment Analysis

Potential targets of Chandanasava compounds correlated to disease, including CKD and CVD, were identified using the KEGG pathway module. KEGG pathway analyses of protein targets and pathway-target interaction networks were represented (Fig. 6). According to the study, the Chandanasava metabolites and target proteins network have 64 clusters and 243 edges. Appropriate p-values with <0.05 correspond to 35 pathways for 29 potentially target genes screened with BH. These results exhibited that the target genes were primarily involved in TNF signaling, cGMP-PKG signaling, Toll-like receptor signaling, MAPK signaling, Adrenergic signaling in cardiomyocytes,
Table 3  The interaction information of *Chandanasava* compounds with 42 target proteins of chronic kidney and cardiovascular diseases

| UniProt ID | Name of the target protein                                    | Gene symbol |
|------------|----------------------------------------------------------------|-------------|
| P22303     | Acetylcholinesterase                                           | ACHEx       |
| P00326     | Alcohol dehydrogenase 1C                                       | ADH1C       |
| P35348     | Alpha-1A adrenergic receptor                                   | ADRA1A      |
| P08913     | Alpha-2A adrenergic receptor                                   | ADRA2A      |
| P08588     | Beta-1 adrenergic receptor                                     | ADRB1       |
| P07550     | Beta-2 adrenergic receptor                                     | ADRB2       |
| O95433     | Activator of 90 kDa heat shock protein ATPase homolog 1        | AHSA1       |
| P15121     | Aldose reductase                                                | AKR1B1      |
| P10275     | Androgen receptor                                              | AR          |
| P42574     | Caspase-3                                                      | CASP3       |
| P13500     | C–C motif chemokine 2                                          | CCL2        |
| P16220     | Cyclic AMP-responsive element-binding protein 1                | CREB1       |
| P16444     | Beta-lactamase                                                 | DPEP1       |
| P21728     | Dopamine D1 receptor                                           | DRD1        |
| P14416     | D(2) dopamine receptor                                         | DRD2        |
| P03372     | Estrogen receptor                                              | ESR1        |
| P00742     | Coagulation factor Xa                                           | F10         |
| P13726     | Tissue factor                                                  | F3          |
| P01100     | Proto-oncogene c-Fos                                           | FOS         |
| P05362     | Intercellular adhesion molecule 1                              | ICAM1       |
| P01583     | Interleukin-1 alpha                                            | IL1A        |
| P01584     | Interleukin-1 beta                                             | IL1B        |
| P60568     | Interleukin-2                                                  | IL2         |
| P05231     | Interleukin-6                                                  | IL6         |
| P05412     | Transcription factor AP-1                                      | JUN         |
| Q12791     | Calcium-activated potassium channel subunit alpha 1            | KCNMA1      |
| P09960     | Leukotriene A-4 hydrolase                                      | LTA4H       |
| P21397     | Amine oxidase [flavin-containing] A                             | MAOA        |
| Q16539     | Mitogen-activated protein kinase 14                            | MAPK14      |
| P45452     | Collagenase 3                                                  | MMP13       |
| P22894     | Neutrophil collagenase                                          | MMP8        |
| P25963     | NF-kappa-B inhibitor alpha                                      | NFKB1A      |
| P29474     | Nitric oxide synthase, endothelial                             | NOS3        |
| P08235     | Mineralocorticoid receptor                                     | NR3C2       |
| Q14432     | CGMP-inhibited 3',5'-cyclic phosphodiesterase A                 | PDE3A       |
| P00749     | Urokinase-type plasminogen activator                            | PLAU        |
| P35354     | Prostaglandin G/H synthase 2                                   | PTGS2       |
| P00797     | Renin, renal                                                   | REN         |
| Q14524     | Sodium channel protein type 5 subunit alpha                     | SCN5A       |
| P23975     | Sodium-dependent noradrenaline transporter                     | SLC6A2      |
| O00206     | Toll-like receptor 4                                            | TLR4        |
| P01375     | Tumor necrosis factor                                          | TNF         |
cAMP signaling, NF-kappa B signaling, MAPK signaling, Renin secretion, T cell receptor signaling pathway, and so on. The target proteins involved in the TNF signaling pathway (TNF) were CCL2, FOS, JUN, NFkBIA, CREB1, CASP3, ICAM1, IL1B, IL6, MAPK14, PTGS2, and TNF; the cAMP signaling pathway interacted with FOS, JUN, NFkBIA, ADRB1, ADRB2, CREB1, DRD1, DRD2, and PDE3A; the protein targets involved in the cGMP-PKG signaling pathway such as ADRA1A, ADRA2A, ADRB1, ADRB2, CREB1, NOS3, PDE3A, and KCNMA1. The target proteins involved in Toll-like receptor signaling pathway were FOS, JUN, NFkBIA, IL1B, IL6, MAPK14, TLR4, and TNF; MAPK signaling pathway mainly interacted with the target proteins including FOS, JUN, CASP3, IL1A, IL1B, MAPK14, and TNF; NF-kappa B signaling pathway primarily consisted of NFkBIA, ICAM1, IL1B, PLAU, PTGS2, TLR4, and TNF; the protein targets in Adrenergic signaling in cardiomyocytes such as ADRA1A, ADRB1, ADRB2, CREB1, MAPK14, and SCN5A; T cell receptor signaling pathway of target proteins involved in FOS, JUN, CASP3, IL1A, IL1B, MAPK14, and TNF; Renin secretion target proteins consisted of ADRB1, ADRB2, CREB1, PDE3A, KCNMA1, and REN. NOD-like receptor signaling pathway mainly networked with CCL2, NFkBIA, IL1B, IL6, MAPK14, and TNF, the target proteins involved in Estrogen signaling pathways such as FOS, JUN, CREB1, ESRI, and NOS3. Multiple target genes could be found in a single pathway at any given time, and a single specific target gene can be found in various pathways. Analysis proves that a single pathway involving numerous protein targets is more significant than a single protein target interacting with multiple pathways. This is due to the fact that the impact of a single-target protein on the entire pathway may be minimal, whereas many target protein interactions can have a significant effect. The main target proteins are TNF, IL1B, IL6, JUN, FOS, NFkBIA, TLR4, MAPK14, CREB1, IL1A, CASP3, CCL2, IL2, and ICAM1. They were highly interacted/involved in these pathways and can be considered potential targets to treat various diseases. From the above-categorized results, the effective bioactive ingredients of polyherbal fermented medicine, Chandanasava, shows insight into potential target proteins in treating inflammation, chronic inflammation conditions,
such as kidney-related diseases, cardiac diseases, rheumatoid arthritis, type 2 diabetes, bacterial and viral infections, boosting immunity, and other diseases.

