Modular characteristics and the mechanism of Chinese medicine’s treatment of gastric cancer: a data mining and pharmacology-based identification

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Background: Traditional Chinese medicine (TCM) is increasingly extensively being applied as a complementary and alternative therapy for gastric cancer (GC); however, there is a lack of large-scale evidence-based deep learning for the guidance of its clinical prescription.

Methods: The combinational search terms of “Gastric cancer and/or gastric malignancy” and “Traditional Chinese Medicine” were used to retrieve clinical study-based herbal prescriptions from public database over the past 3 decades [1990–2020]. Association rules mining (ARM) was used to analyze the prescription patterns of the herbs extracted from the eligible studies. Deep machine learning and computational prediction were conducted to explore candidate prescriptions with general applicability for GC. The action mechanism of the preferred prescription was investigated through network pharmacology, and further validated via in vivo and in vitro experiments.

Results: A total of 194 clinical study-based herbal prescriptions with good efficacy for GC were collected. TCM with focus on invigorating the Spleen and tonifying the vital-Qi is a promising adjuvant therapy for GC. The preferred prescription is composed of Atractylodis Macrocephalae Rhizoma, Astragali Radix, Pinelliae Rhizoma, Citri Reticulatae Pericarpium, Herba Hedyotidis Diffusae, Crataegi Fructus, and so on. We screened 74 bioactive compounds and 2,128 predictive targets of the preferred prescription from public databases. Eventually, 135 GC-related genes were identified as the targets of the preferred prescription. The compound-target network revealed that the crucial substances in the preferred prescription are quercetin, kaempferol, baicalein, and nobiletin. Experimentally, the preferred prescription was validated to modulate GC cell survival and inhibit tumor progression mainly via the hTERT/MDM2-p53 signaling pathway in vivo and in vitro.

Conclusions: TCM aimed at invigorating the Spleen and tonifying the vital-Qi is a promising adjuvant therapy for GC, which offers a guidance for worldwide use of TCM in the treatment of GC.

Keywords: Traditional Chinese medicine (TCM); data mining; machine learning; gastric cancer (GC); network pharmacology

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Introduction

Globally, gastric cancer (GC) is the fourth most common cancer and the second leading cause of cancer deaths (1). Current approaches to GC management largely consist of endoscopic detection followed by gastrectomy and chemotherapy (CT) or chemo-radiotherapy (CRT); however, the available treatments have adverse side effects and are associated with high recurrence rates (2). Therefore, there is a need to address the current limitations of the various therapeutic strategies to facilitate possible clinical applications.

With the development of personalized and complementary medicine, multi-compound and multi-targeting traditional Chinese medicine (TCM) has been shown to be clinically effective in treating GC (3,4). However, due to the lack of large-scale evidence-based medicine, the extensive application of TCM remains inhibited. The use of TCM as an adjuvant therapy is greatly subjective because understanding on GC treatment varies among physicians in terms of etiology, syndrome differentiation, and medicinal prescriptions. Generally, the principal theory of Chinese traditional medicine for GC is invigorating the Spleen and tonifying the vital-Qi, and eliminating blood stasis and removing toxins. The largely unknown mechanism of these empirical prescriptions is another limiting factor for the use of TCM. Therefore, it is important and innovative to screen clinical prescriptions with good efficacy, based on which the obtainment of a basic prescription with general applicability for treating GC could be achieved by machine learning. What's more, the elucidation of prescription patterns by data mining may promote both clinical application and basic researches on herbal pairs. To the best of our knowledge, another study with such an aim has not been previously reported.

In this study, we proposed a method of combining data mining and network pharmacology to systematically elucidate the prescription patterns of TCM, and unravel the modular functions and potential action mechanisms of TCM for treating GC. Additionally, the effects of the machine learning-based preferred prescription were validated in vivo and in vitro. We present the following article in accordance with the ARRIVE reporting checklist (available at https://dx.doi.org/10.21037/atm-21-6301).

Methods

Big data mining and machine learning

Source of literature, inclusion and exclusion criteria

All literature was obtained from the China National Knowledge Infrastructure (CNKI) database, which is the world’s largest Chinese knowledge portal website. The sources of the literature included the Academic Journals Full-text Database, Doctoral Dissertations Full-text Database, and Masters’. These Full-text Database (01/1990–12/2020). The combinational search terms were “Gastric cancer and/or gastric malignancy” and “Traditional Chinese Medicine”. Literature with the following criteria were included: (I) relevant to clinical research on using TCM in treating GC; (II) containing randomized controlled trial (RCT) as the study design; (III) containing prescriptions with complete and specific names of Chinese herbs; (IV) studies must have reported one or more of the following efficacy endpoints: progression-free survival (PFS), overall survival (OS), objective response rate (ORR), or adverse events (AEs) (5). The exclusion criteria were as follows: (I) duplicate publications reporting the same group of participants; (II) non-clinical studies including experimental research on cell lines, xenografts or animal models, or theoretical studies; (III) where TCM and western medicines were integrated as a therapeutic regimen; (IV) non-oral administrations including injection and nasogastric tube nutrition; (V) non-decoction dosage types including Chinese patent medicines and TCM for external use; (VI) use of prescriptions composed of an unspecified or single herb.

Data extraction

Firstly, the names of the prescriptions (ancient prescriptions or recombinant personalized prescriptions) and their constituent herbs were extracted from the eligible literature. Secondly, we referred to the Chinese Pharmacopoeia (2020 Edition) Volume I to standardize the names of each herb (6). Furthermore, the basic information on each herb was extracted from the Chinese Pharmacopoeia, including its Latin name, property, taste, and meridian tropism. The 5 properties of TCM herbs include cold, hot, warm, cool, and neutral. The 5 tastes of TCM herbs include sour, bitter, sweet, pungent, and salty. The various combinations of property and taste determine the herbs’ specific attributes, which can influence the Yin and Yang of the body. For example, herbs with warm and hot properties are used to invigorate the Yang in patients with heat-deficiency disorders. Likewise, sour, bitter, and salty tastes are related to Yin, whereas pungent and sweet pertain to Yang. The meridian serves as the pathway for the transportation of Qi and Blood throughout the body, and its tropism represents the selective therapeutic effects of a medicinal herb on
Based on their attributes (14). In our study, k-means cluster analysis was performed to identify the preferred regroups of the most frequently used herbs from the prescriptions used in various studies, ARM, an in-silico screening process, was applied. In this scheme, the dataset and the association rules are defined as follows: an association rule has the form left hand side (LHS) ⇒ right hand side (RHS), where LHS and RHS are sets of items, with the likely occurrence of the RHS whenever the LHS set occurs (9). The Apriori algorithm was used to extract the significant associations from all possible combinations of the items from the main dataset (10). There are 3 evaluation metrics which are critical in describing the power and significance of the rules generated by ARM (11). Support is the frequency of the rule occurrence in the total dataset, measuring whether an association between the LHS and the RHS happens by chance. Confidence is the frequency of rule occurrence in the cases of the dataset fulfilling the LHS of the rule, thus, representing the reliability of the association. Lift is the ratio of observed support to the expected support when the LHS and the RHS are independent, indicating the dependency of the occurrences of the 2 items when its value is larger than 1 (12). To establish a proper threshold, we detected the central tendency of the association rules to be more obvious at the support of 0.1 and confidence of 0.6 in the correlation analysis of herbal combination patterns. Then, the herbs were categorized in Microsoft Excel 2010 (Microsoft Corp., Redmond, WA, USA) according to their properties, tastes, meridian tropisms, and functions. The software platform IBM SPSS Modeler 18.1 (IBM Corp., Armonk, NY, USA) was used to analyze the categorization-based frequency and the correlations of the prescription patterns and to generate a visual network diagram.

