Mining of Nutritional Ingredients in Food for Disease Analysis

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ABSTRACT Suitable nutritional diets have been widely recognized as important measures to prevent and control non-communicable diseases (NCDs). However, there is little research on nutritional ingredients in food now, which are beneficial to the rehabilitation of NCDs. In this paper, we profoundly analyzed the relationship between nutritional ingredients and diseases by using data mining methods. First, more than 7000 diseases were obtained, and we collected the recommended food and taboo food for each disease. Then, referring to the China Food Nutrition, we used noise intensity and information entropy to find out which nutritional ingredients can exert positive effects on diseases. Finally, we proposed an improved algorithm named CVNDA_Red based on rough sets to select the corresponding core ingredients from the positive nutritional ingredients. To the best of our knowledge, this is the first study to discuss the relationship between nutritional ingredients in food and diseases through data mining based on rough set theory in China. The experiments on real-life data show that our method based on data mining improves the performance compared with the traditional statistical approach, with the precision of 1.682. In addition, for some common diseases, such as diabetes, hypertension and heart disease, our work is able to identify correctly the first two or three nutritional ingredients in food that can benefit the rehabilitation of those diseases. These experimental results demonstrate the effectiveness of applying data mining in selecting of nutritional ingredients in food for disease analysis.

INDEX TERMS Attribute reduction, data mining, disease analysis, food, information entropy, nutritional ingredients, rough sets.

I. INTRODUCTION

NCDs are chronic diseases, which are mainly caused by occupational and environmental factors, lifestyles and behaviors, including Obesity, Diabetes, Hypertension, Tumors and other diseases. According to the Global Status Report on Non-communicable Diseases issued by the WHO, the annual death toll from NCDs keeps adding up, which has caused serious economic burden to the world. About 40 million people died from NCDs each year, which is equivalent to 70% of the global death toll [1]–[3]. Statistics of Chinese Resident’s Chronic Disease and Nutrition shows that, the number of the patients suffering from NCDs in China is higher than the number in any other countries in the world, and the current prevalence rate has blown out. In addition, the population aged 60 or over in China has reached 230 million and about two-thirds of them are suffering from NCDs according to the official statistics [4], [5]. Therefore, relevant departments in each country, especially in China, such as medical colleges, hospitals and disease research centers all are concerned about NCDs.

Suitable nutritional diets play an important role in maintaining health and preventing the occurrence of NCDs [2], [6]. With the gradual recognition of this concept, China has also repositioned the impact of food on health. However, research on nutritional ingredients in food via data mining, which are conducive to the rehabilitation of diseases is still rare in China. At present, China has just begun the IT (Information Technology) construction of smart healthcare. Most studies on the relationship between nutritional ingredients in food and diseases are still through expensive precision instruments or long-term clinical trials. In addition, there are also many prevention reports, but they studied only
one or several diseases [7]. In China, studying the relationship between nutritional ingredients and diseases using data mining is immature. Most doctors only recommend the specific food to patients suffering from NCDs, without giving any relevant nutrition information, especially about nutritional ingredients in food [8].

The solutions for NCDs require interdisciplinary knowledge [2], [9]. In the era of big data, data mining has become an essential way of discovering new knowledge in various fields, especially in disease prediction and accurate health-care (AHC). It has become a core support for preventive medicine, basic medicine and clinical medicine research [10]. With respect to the disease analysis through the mining of nutritional ingredients in food, we mainly make the following contributions: (i) We extracted data related to Chinese diseases, corresponding recommended food and taboo food for each disease as many as possible from medical and official websites to create a valuable knowledge base that are available online; (ii) Applying noise-intensity and information entropy to find out which nutritional ingredients in food can exert positive effects to diseases; (iii) In this paper, the data is continuous and has no decision attributes. To address this problem, we proposed an improved algorithm named CVNDA_Red based on rough set theory, which can better select corresponding core ingredients from the positive nutritional ingredients in food.

The structure of this paper is organized as follows: Section II reviews the related work in the field of disease analysis and data mining. Section III describes the specific data mining algorithms used in this paper, reasons why we select the algorithms, as well as two evaluation indexes. Section IV elaborates the data, experimental results and analysis in detail. Section V presents discussions between methods. Some conclusions and potential future research directions are also discussed in Section VI.

II. RELATED WORK

With the continuous expansion of the impact of NCDs on people, various countries such as South Korea, the United States, Germany, Netherlands and China have conducted research on relationship between food and diseases in recent years. The United States used the Framingham Cohort to find out the relationship between the incidence of Coronary disease and nutritional diets [8], [11]. South Korea used factor analysis, reduced rank regression and the rating method to study the relationship between dietary patterns and nutrition-related health problems [12], [13]. Computational results in [14] showed, a daily intake of 100 grams of fresh fruits may reduce almost one third of the risk of Cardiovascular diseases. Li et al. [15] found that Vitamin D supplementation in Chinese population cannot completely improve the lack of Vitamin D. Agapito et al. [16] proposed that we can recommend food according to the body’s Creatinine values. However, the above studies are basically carried out through long-term clinical trials, which just recommend food for certain specific diseases and they seldom study the relationship between nutritional ingredients and diseases by data mining techniques.

