Preparation of Total Flavonoids Tablets of Portulaca Oleracea L
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Abstract
In this study, total flavonoids of portulaca oleracea were extracted by ultrasonic method and prepared as tablets, single factor test was used to screen the optimal preparation process of tablets. Results show that, when the pressure was 7.5, the adhesive was 4% cross-linked povidone dry powder, and the excipient formula was lactose: dextrin 1:1, all the tablets obtained were in accordance with the indicators under the tablet of Chinese pharmacopoeia.

Keywords: Portulaca flavone tablets; content determination; adhesive.

INTRODUCTION
Portulaca oleracea L. is a kind of wild plant with the same origin of medicine and food. It contains a variety of medicinal components, which are mainly used in treating diarrhea, regulating blood glucose and detoxifying decrease internal heat in clinical practice. Its main components include polysaccharides, flavonoids and alkaloids [1]. Among them, the content of total flavonoids accounts for 7.67% of the total weight of purslane, which has a variety of medicinal activities [2]. As a kind of medicinal and edible homologous plant, its potential great use value, in order to make it more convenient to play a variety of value, the researchers have conducted a variety of studies on its natural pharmaceutical preparations. Currently, the dosage forms have been reported as: portulaca cinnamomum granules [3], portulaca oleracea compound oral liquid, portulaca oleracea polysaccharide granules [4] and portulaca oleracea polysaccharide injection. However, no natural pharmaceutical preparation of portulaca oleracea flavonoids has been reported. Therefore, in this experiment, portulaca oleracea flavones were prepared as functional food with tablet as carrier, laying a foundation for the research on the development of modern dosage forms of flavonoids.

MATERIALS AND METHODS
Chemical and Reagents
Rutin standard (purity≥98%) was purchased from baicaoyuan biotechnology Co., Ltd. (Jiangxi, China). Povidone K30 was purchased from sanland-chem International Inc. (USA). Lactose, dextrin, microcrystalline cellulose, starch and magnesium stearate are all biological agent provided by aubostar biotechnology Co., Ltd. (Beijing, China). Crosslinked polyvidone xl-10 purchased from Shanghai changwei pharmaceutical accessories technology Co., Ltd. (Shanghai, China). All other reagents are of analytical grade purchased from Liaoning quanrui reagent Co., Ltd. (liaoning, China).

Preparation of Flavonoids from Portulaca Oleracea L
Dried purslane, crushed into evenly sized segments. Petroleum ether was used as degreasing solvent, the ratio of material to liquid was 1:10, and ultrasonic degreasing occurs twice at 40℃, each time for 0.5h. Filter and ventilate purslane until completely dry. Set aside. 60% ethanol was used as the extraction solvent, the ratio of material to liquid was 1:30, and it was sealed and soaked overnight. Ultrasonic extraction was conducted at 50℃ for 2 times, each time for 1 hour. Filtration, collection of filtrate, concentration of 10 times volume, extraction of chloroform, ethyl acetate, n-butanol, to obtain concentrated solution of oleracea flavones. Freeze overnight at -80℃ and freeze dry for 48 h to obtain the dry powder of oleracea flavones.

Preparation of Flavonoids Tablets of Purslane
Because the tablets have the characteristics of small volume, high bioavailability, low requirements for storage conditions and stable quality, this experiment used the single-factor test method to prepare portulaca oleracea flavones as tablets, and screened the prescription process according to the quality standards.

**Tablet Preparation Process**

The raw materials and auxiliary materials were mixed evenly at a ratio of 1:1.5, then 6% cross-linked povidone was added in internal addition, 95% ethanol was wetted, and adhesive was added to prepare soft materials. Granulation was made by 12-mesh sieve, dried at 55°C for 30 min, and the grains were integrated by no. 1 sieve and no. 2 sieve, and 5% magnesium stearate was mixed evenly and immediately pressed into pieces [5].

**Screening Pressure**

Preparation pressure is one of the basic factors influencing the quality standard of traditional Chinese medicine tablets. Too much stress leads to slow disintegration; Too little pressure, produce loose pieces. Therefore, in order to ensure the quality of tablets, film pressure screening. Lactose was used as a single excipient, and the tablet appearance, weight variations, hardness and disintegration time were taken as indicators [6] to determine the optimal pressure.

**Screening Filler**

According to the properties of the flavonoids of purslane, lactose, dextrin, starch and microcrystalline cellulose (MCC) with low moisture absorption and good compressibility were selected as fillers, and the better fillers were screened by taking granulation status, difference in tablet weight, hardness, yield and disintegration time as investigation indexes.

**Screening Adhesive**

Because of the poor adhesion of the dry powder of portulaca oleracea, which affects the particle formation rate, the different adhesives were screened. The optimum adhesives were selected based on the conditions of granulation, difference in weight, hardness, yield and disintegration time.