**Cluster analysis**

Clustering is central to many data-driven bioinformatics research and serves a powerful computational method. Deep learning can be effective means to transform mappings from a high-dimensional data space into a lower-dimensional feature space, leading to improved clustering results (13). In this study, we used IBM SPSS Modeler software platform to perform deep learning-based cluster analysis to identify the preferred regroups of the most frequently used herbs based on their attributes (14). In our study, k-means cluster analysis was considered since the variables were quantitative at the interval or ratio level rather than being binary or counts. To avoid unreliable results through omitted variable bias, we included all the attributes, including the 5 properties, 5 tastes, and meridian tropism, and investigated the therapeutic preferences of the candidate clusters. To assess the reliability of a given solution, we compared the results from analyses with different permutations of the initial center values to ensure an appropriate number of clusters.

**Mechanism investigation of the candidate formulae by network pharmacology**

**Compounds library construction and active components screening of the candidate formulae**

To build a compound library of the core herbs for GC, we extracted all the compounds of the candidate formulae from the Traditional Chinese Medicines for Systems Pharmacology Database and Analysis Platform (TCMSP; http://lsp.nwu.edu.cn/index.php), Traditional Chinese Medicines Integrated Database (TCMID; http://bionet.ncpsb.org/batman-tcm/), Bioinformatics Analysis Tool for Molecular Mechanism of Traditional Chinese Medicine (BATMAN-TCM; http://bionet.ncpsb.org/batman-tcm/), and wide-scale literature mining (15,16). To optimize the use of the high cost and time-consuming biological experiments and clinical research, absorption, distribution, metabolism, and excretion (ADME) evaluations are critical procedures for active components screening (17). In this study, oral bioavailability (OB) ≥30% and drug-likeness (DL) ≥0.18 were set as the threshold; however, compounds that did not meet these inclusion criteria but were supported by the literature were retained.

**Therapeutic targets prediction of the candidate formulae**

Computational predictions of bioactive molecule targets based on similarity with known ligands are powerful in narrowing down the number of potential targets and the rationalization of possible side effects of the known molecules (18). The prediction algorithms of the ligand-based strategies include systematic drug targeting (SysDT) (19) and weighted ensemble similarity (WES) models (20). The SysDT model was developed based on random forest (RF) and support vector machine (SVM), which performed impressively on systematic predictions for drug-target associations and interactions involving enzymes,
ion channels, nuclear receptors, and G-protein coupled receptors (19). In the WES model, the standardized ensemble similarities (Z score) by Bayesian network are utilized and the targets are predicted using the multivariate kernel approach (21). In our study, the predictive therapeutic targets of the candidate formulae were obtained from web tools including Search Tool for Interacting Chemicals (STITCH; http://stitch.embl.de/), similarity ensemble approach (SEA; http://sea.bkslab.org/) and SwissTargetPrediction (www.swisstargetprediction.ch) (22-24). Targets with RF ≥0.7, SVM ≥0.8, or Z score ≥7 were considered for further analysis and standardized to corresponding genes for homo sapiens through the UniProt database (https://www.uniprot.org/uploadlists/).

To evaluate the performances of the candidate formulae in treating GC, we mapped the predictive therapeutic targets to the GC-related genes/proteins, which were comprehensively collected from online databases including MalaCards (https://www.malacards.org/), Online Mendelian Inheritance in Man (OMIM; https://omim.org/) and DisGeNET v7.0 (https://www.disgenet.org/home/) (25,26). We visualized the results and generated an additional protein-protein interaction (PPI) network using Metascape (https://metascape.org/gp/index.html).

Construction and topological analysis of the compound-target network of the preferred prescription
Cytoscape v3.7.2 (https://cytoscape.org/) was used to construct a compound-target (C-T) network of the preferred prescription, and to analyze its degree, a key topological parameter for evaluation (15). In the C-T network, compounds sharing interactions with GC-related genes were determined as components that were beneficial for GC. Moreover, we considered the targets (compounds) with degree values equal to or above the mean value to be the predominant therapeutic targets (crucial substances).

Gene Ontology (GO) and pathway enrichment analysis of the preferred prescription
GO analysis, Kyoto Encyclopedia of Genes and Genomes (KEGG), and Reactome pathway enrichment of the preferred prescription were carried out using the Database for Annotation, Visualization, and Integrated Discovery system v6.8 (DAVID; https://david.ncifcrf.gov/) (27). We also used ClueGO, a Cytoscape v3.7.2 plug-in to identify the interactions among the various signaling pathways by generating a functionally grouped network (28,29). Based on the mechanism of GC, we further constructed a multi-regulation map of KEGG pathways of the crucial components in the core herbs.

Moreover, the modular functional characteristics of TCM in GC treatment were demonstrated in PPI networks, which visualized the interactions among significant targets that could be regulated by the crucial components in the preferred prescription. The PPI networks were generated by the GeneMANIA web site (http://genemania.org/) which offers a fast prediction on the functions of the given gene sets through the application of a guilt-by-association approach (30).

Molecular docking
The three-dimensional (3D) structures of the predominant targets of the preferred prescription were collected from protein data bank (PDB; http://www.rcsb.org). AutoDock Tools 1.5.6 software (https://autodock.scripps.edu/) was used to remove the water molecules, isolate proteins, add nonpolar hydrogen, and calculate Gasteiger charges for the structure (31). The preprocessed structures were saved as PDB with partial changes and AutoDock 4 atom types (PDBQT) files. The PubChem database (https://pubchem.ncbi.nlm.nih.gov/) was applied to download the two-dimensional (2D) structures of the crucial substances of the preferred prescription. The 2D structure was processed and transformed into PDB format via Open Babel (32), and then saved in PDBQT format as docking ligands in AutoDock Tools 1.5.6 software. The target proteins were used as receptors while the substances were used as ligands. The active site of molecular docking was determined by the complex of ligand and target protein. Autodock Vina 1.1.2 (https://vina.scripps.edu/) was used to dock small molecules with their target proteins. The conformation with the best affinity was selected as the final docking conformation and visualized in Pymol 2.5 (https://pymol.org/2/).

Experimental validation
Preparation of the preferred prescription and components identification
Crude TCM herbs [dried roots of Atractylodes macrocephala Koidz. 12 g, dried roots of Astragalus membranaceus (Fisch.) Bge. 30 g, dried mature pericarp of Citrus reticulata Blanco 10 g, dried tuber of Pinellia ternate (Thunb.) Breit. 9 g, dried root of Aucklandia lappa Decne. 6 g, dried mature fruits of Amomum villosum Lour. 3 g, dried immature fruits of Citrus aurantium L. 10 g, dried gizzard lining of Gallus gallus domesticus Brisson 10 g, dried mature fruits of
of *Crataegus pinnatifida* Bge. 12 g, dried mature fruits of *Hordeum vulgare* L. 15 g, *Radix Actinidiae Chinensis* 15 g, and *Herba Hedysotidis Diffusa* 15 g] were provided by Sanyue Chinese Traditional Medicine Co. (Nantong, China). All the herbs were soaked for 30 min in 1,800 mL double-distilled water and then boiled at minimum temperature for 30 min before being reflushed and extracted. The boiling process was repeated with 1,800 mL double-distilled water for 30 min. Then, 2 parts of the extracted solutions were mixed and vaporized to 60 mL. The decoction was finally concentrated to 1 g/mL and stored at −20 °C after being sterilized and filtered through a 0.22 μm filter. The extracts of the preferred prescription were detected and analyzed using high-performance liquid chromatography diode array detection (HPLC-DAD) (detailed information shown in the Supplementary materials).