Large-scale and high-dimensional data that we extracted from online have usually a noise, hence we must use some techniques to pre-process the data. For example, the data denoising method based on wavelet transform [17] was proposed in the field of signal and data compression. Zhang and Zhang [18] proposed a new SVM-GARCH prediction model based on time series data to select features by mutual information. Diogo et al. [19] applied noise reduction to exchange rate chaotic data. Actually, the noise also can measure the fluctuations in gene expression, which is one of the important biology indicators. Therefore, many literatures indicate that they are currently utilizing the noise-intensity to discover new knowledge [20]. For example, the concrete expression of protein noise and noise-intensity at equilibrium were described [21], [22], and they also analyzed kinetic behavior by the detailed expression at non-equilibrium. Lei et al. [23] used noise-intensity to improve the LDA algorithm to achieve better performance on a protein classification task according to the protein distribution structure.

Information entropy is a measurement to eliminate uncertainty from given information. Since the concept of entropy was introduced into information theory, it has been widely used in different fields. Yang and Jiang [24] applied the maximum fuzzy entropy and fuzzy c-means segmentation [25] to select the thresholds automatically. An image classification method of invariant pixel region texture based on wavelet packet entropy was also proposed [26]. Bakhshali [27] combined the MFE criterion with the MMI criterion to present a maximum fuzzy segmentation algorithm based on mutual information. A method of recovering short vowels and other phonetic symbols from Arabic phonetic symbol documents was proposed by using the maximum entropy model [28]. Tan and Taniar [29] proposed a distribution estimation algorithm based on maximum entropy, and they demonstrated experimentally the superior performance in dealing with clustering problems. An optimization strategy based on maximum entropy in [30] was proposed to process the parallel resource. Chen [31] described a MAODV protocol based on entropy in Ad Hoc networks and proposed a new measure to express the path stability. An extraction method in [32] was presented for pruning larger nodes with information entropy. John et al. [33] used the utility value reflected by information entropy to calculate the weight of evaluation indexes and proposed an AHP fuzzy comprehensive evaluation model. This model can provide a flexible tool for analysts to strengthen security of the port systems.

In rough set theory, the attribute reduction of decision tables is an interesting research topic. In general, we used heuristic methods to reduce attributes. Firstly, the core attributes must be identified, from these the most important attributes must be selected, and finally added to the reduction set [34]. For example, a method based on discernibility matrix and logic operation was given [35]. Nguyen [36] proposed two fusion methods based on logical
reasoning and discernibility matrix. An inductive learning method based on rough sets was proposed for SAR analysis [37]. Zhang et al. [38] proposed a multivariate decision tree method by using rough set theory. An improved algorithm for finding the minimum attribute reduction set was presented by Jia et al. [39] using the decision table’s generalized information. In addition, Zheng [40] defined the attribute significance from the perspective of mutual information, and proposed a MIBK algorithm to reduce attributes. The idea of functional decomposition in [41] and [42] was applied to the decomposition of decision tables. They established a hierarchical decision model by gradually delaminating from an inaccurate single large-scale objective into sub-goals, which improves the efficiency of data analysis. However, most of the above methods are basically based on the decision tables, which are discrete and have decision attributes. These methods will not work well if there is a decision table with continuous values and non-decision attributes. To solve this problem, we proposed an algorithm named CVNDA_Red based on rough sets, which is suitable for continuous values as well as non-decision attributes and can better select corresponding core nutritional ingredients in food for diseases.

### III. METHODOLOGY

To show whether data mining methods can be applied to the disease analysis or not, we adopted different methods for research purpose. Since this paper is trying to solve a new problem from a real-world application, there is no relevant work for comparison. But we used an exploring way to solve this problem, in other words, we conducted an effect comparison between different methods to select the best one. In order to show the characteristics of nutritional ingredients in food, we draw nutritional ingredient values in recommended food and taboo food for four common diseases. In Fig.1, it shows the nutritional ingredient values in the recommended food and taboo food for four diseases, such as Hypertension, Coronary disease, Kidney calculi and Apoplexy. The X-axis and Y-axis represent nutritional ingredients and corresponding values respectively. The meaning of each letter of X-axis is expressed in Table 2. In addition, blue and red represent recommended food, taboo food respectively. It is clear from Fig.1 that some nutritional ingredient values are very high and the others are very low, no matter in recommended food or taboo food. Generally, there are few taboo food for most diseases in China, as a result we can not collect more. Given the less credible knowledge for small sample sizes by data mining, we are currently focusing our research on recommended food to find which nutritional ingredients can exert positive effects for a certain disease. In this paper, the nutritional ingredients that benefit the rehabilitation of diseases are called positive nutritional ingredients (PNIs). On the contrary, the nutritional ingredients that aggravate diseases are named negative nutritional ingredients (NNIs).