Identification of Hub Genes Associated with Targeted Diseases

Hub genes are the essential genes that control and express all associated sub-proteins to perform their molecular and biological functions whenever required [25, 38]. Three hub genes from the top-hit ranking are associated with the target diseased gene network. Such genes are tumor necrosis factor (TNF, PDB: 2E7A:A) in rank 1 and interleukin-6 (IL6)
Based on the DisGeNET and GeneCards databases search, 21 compounds have the best drug-likeness interacting properties against the three hub proteins. Such interactions are TNF (1 compound), IL6 (2 compounds), and PTGS2 (19 compounds). Therefore, these 21 compounds were further utilized for the molecular docking experiments with the three hub proteins.
Performance of Molecular Docking Studies

Piperine shows the best interaction with TNF and IL6 proteins (Fig. 8, Table 4). The binding score was confirmed as -7.08 for TNF. Ten hydrophobic and one hydrogen bonding interactions were found in Asn46 with a bonding distance of 2.55 Å. IL6 shows six hydrophobic, one hydrogen bonding interactions at Glu80 (2.54 Å), with a binding score of -5.47. The receptor protein PTGS2 against melatonin illustrates 11 hydrophobic and three hydrogen bonding interactions at Gly13 (2.62 Å) and Arg438 (3.12 Å, 2.50 Å) with...
Table 4  Representation of macromolecular interaction of selected disease-associated hub genes against the ingredients of Chandanasava

| Receptor          | Compounds | Binding Energy | Hydrogen bond interactions | Hydrophobic Interaction |
|-------------------|-----------|----------------|-----------------------------|-------------------------|
|                   |           |                | Interacted residues         |                         |
|                   |           |                | Distance (Å)                |                         |
| TNF (2E7A:A)      | Piperine  | -7.08          | Asn46                       | Asn19, Pro20, Gln21, Trp28, Leu29, Arg31, Arg32, Leu43, Arg44, Asp45 |
| IL6 (4CNI:C)      | Piperine  | -5.47          | Glu80                       | Lys57, Thr69, Thr76, Thr124, Thr125, Pro126 |
| PTGS2 (5F19:A)    | Melatonin | -10.5          | Gly13                       | Cys9, Gln10, Asn11, Arg12, Asp94, Thr98, Tyr99, Ala120, Leu121, Pro122, Lys437 |

ACE score indicates the solvation Delta Gs (Gibbs free energy)
the best binding score of -10.5. Among the two compounds, melatonin acts as a potential inhibitor for cardiovascular and chronic kidney disease. Overall, three compounds showed the best hydrogen bonding interaction and binding score out of 21 selected compounds.