**In vitro**, to determine the decoction dose, the half-maximal inhibitory concentration (IC50) of different GC cell lines were assessed by 3-(4,5-Dimethyl-2-thiazolyl)-2,5-diphenyltetrazolium bromide (MTT) assay (detailed information shown in the Supplementary materials), and the dose range of 2, 4, 8 mg/mL was selected.

**Cell apoptosis and cell cycle analyses**

Human GC cell lines AGS, HGC27, MKN28, and SGC7901 were purchased from the Cell Bank of the Chinese Academy of Sciences (Shanghai, China). All cell lines were kept in a humidified atmosphere of 5% CO2 at 37 °C. For apoptosis analysis, the cells were measured using Annexin V-FTTC/PI apoptosis detection kit (Keygen Biotech Co., Nanjing, China) by flow cytometry [Becton, Dickinson, and Co. (BD) Biosciences, Franklin Lakes, NJ, USA] according to the manufacturer’s instructions. Cell cycle distributions were determined using a cell cycle and apoptosis analysis kit (Beyotime Biotech Co., Shanghai, China) by flow cytometry (BD Biosciences).

**Wound-healing assay**

Cells (1,000x10^5 cells/well) were seeded into 6-well plates for 24 h, and scraped with a sterile pipette tip when 80% of the cells were adherent to the walls. Cells were treated with various concentrations of the preferred prescription after removing debris by phosphate-buffered saline (PBS). The scratch area was observed by microscopy at 0, 12, 24, and 48 h, respectively.

**Invasion assay**

The upper surface of the Transwell inserts (8 μm pore size, Merck & Millipore, Darmstadt, Germany) were coated with Matrigel (100 µL, diluted 1:29 with PBS) (Corning, Corning, NY, USA) before serum-free medium containing 2x10^5 cells were loaded. The lower chamber included 500 µL media containing 10% fasting blood sugar (FBS) and various concentrations of the preferred prescription. After 48 h, the chambers were removed, and nonpenetrative cells were washed from the top chamber with PBS. The invaded cells were fixed with 95% ethanol and stained with crystal violet. Image J (https://imagej.nih.gov/ij/) was applied to count the number of cells in images randomly taken under a microscope.

**Western blot assay**

Protein lysates were separated using sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to polyvinylidene difluoride (PVDF) membranes. Membranes were blocked with 5% bovine serum albumin (BSA) for 1 h and incubated with primary antibodies at 4 °C overnight. The primary antibodies included β-actin, Bax, Bel2, N-cadherin, Snail, Slug, hTERT, MDM2, p53, p21, cyclinE, and CDK2 [all antibodies were purchased from Cell Signaling Technology (CST) Danvers, MA, USA]. The secondary goat anti-rabbit horseradish peroxidase-conjugated antibody (ZSGB-BIO, Beijing, China) was incubated at room temperature for 1 h. Signals were examined using the Image Lab system, version 5.1 (Bio-Rad, Hercules, CA, USA).

**In vivo study**

Male BALB/c athymic nude mice (4–6 weeks old, 18–20 g) obtained from Charles River Co. (Beijing, China) were housed in a specific pathogen-free (SPF) environment. An appropriate amount of the preferred prescription extracts was collected and prepared into the 0.735 g/mL solution with distilled water, and used for the intragastric administration of the experimental animals. The MKN28 cells were collected and cultured in the logarithmic growth phase, and the density was adjusted to 5x10^4/mL. Each mouse was inoculated with 0.2 mL of cell suspension in the right armpit after disinfection. After 10 days, the diameter of the induration reached 3–7 mm, suggesting the establishment of a successful model. The 20 nude mice were divided into 4 groups (n=5 each) as follows: (I) model group with transplanted tumors given the gavage of distilled water; (II) 5-fluorouracil (5-FU) group with transplanted tumors given the intraperitoneal injection at a dose of 20 mg/kg body weight (BW) every 3 days; (III) preferred...
prescription group with transplanted tumors given the gavage of decoction at a dose of 14.7 g/kg BW every day for 14 days; (IV) the 5-FU+ preferred prescription group with transplanted tumors given the intraperitoneal injection at a dose of 20 mg/kg BW every 3 days and gavage of decoction at a dose of 14.7 g/kg BW every day for 14 days. To calculate the volume of the tumors, the dimension was measured by length (L) and width (W) using a caliper every 3 days. Mice were sacrificed by cervical dislocation, and the tumors were excised and weighed.

Ethical statement
This study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). Animal experiment was performed under a project license (No. 2021DW-35-01) granted by the Animal Ethics Committee of Affiliated Hospital of Nanjing University of Chinese Medicine (Nanjing, China), in compliance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. A protocol was prepared before the study without registration.

Statistical analysis
The data were described as means ± standard error of the mean (SEM). Statistical significance was determined using one-way analysis of variance (ANOVA, comparison between multiple groups) and Tukey multiple comparison processing (comparison between the two groups), with a P value <0.05 indicating statistical significance. All experiments were repeated at least three times under the same conditions. The statistical analyses were performed using GraphPad Prism software (GraphPad Software, La Jolla, CA, USA).

Results

Screening of eligible literature, clinical study-based prescriptions, core herbs, and frequency distributions according to herbal attributes and principal functional categorizations

The framework of this study can be summarized as follows: (I) screening of clinical study based TCM prescriptions for GC treatment; (II) data mining of the treatment principles, prescription patterns, and generation of candidate formulae by deep machine learning; (III) prediction of the action mechanism of the preferred prescription by network pharmacology; (IV) validation of the antitumor effects of the preferred prescription by experiments in vivo and in vitro (Figure 1A).

A total of 194 eligible prescriptions and 148 herbs with standardized names were screened from clinical studies spanning from January 1990 to December 2020. The screening process is summarized as a PRISMA flow diagram (33) (Figure 1B). The total cumulative occurrences of the 148 herbs in 194 prescriptions were 2,103 times. Herbs with over 20 times frequency of occurrence were selected as predominant ones used in clinic. The top 24 core herbs and their functional categorizations are listed in Table S1. Descriptive statistics of herbal attributes are shown in Figure 2A. In terms of the 5 properties; herbs with warm property were the most frequently prescribed. With regards to the 5 tastes, herbs with bitter (44.59%), pungent (39.86%), and sweet (36.49%) tastes ranked the top 3 in clinical application. In terms of meridian tropism, herbs with a propensity for the Liver (LR) (43.24%), Stomach (ST) (39.19%), and Spleen (SP) meridians were the most frequently used. The top 3 principal functions of the core herbs are demonstrated in Figure 2B. In summary, the treatment principle of TCM in GC is mainly invigorating the Spleen and tonifying the vital-Qi.