#### A. STATISTICAL ALGORITHM (SA)

Fig.1 clearly shows that nutritional ingredient values vary widely. If a certain disease is caused by the lack of certain nutritional ingredients, then their values in recommended food should be relatively higher in theory. Therefore, we can figure out which nutritional ingredient values are high. The nutritional ingredients with higher values should be the ingredients, which benefit the rehabilitation of that specific disease, i.e., PNIs [43]. This method is referred to as statistical algorithm (SA) in this paper. The common notations are defined as following, let \( n \) be the number of the recommended food for a certain disease, and \( m \) be the number of nutritional...
ingredients. The above idea can be expressed as below:

\[
\text{sort}\{\sum_{i=1}^{n} x_{i1}, \sum_{i=1}^{n} x_{i2}, \ldots, \sum_{i=1}^{n} x_{im}\text{[descend]}\} \quad (1)
\]

where \(x_{i1}\) denotes the first nutritional ingredient value of the \(i^{th}\) recommended food for a certain disease; \(x_{i2}\) indicates the second nutritional ingredient value of the \(i^{th}\) recommended food, and so on. All recommended food for this disease is summed according to nutritional ingredient values. Then, the \(m\) summations are sorted in descending order. So, from the sorted nutritional ingredients the most top ones should be PNIs by this way.

**B. NOISE-INTEGRITY ALGORITHM (NI_SA)**

If a certain disease is caused by the lack of some nutritional ingredients, the recommended food should contain these ingredients. In other words, their certain values may not be the highest, but must be existing in all recommended food (high stability), otherwise these recommended food have no reason to be recommended. Conversely, if these nutritional ingredients are not PNIs for that specified disease, they may or may not exist in different recommended food (poor stability). The SA just considers the level of nutritional ingredient values to determine simply whether they are PNIs or not. Therefore, on the basis of SA, we can further consider about the stability of nutritional ingredient values in food for determining whether they are PNIs or not. In this way, two aspects can be considered, i.e., the level and stability of nutritional ingredient values. The greatest advantage of using this method is that it reduces the error rate of what are considered to be PNIs just because of the level of nutritional ingredient values.

In biology, noise-intensity is a measure of stability. As a matter of fact, the noise-intensity can be considered as reflecting the degree of data dispersion, because it is not only affected by the variance, but also controlled by the mean [44]. Therefore, the purpose of using noise-intensity in this method is to express the stability of nutritional ingredient values. It is worth noting that the noise-intensity calculated here is based on the original data, which is raw data, because the variance of processed data is one. The noise-intensity and above idea are expressed as follows:

\[
\varphi_j = \frac{\delta_j}{\mu_j} (j = 1, 2, 3, \ldots, m) \quad (2)
\]

\[
\text{sort}\{\varphi_1, \sum_{i=1}^{n} x_{i1}, \varphi_2, \sum_{i=1}^{n} x_{i2}, \ldots, \varphi_m, \sum_{i=1}^{n} x_{im}\text{[descend]}\} \quad (3)
\]

where \(\varphi_j\) represents noise-intensity of the \(j^{th}\) nutritional ingredient; \(\delta_j\) indicates the variance of the \(j^{th}\) nutritional ingredient and \(\mu_j\) is the corresponding mean. The noise-intensity takes the stability of each nutritional ingredient into account. In this paper, each noise-intensity is treated as a weight scale factor for each corresponding nutritional ingredient summation. This method based on SA is called noise-intensity algorithm (NI_SA).

**C. INFORMATION ENTROPY ALGORITHM (IE_SA)**

Fig.1 clearly illustrates that there are two or more nutritional ingredients with higher values in a disease. For example, the \(Y\) nutritional ingredient and the \(E\) nutritional ingredient are relatively higher than other nutritional ingredients in Hypertension. Similarly, the stability of nutritional ingredients follows the same statistics. To find out which one is a PNI in more than two nutritional ingredients. Based on their similarity value levels or stability, the validity of each nutritional ingredient is considered. For example, if a nutritional ingredient has a good effect to a certain disease, the corresponding validity should be greater than others.

Information entropy measures the degree of disorder of data systems and calculates the amount of valid information. The smaller the information entropy is, the greater validity it provides and vice versa [45]. In this paper, information entropy calculates the validity of each nutritional ingredient. According to the definition of entropy, the entropy of the nutritional ingredient can be expressed as follows:

\[
H_j = -\frac{1}{\ln n} \sum_{i=1}^{n} f_{ji} \ln f_{ji} \quad (4)
\]

In the definition of entropy, \(f_{ji}\) should be considered as the occurrence probability of the \(j^{th}\) nutritional ingredient of the \(i^{th}\) food. Since our data does not have identical food in a disease (i.e., the occurrence probability of each food in a disease is one), Equation (4) is not acceptable in our paper. However, the above mentioned occurrence probability can be replaced by each nutritional ingredient value. Because it makes sense that if there is a higher nutritional ingredient value in food, the occurrence probability is larger. In order to make \(\ln f_{ji}\) reasonable, if \(f_{ji} = 0\), we would note \(f_{ji} \ln f_{ji} = 0\). Then, \(f_{ji}\) is redefined as follows:

\[
f_{ji} = \frac{x_{ji}}{\sum_{i=1}^{n} x_{ji}} \quad (5)
\]

where \(x_{ji}\) represents the value of the \(j^{th}\) nutritional ingredient of the \(i^{th}\) food. The validity of the \(j^{th}\) nutritional ingredient can be defined as follows:

\[
W_j = \frac{1 - H_j}{m - \sum_{j=1}^{m} H_j} \quad (6)
\]

\[
\text{sort}\{W_1, \sum_{i=1}^{n} x_{i1}, W_2, \sum_{i=1}^{n} x_{i2}, \ldots, W_m, \sum_{i=1}^{n} x_{im}\text{[descend]}\} \quad (7)
\]

In this paper, we represent the validity as a weighted scale factor for each corresponding nutritional ingredient summation. This method based on SA is called information entropy statistical algorithm (IE_SA).