**Discussion**

Ayurvedic pharmacopeia includes fermented traditional medicines (FTM) as described earlier for the effective polyherbal formulations that have been fermented by self-generated/ native microbes [14]. *Arishta* and *Asava* are traditionally valuable therapeutic Ayurvedic formulations due to their superior storage efficacy, improved pharmacological value, improved drug molecule extraction from herbs, and efficacy of drug transport in the body [10, 11]. The phytocompounds of *Chandanasava* showed activity against arthritis using molecular docking and dynamic analysis [39]. Previous reports suggested that *Chandanasava* scavenges free radicals by the presence of novel biotransformed phytochemicals [40].

In this study, 61 bioactive metabolites in polyherbal fermented traditional medicine *Chandanasava* were identified using GC–MS metabolomics analysis. Among those, 40 potential phytochemicals were screened by TCMSP based on ADME properties. These potential compounds were subjected to a network pharmacology approach to treat CKD and CVD. The network chart was developed by 42 potential target proteins mined from different databases. Among the 40 active ingredients, eugenol (<methoxy->, stigmasterol, isonipeotic acid, 1-methyl-4-phenyl-, ethyl ester, piperine, elemicin, 5-hydroxymethylfurfural and 1,2-benzendicarboxylic acid, and dibutyl ester were catalogued. Necessitated activities can corroborate the enhanced use in the treatment of CKD with cardiovascular complications. A single *Chandanasava* component may regulate numerous target proteins, while multiple components may regulate a single-target protein. Therefore, multi-target active compounds have more significant therapeutic potential and lesser adverse effects than single-target active compounds. This present assessment is the first report of a single component effect on target proteins in FTM network assessment. A total of 29 potentially target genes were primarily involved in TNF signaling, cGMP-PKG signaling, Toll-like receptor signaling, MAPK signaling, Adrenergic signaling in cardiomyocytes, cAMP signaling, NF-kappa B signaling, MAPK signaling, Renin secretion, and T cell receptor signaling pathway associated with the respective diseases based on GO and KEGG analysis. Based on a system pharmacology study, such bioactive components possess therapeutic potential on the inflammatory response, kidney, and cardiovascular disease–related health conditions. They can cure or prevent inflammatory diseases, kidney, and CVD complications.

Furthermore, the TNF signaling cascade can activate various intracellular signaling pathways such as transcription factor NF-kappa B, apoptosis, cell survival, inflammation, and immunity [41]. Overexpression of TNFR1 (major receptor) can activate p38MAPK in response to external or internal stimuli, boosts IL-6 production and triggers the inflammatory response [42]. TNFR2 agonistic reactions are expressed in cells only upon health limitations and cause therapeutic implications based on signaling cascades involving endothelial cells, CD4 and CD8 T lymphocytes, oligodendrocytes, neuron subtypes, microglia, cardiac myocytes, thymocytes, etc. [43]. Many genes are activated by TNFR1 signaling, which is principally regulated by the NF-kappa B pathway that subsequently activates the MAPK cascade resulting in necroptosis or apoptosis and necroptosis [41]. On the other
hand, TNFR2 signaling leads to survival via activating the NF-kappa B pathway, including the PI3K-dependent NF-kappa B pathway and the JNK pathway leading to cell death [44]. TNF signaling cascade regulates the expression of genes such as NFKBIA, CREB1, CASP3, ICAM1, IL1B, IL6, MAPK14, PTGS2, and TNF in many inflammatory diseases, including atherosclerosis, diabetics, and chronic kidney diseases affecting the inflammatory cascades [45]. Furthermore, danshen-shanzha decoction can interact and result in modulation of 41 potential targets and 16 signaling pathways correlated with metabolizing lipids that prevent endothelial dysfunction promoting cardiovascular protection against atherosclerosis [46]. MAPK14 (mitogen-activated protein kinase 14) represents the pivotal signal transducing component in proinflammatory cytokines, chemokines, and adhesion factors responsible for signaling from the cell membrane to the nucleus and responsible for intracellular responses and atherosclerosis [47].