To facilitate better application of the core herbs in clinic, we summarized their clinical indications (Table S2). Particularly, herbs with the 3 major functions that embody TCM treatment principles for GC are listed in Table 1.

Frequently prescribed herbal combination patterns by ARM and novel candidate formula prediction by cluster analysis

The ARM method was applied to analyze the combination patterns of the 194 prescriptions. Guided by the theory of synergy and attenuation in TCM, couplet herbs are 2 herbs administered together to enhance therapeutic effects or reduce toxicity. With a threshold of minimum support of 0.1 and confidence of 0.6, the prescribed pairs of couplet herbs with the top 3 confidence included; Atractylodis Macrocephalae Rhizoma (Bai Zhu) paired with Dioscoreae Rhizoma (Shan Yao; 92.31%), Atractylodis Macrocephalae Rhizoma paired with Aucklandiae Radix (Mu Xiang; 90%), and Atractylodis Macrocephalae Rhizoma paired with Codonopsis Radix (Dang Shen; 88.57%) (Table 2). Triplet herbs are a combination of 3 herbs, which interact with each other and are usually contained in a decoction or used as an independent decoction. Based on the established threshold above, the triplet combinations of herbs with the top 3 confidence included Poria (Fu Ling)-Aucklandiae Radix-Glycyrrhizae...
Figure 1 The technical roadmap of the current study. (A) The framework of the current study is summarized as data mining and machine learning combined with network pharmacology and experimental validation. (B) Flow chart of literature mining. A total of 1,333 records were retrieved, and 194 prescriptions were extracted.
A

Neutral
24 (18.32%)
Cool
6 (4.58%)
Warm
50 (38.17%)
Hot
5 (3.82%)

Salty
Bitter
Pungent
Sour
Sweet

Cold
46 (35.11%)

Sour
8 (4.04%)

Bitter
66 (33.33%)

Pungent
59 (29.81%)

Sweet
54 (27.27%)

Cool
48 (35.11%)

Warm
50 (38.17%)

Hot
5 (3.82%)

B

Tonifying and replenishing medicinal

Ligustri lucidi fructus
24 (3.94%)
Dioscoreae rhizoma
26 (4.27%)
Paeoniae radix alba
29 (4.76%)
Pseudostellariae radix
29 (4.76%)
Angelicae sinensis radix
48 (7.86%)
Glycyrrhizae radix
96 (15.75%)

Atractylodis macrocephalae rhizoma
145 (23.81%)

Codonopsis radix
105 (17.24%)

Astragali radix
107 (17.57%)

C

C

D
Radix (Gan Cao) (100%), Atractylodis Macrocephalae Rhizoma-Aucklandiae Radix-Codonopsis Radix (100%), and Atractylodis Macrocephalae Rhizoma-Galli Gigeriae Endothelium Corneum (Ji Nei Jin)-Codonopsis Radix (100%) (Table S3). A network diagram was generated to visualize the association rules among the core herbs (Figure 2C).

Additionally, the core herbs in the 194 prescriptions were regrouped into 4 clusters by machine learning. The cluster analysis result was presented in a 2D scatter diagram (Figure 2D). Cluster 1 (candidate formula 1, CF 1) included; Atractylodis Macrocephalae Rhizoma, Astragali Radix (Huang Qi), Pinelliae Rhizoma (Zhi Ban Xia), Citri Reticulatae Pericarpium (Chen Pi), Herba Hedyotidis (Bai Hua She She Cao), Galli Gigeriae Endothelium Corneum, Aucklandiae Radix, Amomi Fructus (Sha Ren), Hordei Fructus Germinatus (Mai Ya), Aurantii Fructus (Zhi Ke), Radix Actinidiae Chinensis (Mi Hou Tao Gen), Crataegi Fructus (Shan Zha); cluster 2 (candidate formula 2, CF 2) included; Paeoniae Radix Alba (Bai Shao), Scutellariae Barbatae Herba (Ban Zhi Lian), Salviae Miltiorrhizae Radix et Rhizoma (Dan Shen), Ligustri Lucidi Fructus (Nu Zhen Zi); cluster 3 (candidate formula 3, CF 3) included; Poria, Codonopsis Radix, Glycyrrhizae Radix, Cocos Semen (Yi Yi Ren), Pseudostellariae Radix (Tai Zi Shen), Dioscoreae Rhizoma; and cluster 4 (candidate formula 4, CF 4) included; Angelicae Sinensis Radix (Dang Gui), Curcumae Rhizoma (E Zhu). For a clearer understanding of the functions of the candidate formulae, the herbal attributes preferences of the 4 candidate formulae are shown in a distribution histogram (Figure 2E-2H). The CF 1 and CF 2 were composed of more herbs with warm and cold
properties respectively, while the property of CF 3 appeared to be milder. With regards to the 5 tastes, the majority of the herbs in the CF 1 possessed sour and pungent tastes, while most herbs with sweet tastes were clustered in CF 3. Meridian tropism represents the selective therapeutic effects of a Chinese herb on a certain region of the human body (7). The CF 1, CF 2, and CF 3 prescribed more herbs belonging to SP (ST), LR, and KI meridians, respectively. Specifically, the CF 1 and CF 3 clusters were distinguished for tonifying Qi and invigorating the Spleen, and regulating Qi and resolving dampness, which meant both of them could increase appetite, alleviate lassitude, fullness sensation in the upper abdomen, and loose stools, as well as help GC patients feel less depressive. While the CF 2 and CF 4 seemed to play a significant role in tonifying the Blood and promoting blood circulation, which indicated they are more applicable for GC patients with symptoms like pale complexion, dizziness, insomnia, distending pain of the hypochondrium, and so on.

### Active components library construction and therapeutic targets prediction of the candidate formulae

Complied with OB ≥30% and DL ≥0.18, 305 compounds of the 24 core herbs were screened out as bioactive components (Table S4). The numbers of active components

| Herbal nature | Principal functional categorizations | Number of prescriptions using the herbs | Frequency of use (%) | Syndromes | Key signs & symptoms | Treatment principles | Representative herbs |
|---------------|-------------------------------------|----------------------------------------|----------------------|-----------|----------------------|---------------------|---------------------|
| Sweet, warm   | Qi-tonifying                        | 182                                    | 93.81                | Poor appetite, dislike to talk, lassitude, weak limbs, boredom, loose stools, heavy descending sensation in abdominal cavity, prolapse of rectum. | Tonify Middle-Jiao Qi | Atractylodis Macrocephalae Rhizoma, Astragali Radix, Codonopsis Radix, Glycyrrhiza Radix, Pseudostellariae Radix, Dioscoreae Rhizoma |
| Pungent, warm/bitter, warm | Qi-regulating | 119                                    | 61.34                | Mental depression, restlessness, sighing, distension, wandering pain in the costal and hypochondriac region, distress in epigastrium, poor appetite or vomiting, irregular bowel movements, thin greasy tongue coating, wiry pulse. | Disperse Liver Qi | Citri Reticulatae Pericarpium, Aucklandiae Radix, Aurantii Fructus, Fructus Evodiae |
| Sweet, neutral | Food abating                        | 55                                     | 28.35                | Epigastric and abdominal distension and fullness or pain, which are aggravated by food intake, belching with foul smell, anorexia, constipation, acidic regurgitation, nausea, vomiting, diarrhea with foul smell or fermented contents or constipation | Dissolve the stagnation | Galli Gigeriae Endothelium Corneum, Hordei Fructus Germinatus, Crataegi Fructus |

TCM, traditional Chinese medicine; GC, gastric cancer.
in CF 1-4 were 74, 106, 154, and 5, respectively. With 35 compounds hitting no corresponding targets, a total of 2,128 predictive targets were retrieved and normalized via prediction databases and UniProt, with the potential to interact with 305 active components.