**D. NOISE-INTEGRITY AND INFORMATION ENTROPY ALGORITHM (NIE_SA)**

As mentioned above, NI_SA and IE_SA are both based on the SA method, but their starting points are different. One consideration is the stability after the level of the nutritional ingredient values, and the other one is to consider the validity
after the level of the nutritional ingredient values. For integrating the above two methods through a certain way such as voting, the results of each method can be considered as a kind of voting results. If both methods assume a certain nutritional ingredient having positive effects, it is more likely to consider that nutritional ingredient as a positive nutritional ingredient. It is more reasonable than deciding which one of those nutritional ingredients is a PNI by using a single method [46]. The above idea can be expressed as follows:

\[
\text{sort}\left[\begin{array}{c}
\text{sort}_{11} \\
\text{sort}_{12} \\
\vdots \\
\text{sort}_{mn}
\end{array}\right] \text{[ascend]} \tag{8}
\]

where \(\text{sort}_{11}\) represents the ranked serial number of the first nutritional ingredient according to NI_SA; \(\text{sort}_{12}\) shows the ranked serial number of the first nutritional ingredient according to IE_SA, and so on. The sorting strategy is as follows: summing up their ranked serial numbers for the same nutritional ingredients, and then sorting the gained summations in ascending order. According to the above two methods, it is clear that the greater possibility of nutritional ingredients are viewed as positive nutritional ingredients, the smaller summations are sorted, i.e., the top ranked ingredients should be PNIs. In this paper the method based on NI_SA and IE_SA is referred to as NIIE_SA.

**E. ROUGH SET ALGORITHM (RS)**

As can be seen from Table 3, the number of positive nutritional ingredients is different for each disease. For some diseases, 10 out of 26 nutritional ingredients (almost \(\frac{3}{5}\)) are PNIs, while others have only two PNIs. So, can some nutritional ingredients with little positive effects be excluded first? In this way, the rest nutritional ingredients are almost positive to diseases, and the same is achieved for our purposes. In this paper, this method based on attribute reduction is called RS.

The rough set theory is a mathematical tool developed by a Polish computer scientist, Pawlak [47], to study uncertain and inaccurate knowledge [48]. Researchers have proposed lots of attribute reduction algorithms, such as based on positive domain, discernibility matrix, and decision tree. The definition of rough sets is as follows. The form of a decision table in rough sets can be represented by a four-tuples: \(S = (U, A, V, f)\), which is also expressed as \(S = (U, V)\). \(U\) called the universal set represents a non-empty, finite set of objects. \(A\) denotes a non-empty, finite set of attributes, \(C \subseteq D\), where \(C\) is a set of condition attributes, and \(D\) is a set of decision attributes. If \(D = \phi\), then, \(S\) represents a decision table without decision attributes. \(V = \cup_{a \in A} V_a\) is a set of attribute values, \(V_a\) denotes values that the attribute \(a\) may take. \(f : U \times A \rightarrow V\) is an information function that assigns a value to each attribute of each object, i.e., \(\forall a \in A, x \in U, f(x, a) \in V_a\).

Let \(P\) be a subset of \(A\) \((P \subseteq A)\), the associated equivalence relation \(\text{ind}(P)\) can be expressed as: \(\text{ind}(P) = \{(x, y) \in U \times U | \forall a \in P, f(x, a) = f(y, a)\}\). \(\text{ind}(P)\) is called a \(P\)-indiscernibility relation on \(U\). \(U/\text{ind}(P)\) or \(U/P\) represents a set, which uses \(\text{ind}(P)\) to classify \(U\). The equivalence classes of the \(P\)-indiscernibility relation are denoted as \([x]\_P = \{y | y \in U, (x, y) \in \text{ind}(P)\}\). Dependency of the condition attribute \(P\) in the decision attribute \(D\) can be expressed as \(|\text{POS}_P(D)|/|U|\). It represents the positive domain of \(P\) in \(D\). In these sets, \(P\) can uniquely classify all elements in \(U\) as a subset \(U/D\). The significance degree \(\text{sig}(a)\) of the attribute \(a\) in the attribute set \(P\) can be defined as follows:

\[
\text{sig}(a) = |\text{POS}_P(D) - \text{POS}_{P - a}(D)|/|U| \tag{9}
\]

\[
\text{tol}(S) = |\text{POS}_C(P(D))|/|U| \tag{10}
\]