The TCM Yangxinshi tablet was proven efficient for abating cardiovascular complications and enhancing immune power during heart failure [48]. Similarly, QiShenYiQi, Shenmai and Xuesaitong have been used to treat myocardial infarction and myocardial ischemia by promoting anti-apoptosis, anti-inflammation, antioxidant, and angiogenesis using animal models [49–51]. Furthermore, Buchang Naoxintong, Fufang Danshen formula have been traditionally used to treat CVD and stroke by regulating the corresponding inflammatory markers using literature and multi-compound target disease network [52]. In Indian traditional knowledge, *Arjunarishta* is cardiotonic; it rejuvenates and boosts the cardiovascular system by accelerating the blood flow in coronary arteries and controlling blood cholesterol levels and myocardial protection against ischemic damage [11]. There is substantial experimental evidence that it is beneficial in preventing cardiovascular disease alone or in combination with statins [53, 54]. *Dashamoolarishta* exhibited several therapeutics for anti-inflammation, analgesic, and anti-platelet properties effects by the animal model compared with standard drug aspirin [55]. Chronic kidney disease is thought to be caused by an immune response. It has a strong correlation with blood coagulation. MAPK14, NFKB1, cAMP, TLR4, TNF, IL6, TCR (CD28), and JUN are closely associated with inflammation, ECM, hypoxia, angiogenesis, and immune response. Previous studies reported that the TCM NE-THCQ exhibited anti-renal fibrosis activity by suppressing ECM deposition, decreasing inflammation, and regulating hypoxia through mechanistic signaling pathways [56]. Bu-shen-Huo-xue formula, a similar TCM, has been used to regulate coagulation and fibrinolytic balance to treat chronic kidney disease expression of the inflammatory marker using multi-target multi-components system pharmacology study [57]. A previous report demonstrated that *Astragalus* has medicinal properties in patients with diabetes, improving renal function [58]. Among the 29 target proteins, 13 (TNF, IL1B, IL6, JUN, FOS, NFKB1A, TLR4, MAPK14, CREB1, IL1A, CASP3, CCL2, IL2, and ICAM1) were highly involved in various pathways. The predicted PPINs were functionally validated based on gene ontology terms, experimentally determined interaction score (edges), and bonding interactions [59, 60]. Moreover, several recent experimental and molecular dynamic simulation studies were conducted for piperine and melatonin among selected hub proteins (TNF, IL6, and PTGS2), which show good interaction with each other [61–63]. Thus, the findings are functionally validated and associated with previous research.

The potential target proteins stimulate overexpression of cytokine levels, apoptosis, immune response, and inflammatory markers leading to chronic inflammation-associated diseases such as type 2 diabetes, CVD, chronic kidney failure, rheumatoid arthritis, insulin resistance, and cancer. Based on the analysis by network pharmacology, those bioactive ingredients regulate or alleviate signal pathways that have therapeutic effects on inflammation and kidney and cardiac diseases–related symptoms. Such compounds can treat or
alleviate the symptoms of inflammation and kidney and cardiac diseases through the role of potentially interacted target proteins in metabolic pathways caused by the occurrence of diseases. The molecular docking analysis demonstrates that the compounds piperine and melatonin act as potent inhibitors for CKD and CVD. Furthermore, the compounds piperine and melatonin also exhibited antibiofilm activity, indicating evidence for arresting urinary tract infections that can complicate the CKD incidence and aggravation. Similarly, docking piperine with cyclin-dependent kinases is effective against oral cancer [37]. Moreover, the small molecule has been proven effective in controlling the COVID-19 pandemic [64]. Furthermore, melatonin has been confirmed for renal management during chronotherapy by targeting the renin-angiotensin system (RAS) through intra-renal targeting [65]. Hence, the small molecule interaction with the protein targets is affirmative of protective roles in chronic kidney disease, thereby could be indirectly attributed to managing cardiovascular implications through signaling cascades. However, further protein interaction crosstalk will effectively affirm the Chandanasava effects in infectious disease management. Hence, chronic kidney disease and cardiovascular complications in the perspectives of the FTM are addressed. Thus, the conceptualization of disease Vs disorder management for efficient clinical outcomes and mortality mitigation are attributed to FTM.

Conclusion

Bioactive potentials of phytomedicinal herbs such as Chandanasava in the arena of FTM are constituted of multiple active components. Still, the functional properties of disease management have yet to be revealed. Mechanisms of action for combating the versatile arresting modes require more molecular dissection modalities with interdisciplinary research. This analysis demonstrates that the identified compounds, piperine and melatonin from Chandanasava, have protective potentials to inhibit biofilm formation of infectious diseases involved in CKD associated with CVD as affirmative from comprehensive network analysis coupled with docking perspectives. The study indicates the scope of validating the claims of the traditional Ayurvedic medicines using the modern tools of biology, particularly FTM, which are least understood. Such approaches could yield new avenues and drug molecules. The study also highlighted that the multi-target active compounds have better therapeutic and low adverse effects than the single-target active compounds. It also demonstrates that a single phytochemical compound has the potential to interact with multiple cellular targets and is likely to be a therapeutic measure for simultaneously treating multiple disorders or syndromes. However, further experimental validation, such as in vitro and in vivo analysis, is warranted for better understanding leading to drug development.

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Author Contribution A.V is the major contributor to this manuscript. A.V conducted the analysis, drafted the first version of the manuscript, and finalized the manuscript. A.V, R.P, and T.S collected the data and processed the graph and the table in the manuscript. Y.M and S.S conceptualized and supervised the investigation, as well as systematically evaluated the findings. The final manuscript was read and approved by all contributors.
Data Availability  All datasets generated for this study are included in the article/Supplementary Material.

Declarations

Ethics Approval  Not applicable.

Consent to Participate  Not applicable.

Consent for Publication  Not applicable.

Conflict of Interest  The authors declare no competing interests.

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