To investigate the relationship between the predictive targets and GC, 429 GC-related genes were screened, and 136 targets overlapped. As shown in Figure 3A, CF 1 targeted the most GC-related genes and was defined as the “preferred prescription” in our study. The CF 1-4 contributed to 135, 12, 13, and 4 genes/proteins, respectively. As shown in Figure 3B, the shared genes/proteins among the 4 formulae included human telomerase reverse transcriptase (hTERT), tyrosine-protein phosphatase non-receptor type 11 (PTPN11), estrogen receptor (ESR1), and sonic hedgehog protein (SHH), G2/mitotic-specific cyclin-B1 (CCNB1), fibroblast growth factor 2 (FGF2), and so on.

Construction and topological analysis of the compound-target network of the preferred prescription

Topological analysis of the C-T network was conducted to identify the crucial components and targets in the preferred prescription. As shown in Figure 3C, the network embodied 505 nodes (11 herbs, 61 active components, and 429 target genes/proteins), and 952 C-T interactions. The mean degree of the active components was 15.61. There were 23 compounds with a degree value higher than 15.61. In this network, crucial substances quercetin (Astragali Radix, Herba Hedyotidis, Radix Actinidiae Chinensis, Crataegi Fructus), kaempferol (Astragali Radix), baicalein (Pinelliae Rhizoma), and nobiletin (Citri Reticulatae Pericarpium, Aurantii Fructus) targeted 256, 191, 144, and 131 GC-related genes, respectively. There were 49 targets with degree values higher than 6.57, the mean degree of the predicted targets. The TP53, bTERT, vascular endothelial growth factor A (VEGFA), caspase-3 (CASP3), murine double minute 2 (MDM2), matrix metalloproteinase 2 (MMP2), and apoptosis regulator Bcl-2 (BCL2) genes were targeted by 26, 19, 16, 16, 14, 14, and 12 compounds respectively, which indicated they may be involved in the underlying mechanisms of the preferred prescription.

Table 2 Top 10 pairs of couplet herbs used in clinical prescriptions

| Herb (LHS)                  | Number of prescriptions | Herb (RHS)                  | Number of occurrences | Support (LHS) (%) | Confidence (LHS ≥ RHS) (%) | LIFT  |
|-----------------------------|-------------------------|-----------------------------|-----------------------|------------------|---------------------------|-------|
| Atractylodis Macrocephalae Rhizoma | 145                      | Dioscoreae Rhizoma          | → 26                  | 13.40            | 92.31                     | 1.24  |
| Atractylodis Macrocephalae Rhizoma | 145                      | Aucklandiae Radix           | → 30                  | 15.46            | 90.00                     | 1.20  |
| Atractylodis Macrocephalae Rhizoma | 145                      | Codonopsis Radix            | → 105                 | 54.12            | 88.57                     | 1.19  |
| Astragali Radix             | 107                      | Liguistri Lucidi Fructus    | → 24                  | 12.37            | 87.50                     | 1.59  |
| Atractylodis Macrocephalae Rhizoma | 145                      | Poria                       | → 126                 | 64.95            | 87.30                     | 1.17  |
| Atractylodis Macrocephalae Rhizoma | 145                      | Coicis Semen                | → 63                  | 32.47            | 87.30                     | 1.17  |
| Poria                       | 126                      | Pseudostellariae Radix      | → 29                  | 14.95            | 86.21                     | 1.33  |
| Poria                       | 126                      | Amomi Fructus               | → 28                  | 14.43            | 85.71                     | 1.32  |
| Atractylodis Macrocephalae Rhizoma | 145                      | Amomi Fructus               | → 28                  | 14.43            | 85.71                     | 1.15  |
| Codonopsis Radix            | 105                      | Hordei Fructus Germinatus   | → 25                  | 12.89            | 84.00                     | 1.55  |

LHS, left hand side; RHS, right hand side.

GO and pathway enrichment analysis of the preferred prescription

To explore the potential mechanism of the preferred prescription, we utilized the DAVID database to decipher the information related to gene ontology. The GO analysis on the targets of the preferred prescription is shown in Figure 4A. The significant biological processes (BP) (P<0.05) included apoptotic process (GO: 0006915), cell adhesion (GO: 0007155), cell cycle arrest (GO: 0007050), and signal transduction (GO: 0007165). The significant molecular functions (MF) (P<0.05) included protein kinase activity (GO: 0004672), cadherin binding involved in cell-cell adhesion (GO:0098641), enzyme binding (GO:0019899), and ubiquitin-protein ligase binding (GO:0031625). The
Figure 3 Target mapping of the candidate formulae to GC-related genes and compound-target network construction of the preferred prescription. (A) Overlapping diagram. The segments of the outside circle represent GC-associated genes (light orange), CF 1 targets (red), CF 2 targets (blue), CF 3 (green) and CF 4 targets (purple). The inside circle, specifically, the dark orange segments represent the overlapping parts. (B) PPI network of GC-related genes of the candidate formulae. The nodes represent GC-related genes/proteins from CF 1 targets (red), CF 2 targets (blue), CF 3 (green), and CF 4 targets (purple). Nodes with more than 1 color represent the shared genes/proteins among different formulae. (C) Compound-target network of the preferred prescription. The nodes represent Chinese herbs (red ellipse), active components (yellow ellipse), GC-associated genes (light blue ellipse), GC-related (turquoise ellipse), and GC-unrelated (green ellipse) predicted targets of the preferred prescription. GC, gastric cancer; CF, candidate formula; PPI, protein-protein interaction.
significant cellular components (CC) (P<0.05) included nucleus (GO:0005634), cytoplasm (GO:0005737), plasma membrane (GO:0005886), and cytosol (GO:0005829).

The significant KEGG pathways were mainly the pathways in cancer, T cell receptor signaling pathway, Toll-like receptor signaling pathway, apoptosis, and the VEGF signaling pathway (Figure 4B). Cross-talk pathways network of the preferred prescription is shown in Figure 4C. Moreover, a multi-regulation map of KEGG pathways was demonstrated, indicating the preferred prescription may exert inhibition on both tumorigenesis and progression of GC (Figure 4D).