where \(\text{tol}(S)\) is called the compatibility degree of \(S\), if \(|\text{POS}_C(D)| = |U|\), the decision table is considered to be a compatible one, otherwise it is incompatible. Obviously, the necessary and sufficient condition of a compatible decision table is that its compatibility degree equals to one. For all decision tables, contribution of each condition attribute on the decision attributes is different. The contributions of condition attributes to decision attributes are called the attribute significance. The attribute reduction algorithms are reflected by the changes of the classification ability after the deletion of a certain attribute. However, the above attribute reduction algorithms are based on decision tables whose attribute values are discrete and which have decision attributes. The data in this paper is obviously not suitable for those methods. At present, attribute reduction methods with continuous values are through discretization processing, and then to select features without supervision. But the results of discretization are bound to lose a lot of information in this way. So, direct reduction of attributes in the case of continuous values will be more realistic. To this end, we presented an improved algorithm (i.e., called continuous value and no decision attributes reduction, CVNDA_Red) to suit this kind of data. Firstly, establishing food fuzzy similar matrix is needed [49]:

\[
\text{Sim}(o_s, o_t) = \min\{1 - \frac{|x_{sj} - x_{tj}|}{\max x_{tj} - \min x_{tj}}\} \tag{11}
\]

where \(\text{Sim}(o_s, o_t)\) denotes the similarity of food \(s\) and \(t\). The \(\max x_{tj}\) and \(\min x_{tj}\) represent the maximum value and the minimum value of the \(j^{th}\) nutritional ingredient in food for a disease respectively. Then, the square method is used to find the transitive closures of the fuzzy similarity matrix. The aim is to extract the different values of \(\lambda\) from transitive closures. After sorting them in descending order, it needs to calculate the corresponding compatibility degree based on \(\lambda\). This step will be terminated until it is found that the compatibility degree of decision tables is less than one. At this time, \(\lambda\) can be viewed as a next step threshold. In this paper, since there is no decision attributes in data, we treated all condition attributes as decision attributes.

For each condition attribute \(a (a \in C)\), after deleting \(a\), the calculation of the corresponding similarity matrix and transitive closures is needed by the same ways as above in the case of the obtained threshold, and then to cluster the results \((C - \{a\})\). Finally, the significance degree of condition attribute \(a\) is calculated, and other attributes are followed
The relationship among methods used in this paper.

Algorithm 1: CVNDA Red

Input: Food-nutritional ingredients matrix \((n \times m)\)

Output: \(sn\) % the serial numbers of selected nutritional ingredients

1. \(R \leftarrow Sim(Data)\); % R is a symmetric matrix \((n \times n)\)
2. \(S \leftarrow transClosure(R)\); % obtaining transfer closure matrix by square method
3. \(\lambda \leftarrow sort(unique(S), \text{‘descend’});\)
4. for \(i = 1 : size(\lambda)\) do
5. \(xrd = tol(\lambda_i);\)
6. if \(xrd < 1\) then
7. \(yz = \lambda_i;\) % getting the threshold
8. break;
9. end
10. end
11. \(SIG = zeros(1, m);\)
12. for \(i = 1 : m\) do
13. \(R \leftarrow Sim(Data - a);\) % obtaining fuzzy similar matrix after deleting attribute \(a\)
14. \(S \leftarrow transClosure(R);\)
15. \(\lambda \leftarrow sort(unique(S), \text{‘descend’});\)
16. for \(i = 1 : size(\lambda)\) do
17. if \(\lambda_i > yz\) then
18. % nothing need to do!
19. else
20. \(U = union(U, cluster(\lambda_i));\)
21. end
22. end
23. \(SIG(1, i) = sig(U);\)
24. end
25. \(sn = greatZero(SIG);\)
26. return \(sn\)

In Algorithm 1, Line 3 calculates different values of \(\lambda\) according to the transitive closure matrix obtained from Line 2; Line 4-10 obtain the threshold by compatibility degree of decision tables (less than one); Line 16-22 get the equivalence classes according to the \(\lambda_i\), and then cluster the results. The last clustering results are unionized with the next clustering results until \(\lambda_i\) is greater than the threshold \(yz\); Line 25 chooses the serial numbers of nutritional ingredients whose significance degrees are greater than zero.

F. COMBINATION OF NIIE_SA AND ROUGH SETS (RSNIIE_SA)

It is clear from RS that the obtained results are only nutritional ingredients, which are viewed as positive nutritional ingredients, but they are not sorted according to the possibility of judging as PNIs. So, using these results directly on patients will not be effective. In order to more fully identify nutritional ingredients, which have the most effects on the rehabilitation of diseases, results obtained by RS can be sorted. The sorting rule is based on the ranked serial numbers of nutritional ingredients of NIIE_SA. For example, if the nutritional ingredients obtained by RS for Hypertension are A, B and C nutritional ingredients, and the ranked serial numbers of those ingredients obtained by NIIE_SA are the third, the second, and the first respectively, then, the results obtained using this method are C, B and A. In this paper, this method based on RS and NIIE_SA is called RSNIIE_SA.

Fig.2 gives better understanding of the relationship among the above methods intuitively. According to the definition of RS, the results are not sorted according to the possibility of judging as PNIs. Therefore, the results of RS are not presented as the final results here.

IV. EXPERIMENTS

A. DATA SET

We mainly used the combination of Spider technology [50] and Selenium 2 technology [51] based on the Python language to get the data. The main form to get Spider data is the texts that we extracted from web pages. Firstly, we downloads the relevant web pages according to specified URLs, and then uses regular expressions to locate the desired texts. But nowadays it is not easy to get data using Spider as many websites deploy anti-crawling technology. However, another technology called Selenium 2 can be used to extract the data as it is automatic to simulate the people’s behaviors to access to websites. We can also get data from academic papers or articles. In order to obtain high quality data, we have mainly chosen Chinese medical and official websites, including:

- 360baike (https://baike.so.com);
- CDC (http://www.chinacdc.cn);
- XYWY (http://jib.xywy.com);
The instances of data used in this paper:

<table>
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<th>Disease</th>
<th>Recommended food</th>
<th>Taboo food</th>
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<td>Mushroom.</td>
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</tbody>
</table>

All these websites are available online. We follow two steps to obtain the data. Firstly, all the possible diseases were collected, and then the recommended and taboo food were fetched according to the corresponding diseases. We have collected about 12,100 diseases, out of them 9,100 diseases were selected by removing repeated diseases. Based on 9,100 diseases, we have also collected 56,000 data that contains a number of recommended and taboo food. While selecting recommended food or taboo food for each disease, we adopted the principle of intersection to deal with the data, i.e., it is selected if a certain food showed up in each piece of data for a disease. Due to the existence of diseases without recommended and taboo food, the total instances of data become 7,594 after we remove these diseases. Each one of these data includes three items: the disease name, recommended food, and taboo food. Here are several examples:

The referred nutritional ingredient values for each food are from the China Food Composition 2002 [52] and the China Food Composition 2004 [53], which are compiled by the National Institute for Nutrition and Health Chinese Center for Disease Control and Nutrition and Food Safety. The former version includes sixteen nutritional ingredients, whereas the later version contains seventeen nutritional ingredients. The common data we got in the two versions have been deleted and we summed up the remaining data that results 26 nutritional ingredients in food, which finally are presented as follows:

The capital letters represent 26 nutritional ingredients. Since the representation of Vitamin A and Retinol equivalence in China is not consistent and they have different values for some certain food in the two versions, two different nutritional ingredients were considered. In order to avoid missing any possibility, we assumed them as two PNIs if any one of them has positive effects for diseases. If nutritional ingredient values of a certain food are not available in the two versions, a weighted average value of similar food will be filled, otherwise that food will be deleted.

In order to verify the rationality of the above methods, ten common diseases are selected as testing samples as shown in Table 3. These diseases were selected because of their unquestionable PNIs and NNIs, either in China or abroad. Of course, we also proved the correctness of the testing samples below by the relevant medical experts. Since there are no unified conclusions about other diseases, those cannot be verified yet. But for future work, many other examples can be taken to implement the above methods and to verify the feasibility. In this paper, the first letter of the word Disease as well as the word Name and serial number are used to indicate the corresponding diseases.

Vitamin is a set that includes F, G, H, J, N nutritional ingredients; Vitamin B contains G, H nutritional ingredients. Since there are not statistics about Vitamin D and Mineral salt nutritional ingredient values in food, this paper does not discuss whether these two nutritional ingredients are helpful to rehabilitate the diseases or not.

### B. EVALUATION MEASURE

In order to explain and compare the validity of data mining methods applied in our study, two evaluation indexes, individual precision (IP) and over precision (OP), are proposed. They are defined as follows, which are based on the idea that the top ranking nutritional ingredients sorted by a certain method have the greater possibility of being judged as PNIs. In order to do the mathematical calculation, each value of positive nutritional ingredients is noted as one from the obtained results (i.e., the blue color in the experimental results below), and each of the others is noted as zero. For RS and RSNIIIE_SA algorithm, the results of each disease may only
TABLE 3. The positive nutritional ingredients and negative nutritional ingredients for ten diseases.

<table>
<thead>
<tr>
<th>Label</th>
<th>Disease</th>
<th>PNIs</th>
<th>NNIs</th>
</tr>
</thead>
<tbody>
<tr>
<td>DN1</td>
<td>Nearsightedness</td>
<td>Calcium, Vitamin B, Vitamin A, Vitamin C, Vitamin E, Vitamin D</td>
<td>Carbohydrate</td>
</tr>
<tr>
<td>DN2</td>
<td>Caries</td>
<td>Magnesium, Phosphorus, Vitamin</td>
<td>Carbohydrate</td>
</tr>
<tr>
<td>DN3</td>
<td>Chronic cold</td>
<td>Vitamin, Zinc, Selenium</td>
<td>Sodium, Fat, Carbohydrate</td>
</tr>
<tr>
<td>DN4</td>
<td>Hypertension</td>
<td>Fiber, Potassium, Magnesium, Calcium</td>
<td>Sodium, Carbohydrate</td>
</tr>
<tr>
<td>DN5</td>
<td>Acne</td>
<td>Vitamin B, Vitamin A, Zinc</td>
<td>Fat, Energy, Carbohydrate</td>
</tr>
<tr>
<td>DN6</td>
<td>Coronary disease</td>
<td>Vitamin C, Calcium</td>
<td>Cholesterol, Fat</td>
</tr>
<tr>
<td>DN7</td>
<td>Kidney stone</td>
<td>Vitamin, Magnesium</td>
<td>Sodium, Calcium</td>
</tr>
<tr>
<td>DN8</td>
<td>Diabetes</td>
<td>Fiber, Protein, Vitamin, Mineral salt</td>
<td>Fat, Energy</td>
</tr>
<tr>
<td>DN9</td>
<td>Apoplexia</td>
<td>Potassium, Vitamin C, Magnesium</td>
<td>Cholesterol, Sodium</td>
</tr>
<tr>
<td>DN10</td>
<td>Anemia</td>
<td>Iron, Copper, Zinc, Protein, Vitamin B, Renedatene</td>
<td>Calcium, Phosphorus</td>
</tr>
</tbody>
</table>