**Modular characteristics and molecular mechanism of the preferred prescription for GC treatment**

To elucidate modular characteristics of the preferred prescription, we summarized the significantly enriched BP, KEGG signaling, and reactome pathway of the decoction, which was mainly distributed in the modules of immune regulation, epithelial-mesenchymal transition (EMT), and cell apoptosis/cell cycle (Table 3). Then, an herb-crucial compound-biological functional module-molecule network was constructed to determine the relationships among these elements (Figure 5A). For example, *Atractylodis Macrocephalae Rhizoma*, a core herb in different combinational patterns, is known to invigorate the Spleen and tonify the vital-Qi, which predominantly regulates the immune module. *Astragali Radix, Pinelliae Rhizoma, Citri Reticulatae Pericarpium, Amomi Fructus, Hordei Fructus Germinatus*, and *Aurantii Fructus* are known to regulate the movement of Qi, promote blood circulation, and disperse blood stasis, which mainly regulate the EMT module. In addition, GeneMANIA was used to analyze the interactions among the significant targets, which were enriched in the pathways of each functional biological module. The results

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**Table 3**

| GO:0005634 | nucleus |
|------------|---------|
| GO:0005737 | cytoplasm |
| GO:0005886 | plasma membrane |
| GO:0005829 | cytosol |

**Figure 4**

- **A**
  - GO:0005886: Plasma membrane
  - GO:0005737: Cytoplasm

- **B**
  - hsa05200: Pathways in cancer
  - hsa04060: Cytokine-cytokine receptor interaction

- **C**
  - BP: Biological process
  - MF: Molecular function

- **D**
  - GO:0005576: Extracellular region
  - GO:0005654: Nucleoplasm
  - GO:0005634: nucleus
  - GO:0005737: cytoplasm
  - GO:0005886: plasma membrane
Figure 4 GO and pathway enrichment analysis of the preferred prescription. (A) GO analysis of significant BP, MF, and CC (P<0.05). (B) The significant KEGG pathways (P<0.05). (C) Cross-talk pathways network of the preferred prescription. The nodes represent KEGG pathway terms (P<0.05), and the closer colors they have, the more similar potential functions they possess. The size of nodes represents the enrichment significance of KEGG pathway terms. (D) Multiregulation map of KEGG pathways reflects interactions among crucial components and targets overlapped with GC-related genes. GO, Gene Ontology; BP, biological process; MF, molecular function; CC, cellular component; KEGG, Kyoto Encyclopedia of Genes and Genomes; GC, gastric cancer; MSS, microsatellite stability; MSI, microsatellite instability; EMT, epithelial-mesenchymal transition.
Table 3 Modular functional pathways of the targets in the preferred prescription

| Module                              | Type          | GO biological process/KEGG signaling pathway/reactome pathway                              | P value (Benjamini adjusted) |
|-------------------------------------|---------------|-------------------------------------------------------------------------------------------|------------------------------|
| Immune                              | Reactome      | Immune system                                                                             | 6.18E-05                     |
|                                     | KEGG          | T cell receptor signaling pathway                                                          | 5.71E-09                     |
|                                     | KEGG          | Toll-like receptor signaling pathway                                                       | 8.98E-08                     |
|                                     | KEGG          | NOD-like receptor signaling pathway                                                        | 2.66E-04                     |
| Cell apoptosis/cell cycle           | KEGG          | Apoptosis                                                                                 | 1.35E-06                     |
|                                     | KEGG          | Cell cycle                                                                                | 9.49121E-08                  |
| Epithelial mesenchymal transition   | GO            | Angiogenesis                                                                               | 0.000915735                  |
|                                     | Reactome      | Extracellular matrix organization                                                         | 3.78222E-05                  |
|                                     | KEGG          | Focal adhesion                                                                            | 4.85619E-08                  |
|                                     | KEGG          | Wnt signaling pathway                                                                      | 0.005181823                  |

GO, Gene Ontology; KEGG, Kyoto Encyclopedia of Genes and Genomes; NOD, nucleotide-binding and oligomerization domain.

indicated that the preferred prescription has the exact substance basis to regulate the biological modules related to the pathophysiology of GC.

To confirm the molecular mechanism underlying the preferred prescription, molecular docking was performed. Based on the sub-network between the crucial substances and predominant targets of the preferred prescription (Figure 5B), we found that both nobiletin and kaempferol have strong affinity with TERT and MDM2 molecules, and p53 may function as a downstream target (Figure 5C-5F).

The preferred prescription suppressed GC proliferation and induced cell apoptosis

The typical HPLC-DAD chromatogram of all 12 major components in the preferred prescription is shown in Figure 6. The calycosin 7-O-glucoside, rutin, narirutin, naringin, hesperidin, neohesperidin, calycosin, naringenin, kaempferol, formononetin, nobiletin and atractylenolide II contents in the decoction were determined as 0.010, 0.014, 0.229, 0.214, 0.253, 0.239, 0.002, 0.002, 0.004, 0.001, 0.003, and 0.002 mg/g, respectively.

To determine the effect of the preferred prescription on GC cells, the AGS, HGC27, MKN28, and SGC7901 cell activities were assessed by MTT assay. As shown in Figure 7A, the cellular viabilities of the 4 GC cell lines were all significantly inhibited. In nude mouse xenograft models, we further validated that the preferred prescription-treated mice showed dramatically decreased tumor weights compared to the control (Figure 7B,7C). Notably, body mass did not change over the preferred prescription treatment time courses and mice appeared healthy over the duration of the experiments, suggesting that no significant adverse side-effects were experienced (Figure 7D, Figure S1). These results suggested that the preferred prescription treatment suppressed GC proliferation in vivo and in vitro.

Inducing the apoptosis of cancer cells is a vital way for anticancer drugs to take effect. Therefore, the GC cells were treated with different concentrations of the preferred prescription, and Annexin V-positive cells were detected by FITC analysis to evaluate whether the preferred prescription could induce apoptotic cell death. As shown in Figure 7E,7F, the preferred prescription significantly and dose-dependently increased the apoptosis rates of GC cells compared to the control. We also detected that the preferred prescription significantly increased the expression of pro-apoptotic protein Bax and decreased the expression of antiapoptotic Bcl2 protein in a concentration-dependent manner (Figure 7G-7I). Similar results were found in vivo (Figure 7J,7K). Taken together, our findings indicated that the preferred prescription inhibited the growth of GC by inducing apoptosis both in vivo and in vitro.

The preferred prescription induced GC cell cycle arrest via hTERT/MDM2-p53 signaling pathway

From the results of KEGG pathway enrichment and the sub-network among the crucial substances and
Figure 5 Modular characteristics and molecular docking of the preferred prescription. (A) Herb-key compound-biological functional module-molecule network. Gray dotted line stands for the predicted relationship between herb and crucial compound. Black dotted line stands for the predicted relationship between herb and functional module of the predictive targets. The prefuse force directed layout of the PPI network by GeneMANIA is based on edge betweenness score. The black nodes represent queried proteins. The network weighting of relationships between proteins are shown at the left top. (B) Sub-network among the crucial substances and predominant targets. Molecular docking scores of kaempferol and nobiletin with TERT and MDM2 protein targets were −8.1 (C), −7.6 (D), −9.0 (E), −8.4 (F) kcal/mol, respectively. PPI, protein-protein interaction; TERT, telomerase reverse transcriptase; MDM2, murine double minute 2.
Figure 6 The HPLC-DAD chromatogram of the main components in the preferred prescription. 1: calycosin 7-O-glucoside; 2: rutin; 3: narirutin; 4: naringin; 5: hesperidin; 6: neohesperidin; 7: calycosin; 8: naringenin; 9: kaempferol; 10: formononetin; 11: nobiletin; 12: atractylenolide II. The contents of them in the preferred prescription were 0.010, 0.014, 0.229, 0.214, 0.253, 0.239, 0.002, 0.002, 0.004, 0.001, 0.003, and 0.002 mg/g respectively. HPLC-DAD, high-performance liquid chromatography diode array detection.

the predominant targets, the effects of the preferred prescription were evaluated on the hTERT/MDM2-p53 signaling pathway. The activation of the p53 protein initiates a program of cell cycle arrest, cellular senescence, or apoptosis (34). During different phases of cell cycle, p53 controls both the G1 and G2/M checkpoints (35). Therefore, we performed flow cytometry assay to evaluate whether the preferred prescription modulated the cell cycle of GC cells. As shown in Figure 8A,8B, the proportion of MKN28 cells in the G1 phase was increased and the proportion of cells in the S phase was decreased, while there was a significant increment in G2/M in AGS, HGC-27, SGC-7901 cells.