where \( IP_k = \sum_{j=1}^{m} B_{kj} \left(\frac{1}{j}\right)^l \) (12)

where \( IP_k \) represents the individual precision of the \( k^{th} \) disease, \( B_{kj} \) denotes a binary value (one or zero) of the \( j^{th} \) nutritional ingredient. \( l \) represents the value coefficient (\( l \geq 1 \)), which is added to emphasize the high value of the top ranking nutritional ingredients in experimental results [54], [55]. The value of \( l \) in this paper is two. Of course, in order to emphasize the higher value, it could be made larger. According to actual needs, the value of \( l \) can be changed. The \( IP \) index is mainly to compare the ability of different methods to find out PNIs for a certain disease. According to the Equation (12), the increasing rate will be very slow if the first value is zero, especially when \( l \) is larger (i.e., the purpose is to emphasize the significance of accurately choosing nutritional ingredients as PNIs in the first several times). It is the same in real cases. For example, when a doctor is recommending some food to a patient, he must first choose the food that contains rich PNIs to that patient. If the doctor’s first recommended food does not satisfy patients’ condition, the patients will definitely blame that the doctor did not identify the real cause of disease. Therefore, the above proposed index \( IP \) makes sense.

\[
OP = \frac{1}{\sum_{k=1}^{g} \frac{m_{k}'}{m}} \sum_{j=1}^{m} \left( all \cdot \sum_{k=1}^{g} B_{kj} \left(1 - \frac{m_{k}'}{m}\right) \left(\frac{1}{j}\right)^l \right) \] (13)

where \( all = 1/\sum_{k=1}^{g} \left(1 - \frac{m_{k}'}{m}\right) \) represents the reciprocal of proportion summation of positive nutritional ingredients. \( g \) denotes the number of testing samples. \( m_{k}' \) indicates the PNIs number of the \( k^{th} \) disease. The \( OP \) is to compare the ability of different methods to find out positive nutritional ingredients for all diseases. According to Table 3, it can be seen that the number of PNIs for each disease is not equal. It is obviously unreasonable if all selected nutritional ingredients are viewed as one. For example, there are twenty positive nutritional ingredients for one disease and only two PNIs for other diseases. If there is only one chance to select a positive nutritional ingredient, obviously the correct rate of the disease with twenty positive nutritional ingredients is bigger than other diseases that have only two PNIs. Thus, all coefficient is added for avoiding this error.

C. EXPERIMENTAL RESULTS

1) THE RESULTS OF SA

Nutritional ingredients with blue color indicate PNIs (see Table 3, Column three). The expected experimental result shows that the nutritional ingredients on the left side of each row should be positive nutritional ingredients, i.e., the blue ones are transferred to the left. It is clear from Fig.3 that these are not the expected results. Instead, each row of the left side is filled with Y, E and B nutritional ingredients rather than PNIs. However, those results are consistent with Fig.1, in which the highest nutritional ingredient values for each disease is Y. It is not difficult to explain why the ranking results come out like that, i.e. nutritional ingredients with higher values are all Y, E and B in four diseases.

At present, people are not free to claim only a few nutritional ingredients in food (e.g., I just want to eat the Vitamins inside of fruits or vegetables.). Only professional doctors can do that, but the price is too high. Now, doctors may not select a few nutritional ingredients, and we would fill with zero here if other nutritional ingredients had no values (zero or one). Because if a certain nutritional ingredient was not selected, it would mean that the possibility of being assigned as a PNI is zero.
only consider to recommend these food with rich nutritional ingredients what you may lack, regardless of other nutritional ingredients whether you may lack or not. Therefore, SA is not feasible for selecting which ones are PNIs among many nutritional ingredients in first several times. In other words, only relying on statistical methods can’t solve our problem.

2) THE RESULTS OF NI_SA

Fig. 4 clearly shows that considering the stability of nutritional ingredients in food has great effects. Except for DN8 and DN9, the top ranking nutritional ingredients of each disease have become positive nutritional ingredients in the first time, and even the second ranking nutritional ingredients also have turned into what are expected, i.e., PNIs. Especially for DN1 and DN10, three out of seven PNIs (almost 50%), are selected successfully in first three times. Then, the ranked positions of PNIs are basically shifted left, which indicates that considering stability is a more reasonable direction.

3) THE RESULTS OF IE_SA

Fig. 5 clearly shows that considering the validity of nutritional ingredients in food also leads to better results. Every first position of each row is placed with a positive nutritional ingredient except for DN1 and DN10, three out of seven PNIs (almost 50%), are selected successfully in first three times. Then, the ranked positions of PNIs are basically shifted left, which indicates that considering stability is a more reasonable direction.

4) THE RESULTS OF NIIE_SA

Fig. 6 elaborates that the use of voting strategy has better effects in determining which nutritional ingredients are most likely to become positive nutritional ingredients. This strategy turns the first ranked position of all diseases into a PNI, and the overall PNIs have moved to the left. This ensures that those PNIs can be identified correctly in the first time, so it is feasible to apply this strategy to solve our problem.

5) THE RESULTS OF RS

According to the RS algorithm, in each row of the above results, there is no significance difference among the selected nutritional ingredients. As can be seen from Fig. 7, if only the nutritional ingredients are showing up in the experimental results, they are all considered as PNIs. It can be seen that most selected nutritional ingredients are correct. In order to explain the ability of this method in detail, we used the correct ratio (PNIs / selected nutritional ingredients) for each disease to show as follows:

The bold numbers indicate that more than half of the selected nutritional ingredients are correct. It clearly shows that the correct ratio of six diseases is greater than or equal to 50% and three other diseases (italic numbers) are close to 50%. So it can be demonstrated that most selected nutritional ingredients are correct, and this leads to the success of this method in finding out positive nutritional ingredients for diseases.

6) THE RESULTS OF RSNIIE_SA

The final and satisfactory results are shown in Fig. 8. The first column shows that the results obtained in the first time, that
means the PNIs for each disease. The results we obtained for the second time are shown in column two, but the results in the DN6 disease are not expected. Generally, this method gives the most accurate results among all the other methods, because we can accurately find out the first two or three nutritional ingredients out of 26 ingredients for each disease.

It is the best method that meets our actual needs. For example, if the patient is suffering from the above ten diseases, the recommended food must be rich with the first two or three nutritional ingredients.

V. DISCUSSION

The above results are given independently, and no comparison is made among the methods. So, it is difficult to interpret which method is the best to select positive nutritional ingredients for diseases. According to the above two indexes, the next is a comparison of five methods based on the effect.

The methods where IP equals to 1.5 indicate that the second positive nutritional ingredient is also selected correctly in the second time, which can be seen from Equation (12). Even though the results show that the IP of NIEI_SA is very similar to RSNIEI_SA, the RSNIEI_SA is much better than NIEI_SA from an actual application view. In the case of the same IP, the NIEI_SA gives 26 nutritional ingredients, but RSNIEI_SA gives only a few nutritional ingredients, thus, RSNIEI_SA reduces the possibility of wrong recommended food with PNIs. From the view of individual diseases, RSNIEI_SA is the best, because it identifies positive nutritional ingredients correctly in first two times.

As Fig.10 shows, the parts above the reference line indicate that the first position are PNIs for all testing samples. As can be also seen from Fig.10, when we plot the graph using Matlab, we have only used three decimal points. From Fig.10, SA selects successfully zero positive nutritional ingredients for all diseases. For the NI_SA, the value of OP is 1.081, which is almost one, this shows that the PNIs are all selected successfully in the first time. In Fig.9, we do not see big performance difference between NIEI_SA and RSNIEI_SA. However, it is quite obvious in Fig.10 that RSNIEI_SA is the best one, because the OP value of RSNIEI_SA is 0.5 higher than NIEI_SA. Over all, the value of OP using data mining methods is greater than the OP value of the statistical SA method, especially RSNIEI_SA,
it achieved 1.682 improvements, which shows that it can be applied in the disease analysis based on the mining of nutritional ingredients in food.

VI. CONCLUSION

The main work of this paper can be divided into two parts: firstly, we obtained and sorted out more than seven thousand Chinese diseases and corresponding recommended and taboo food from medical and official websites; secondly, we discussed the relationship between nutritional ingredients and diseases, which mainly aims to find out which ingredients play a positive role in the rehabilitation of diseases. To the best of our knowledge, this is the first study in China, which mines the relationship between nutritional ingredients in food and diseases by using data mining technology. Experimental results showed that although we could not completely find all the positive nutritional ingredients for diseases by data mining methods, the first two or three ones were selected accurately. In addition, if our perspective can be combined with taboo food, the results would be likely to be better and in line with reality, which will be our future work direction.

There are two main benefits of this work: 1) You can access to our website\textsuperscript{2} to get diseases, recommended food, taboo food and corresponding nutrition information involved in this paper; 2) We can assist doctors and disease researchers to find out positive nutritional ingredients that are conducive to the rehabilitation of the diseases as accurately as possible. At present, some data is not available, because they are still in the medical verification. Besides, our knowledge base is still gradually improving, if researchers find out something incorrect in our work, we hope to contact us and make our research improved.

REFERENCES

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