By binding to p53, MDM2 inactivates the suppressive function of the tumor in p53 and prevents it from intervening in the cell cycle (36). Cells lacking TERT possessed elevated p53 levels and transcriptional signatures were consistent with p53 up-regulation. Thus, we examined the lysates of the MKN28 cells treated with different concentrations of the preferred prescription using western blot assay. As shown in Figure 8C, the preferred prescription treatment significantly and dose-dependently decreased the expressions of hTERT and MDM2, and significantly increased the expression of p53. A major player in the p53-mediated G1 arrest is the p21 gene product that inhibits cyclin E-cdk2 (34). Therefore, we further detected the expressions of p21, cyclinE, and CDK2. It was found that the preferred prescription significantly increased the expression ratio of p21 and reduced the ratios of cyclinE and CDK2 in MKN28 cells (Figure 8C,8D). Similar results were verified in nude mouse xenograft models (Figure 8E,8F). The above findings implied that the preferred prescription induced cell cycle arrest in GC cells via hTERT/MDM2-p53 signaling pathway. It also supported the causal link between the elevated p53 by the preferred prescription and the induction of pro-apoptosis proteins of Bax, and the depletion of anti-apoptosis proteins Bcl-2.

The preferred prescription inhibited EMT of GC cells via the hTERT/MDM2-p53 signaling pathway

In cancer, EMT is associated with tumor initiation, invasion, metastasis, and resistance to therapy (37). The role of p53 in EMT has been well studied (38). Recently, it has been reported that p53, p21, and MDM2 bind to the EMT-inducing transcriptional factors Snail/Slug, and promote its ubiquitin-mediated proteasomal degradation (39,40). Based on these existing studies and our findings above, we hypothesized that the preferred prescription also exerted inhibition on EMT of GC via the hTERT/MDM2-p53 signaling pathway. We performed wound healing assay, which revealed that the preferred prescription
Figure 7 The preferred prescription suppressed GC cells proliferation and induced cell apoptosis. (A) MTT assay showing a concentration-dependent effect of the preferred prescription on the viability of AGS, HGC27, MKN28, and SGC7901 cells. (B) The inhibitory effect of the preferred prescription on the tumor growth of nude mouse xenograft models. The weights of the tumors (C), and the weights of the nude mice (D) were monitored (n=5). (E,F) Flow cytometry depicting cell apoptosis of GC cells treated with different concentrations of the preferred prescription. (G-I) The pro- and anti-apoptotic proteins were detected in MKN28 cells treated with different concentrations of the preferred prescription by western blotting. (J-K) The pro- and antiapoptotic proteins were detected in tumors of nude mouse xenograft models (n=5) by western blotting; *, P<0.05; **, P<0.01, ***, P<0.001, ****, P<0.0001 vs. Control. GC, gastric cancer; MTT, 3-(4,5-Dimethyl-2-thiazolyl)-2,5-diphenyltetrazolium bromide.
Figure 8 The preferred prescription induced GC cell cycle arrest via hTERT/MDM2-p53 signaling pathway. (A,B) Flow cytometry depicting the effect of the preferred prescription on cell cycle checkpoints in GC cells treated with different concentrations of the preferred prescription. The expressions of hTERT, MDM2, p53, p21, cyclinE, and CDK2 proteins were detected and quantified in MKN28 cells treated with different concentrations of the preferred prescription (C,D), and in extracts from transplanted tumors (n=5) (E,F) by western blotting; *, P<0.05, **, P<0.01, ***, P<0.001, ****, P<0.0001 vs. Control. GC, gastric cancer; hTERT, human telomerase reverse transcriptase; MDM2, murine double minute 2; CDK2, cyclin-dependent kinase 2.
remarkably suppressed the migration of MKN28 cells in a concentration-dependent manner, and notably the wound healing area of the preferred prescription (8 mg/mL) group was still large after culturing for 48 h (Figure 9A,9B). Transwell assay revealed that the preferred prescription dramatically inhibited GC invasion, even at the lowest concentration of 2 mg/mL (Figure 9C,9D). Further, we examined the EMT-related markers, and found that after the preferred prescription treatment, N-cadherin, Snail, and Slug expression ratios were significantly reduced in MKN28 GC cells (Figure 9E,9F) and in vivo (Figure 9G,9H) compared to the controls. Collectively, these data suggest that the preferred prescription may suppress EMT of GC via the hTERT/MDM2-p53 signaling pathway.

Discussion

Globally, GC is the most common cancer and is the leading cause of cancer deaths (1). Therefore, the discovery of novel therapeutic strategies is urgent to enhance the therapeutic effects of existing drugs. Recently, with the development of bioinformatics and network pharmacology, more researchers have applied these methods to unravel the therapeutic effects of TCM formulae (4,41). Elaborately prescribed herbal formulae are being increasingly beneficial for GC patients in relieving adverse events caused by CT or CRT, expediting postoperative recovery, and reducing recurrence or metastasis incidence. However, the modular functional characteristics and molecular mechanisms of TCM in ameliorating GC have remained unclear. In the current study, we conducted a comprehensive data mining of clinical prescriptions, based on which, we obtained a basic TCM prescription with general applicability for GC treatment via machine learning. Based on network pharmacology exploration, the pharmacological mechanism of this preferred prescription against GC was also clarified via experimental verification.

In this study, all the TCM prescriptions for GC treatment from eligible clinical studies over the past 3 decades were collected via CNKI, which is a predominant academic database containing the most comprehensive and authoritative information on TCM. A total of 194 prescriptions were retrieved, among which the most common used couplet herb pairs and triplet herbal combinations were analyzed through ARM. The top 3 recommended herb pairs were all led by Atractylodis Macrocephalae Rhizoma. As one of the most potent herbs to invigorate the Spleen and tonify the Qi, it is especially indicated for poor appetite, loose stools, and diarrhea, which are the common manifestations in GC patients, especially after postoperative CT or CRT. When paired with Dioscoreae Rhizoma, with the potential to nourish the Yin and tonify the Lung and Kidney, it improves appetite, and mitigates loose stools. When paired with Aucklandiae Radix, it promotes digestion and relieves pain. Codonopsis Radix can be used with Atractylodis Macrocephalae Rhizoma in GC patients with weak physique and lassitude, or those experiencing cold pain in the stomach and abdomen, vomiting, or diarrhea (8). In addition, we found that Poria-Pseudostellariae Radix-Atractylodis Macrocephalae Rhizoma was one of the most recommended triplet herbal combinations. Notably, they are the major compositions of Si Jun Zi decoction, a classic traditional Chinese herbal prescription, which is well known for treating digestive function disorders (42). A deep machine learning method-based cluster analysis was further conducted to obtain an optimized prescription. In this preferred prescription, Atractylodis Macrocephalae Rhizoma and Astragali Radix were found to be major herbs responsible for tonifying the Middle-Jiao and invigorating the Spleen, which could relieve fatigue, poor appetite, loose stools, and other symptoms in GC patients. Pinelliae Rhizoma, Citri Reticulatae Pericarpium, Aucklandiae Radix, Anommi Fructus, and Aurantii Fructus worked cooperatively to alleviate symptoms like fullness in the abdomen, belching, nausea, and vomiting in GC patients with dampness stagnation. Dyspepsia is very common among GC patients, especially after surgery. Thus, Galli Gigeriae Endothelium Corneum, Hordei Fructus Germinatus, and Crataegi Fructus in the preferred prescription were used to improve digestive dysfunction. For patients enduring a long course of the disease, Herba Hedyotidis and Radix Actinidiae Chinensis could help in the elimination of internal toxins. Generally, the treatment principles and clinical indications of the preferred prescription are summarized in Figure S2.

Next, we identified the bioactive components and the potential molecule targets of the preferred prescription by network pharmacology. A total of 74 bioactive components were acquired from literature and various public databases, 2,128 genes relevant to the preferred prescription were obtained via target prediction, and 429 GC-related targets were retrieved from disease databases. Eventually, 135 overlapping genes were identified as disease-associated targets. The disease of GC is heterogeneous, whereby the presence of molecular heterogeneity has been described based on anatomic histopathology, the anatomic site, gene expression, and so on (43). The multi-compound and
Figure 9 The preferred prescription inhibited EMT of GC cells via the hTERT/MDM2-p53 signaling pathway. Wound healing assay and Transwell assay detecting the migration (A,B) and invasion (C,D) abilities of MKN28 cells treated with different concentrations of the preferred prescription (scale bar =80 μm). The expressions of EMT-related markers were detected and quantified in MKN28 cells treated with different concentrations of the preferred prescription (E,F), and in extracts of the transplanted tumors (n=5) (G,H) by western blotting; *, P<0.05, **, P<0.01, ***, P<0.001, ****, P<0.0001 vs. Control. EMT, epithelial-mesenchymal transition; GC, gastric cancer; hTERT, human telomerase reverse transcriptase; MDM2, murine double minute 2.
multi-target characteristics of the preferred prescription potentiate its multiple biological functions in treating GC. To acquire an in-depth understanding of the overlapping targets, GO, KEGG, and reactome pathway enrichment were performed. The modular functional network revealed part of the combinational rules of herbs in the context of biological functional molecules (Figure 5A). In this study, the crucial compounds ranked by degree were quercetin, kaempferol, baicalein, nobiletin, and luteolin. Existing studies on these bioactive substances have shown diverse anti-GC mechanisms. For example, luteolin could shift the Bax/Bcl ratio in human GC cells by increasing the expressions of pro-apoptotic proteins (44-46). Treatment with luteolin was also observed to up-regulate p21/cip1 (CDKN1A), a TP53 activity signature (47). The mixture of Radix Actinidiae Chinensis could down-regulate the expressions of stromal cell-derived factor-1 (SDF-1), MMP-2, and MMP-9 in SGC-7901 cells (48). Quercetin was found to restrain transforming growth factor (TGF)-β1-induced EMT by inhibiting Twist1 and regulating E-cadherin expression (49). Also, quercetin-3-methyl ether (Q3ME) is a natural flavonoid compound capable of inhibiting esophageal carcinogenesis by targeting the receptor tyrosine kinases (RTKs) (50). Although there was no literature on some crucial components associated with GC, the efficacy was noteworthy in other cancers. All literature, together with the experimental studies, provided a valuable hint in identifying the action mechanism of the preferred prescription against GC.

From the in vivo results, it was revealed that treatment with the preferred prescription significantly suppressed tumor growth compared to the control. Notably, the preferred prescription did not compromise the mice’s body weights compared to the 5-FU group, indicating that it has a better safety profile, or at least in part, is favorable for patients who are intolerant of 5-FU treatment. To explore the potential mechanism, multiple biological function assays were conducted in vitro in GC cell lines, including AGS, HGC-27, MKN28, and SGC-7901. It was demonstrated that the preferred prescription promoted cellular apoptosis and attenuated the metastatic capability in GC cells.

From the results of target prediction and pathway analysis, the preferred prescription might suppress the survival and metastasis of GC cells via the hTERT/MDM2-p53 signaling pathway. The TERT protein is often overexpressed in tumor cells and mediates cellular immortalization (51). Recent research revealed that cells lacking TERT possessed elevated p53 levels and transcriptional signatures were consistent with p53 up-regulation. The up-regulation of the MDM2 oncogene plays a role in the diffuse type of GC (52). By binding to p53, MDM2 inactivates the anti-tumor function of p53 and prevents it from intervening in the cell cycle (36). The activation of p53 induces p53-dependent cell death and p53- and p21-dependent cell cycle arrests, which is characterized by depletion of the S-phase cells and accumulation at the G1/S and/or G2/M phase boundaries of the cell cycle (53). In the present study, the flow cytometry and western blot results supported the prediction. After treatment with the preferred prescription, the proportion of MKN28 cells in the G1 phase was increased and the proportion of cells in the S phase was decreased, while there was a significant increment in G2/M in AGS, HGC-27, and SGC-7901 cells. Additionally, we validated that the preferred prescription exerted negative modulation on the expressions of hTERT and MDM2, and positively modulated the expressions of p53 and p21. The activation of p53 stimulates the synthesis of the p21 protein, which inhibits cyclin E-cdk2 activity, and this in turn acts upon the retinoblastoma (Rb)-MDM2 complex that promotes p53 activity and apoptosis (34). In this study, we demonstrated that after the preferred prescription treatment, increased p53 activity induced the pro-apoptosis protein Bax and depleted the anti-apoptosis protein Bcl-2. We also detected decreased expressions of Slug and Snail under the preferred prescription treatment, which was possibly due to p53, p21, and MDM2 interacting with the EMT-inducing transcriptional factors, and leading to their ubiquitination (39). Taken together, the preferred prescription might play a role in inducing cell cycle arrest, cellular apoptosis, and inhibiting EMT process of GC via the hTERT/MDM2-p53 signaling pathway.

Meanwhile, our research had several limitations. Firstly, the eligible literature in our study was drawn only from Chinese databases. With the development of TCM, we will be able to include more information from other Asian countries like Japan and South Korea. Secondly, the bioactive substances of the Chinese herbs screened in the existing databases need further preclinical and clinical verification. Lastly, a clinical trial on the preferred prescription is required to reliably assess the roles of TCM in the recurrence and metastasis of GC.

Conclusions
To conclude, data mining and machine learning combined with network pharmacology analysis and experimental verification may elucidate the modular functions and
pharmacological mechanisms of TCM on GC from an innovative perspective. It was demonstrated that the preferred prescription may suppress the survival and metastasis of GC cells via modulating the hTERT/MDM2-p53 signaling pathway. Meanwhile, in-depth pharmacological mechanisms by which the preferred prescription ameliorates GC need to be further explored. Also, as the core concept of TCM, syndrome differentiation cannot be completely replaced by the results of machine learning. Hence, clinically effective combinations of herbs should also be encouraged as individualized strategies for GC patients. This study will facilitate the application of TCM in GC treatment with the purpose of improving therapeutic strategy in clinic.

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Footnote

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. This study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). Animal experiment was performed under a project license (No. 2021DW-35-01) granted by the Animal Ethics Committee of Affiliated Hospital of Nanjing University of Chinese Medicine (Nanjing, China), in compliance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health.

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