**Determine the Excipient Formula**

Based on the above factors, the optimal excipient formula was determined.

**Determination of Flavonoids in Tablets of Purslane**

The content of total flavonoids in flavonoids tablets was determined by the content determination method of common tablet materials in 2015 edition of Chinese pharmacopoeia. Sodium nitrite colorimetry was used to prepare rutin standard solution, and the absorbance value was measured at 475 nm. The standard curve was established to calculate the content of flavonoids and the relative percentage $K$ in the tablet.

The calculation formula is as follows:

$$K(\%) = \frac{C \times V_1 \times V_2}{M \times V_3} \times 100$$

Where, $K$ is the relative percentage (%) of flavonoid raw materials in flavonoid tablets; $C$ is the sample concentration (mg/mL); $V_1$ is the volume (mL) of the tablet aqueous solution; $V_2$ is the fixed volume (mL) of the chromogenic reaction; $M$ is the weight of flavonoids (mg) of purslane; $V_3$ is the volume (mL) used for color reaction.

**RESULT**

Screening results of preparation pressure of tablets.

![Picture-1: Test results of hardness and disintegration time of film pressure](image)

In this study, the preparation pressures of tablets were 6, 7, 7.5 and 8. According to the general rule 0101 of the 2015 edition of Chinese pharmacopoeia, the appearance, weight difference, hardness and disintegration time of the tablets were tested. The results are shown in Picture-1 and Table-1. When the pressure is 7.5, all indicators are in accordance with the provisions of the national pharmacopoeia.

**Table-1: Tablet weight difference and tablet appearance test results of tablet preparation pressure (n=20)**

<table>
<thead>
<tr>
<th>Pressure</th>
<th>Weight difference</th>
<th>Tablet appearance</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>Disqualified</td>
<td>Powdery tablet, complete, uniform, matte</td>
</tr>
<tr>
<td>7</td>
<td>Qualified</td>
<td>Micronized tablet, complete, uniform and slightly shiny</td>
</tr>
<tr>
<td>7.5</td>
<td>Qualified</td>
<td>Glossy tablets, complete and smooth</td>
</tr>
<tr>
<td>8</td>
<td>Qualified</td>
<td>Glossy tablets, complete and smooth</td>
</tr>
</tbody>
</table>

**Screening Results of Tablet Filler**

Lactose, dextrin, microcrystalline cellulose and starch were selected as fillers in the experiment. The results of various indicators were shown in Picture 2 & 3 and Table-2. The results showed that all the indexes of lactose and dextrin were in conformity with the standards of Chinese pharmacopoeia.
Table-2: Tablet weight difference of filler and granulation appearance test results (n=20)

<table>
<thead>
<tr>
<th>Filler</th>
<th>Weight difference</th>
<th>Tablet appearance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactose</td>
<td>Qualified</td>
<td>The particles are easy to form, soft material too sticky, less powder</td>
</tr>
<tr>
<td>Dextrin</td>
<td>Qualified</td>
<td>Particles easy to form, adhesion, small powder, one-sided spot</td>
</tr>
<tr>
<td>MCC</td>
<td>Qualified</td>
<td>Particles are not easy to shape, size is not uniform, too much fine powder</td>
</tr>
<tr>
<td>Starch</td>
<td>Disqualified</td>
<td>Particles are not easy to form, too much fine powder, tablets soft</td>
</tr>
</tbody>
</table>

Screening results of tablet adhesives

Adhesives with different viscosity were selected in the experiment. The viscosity was successively dextrin pulp, glucose syrup, sucrose syrup, polyvidone (PVP) dry powder and polyvidone solution. The test results of various indicators are shown in picture 4 & 5, and Table-3. The results showed that the PVP powder was in accordance with the pharmacopoeia, and was more suitable as a binder for the flavonoids tablets of purslane.

Table-3: Adhesive weight difference and tablet appearance test results (n=20)

<table>
<thead>
<tr>
<th>Adhesive</th>
<th>Weight difference</th>
<th>Tablet appearance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dextrin pulp</td>
<td>Qualified</td>
<td>Powder tablets, particles easy to shape, one-sided smooth</td>
</tr>
<tr>
<td>Glucose syrup</td>
<td>Qualified</td>
<td>Powder tablets, particles easy to shape, sticky</td>
</tr>
<tr>
<td>Sucrose syrup</td>
<td>Qualified</td>
<td>Powder tablets, granules are not easy to shape and granulate</td>
</tr>
<tr>
<td>PVP dry powder</td>
<td>Qualified</td>
<td>Powder tablets, particles easy to shape, one-sided luster</td>
</tr>
<tr>
<td>PVP solution</td>
<td>Qualified</td>
<td>Glossy tablets, smooth, slightly variegated, not easy to granulate</td>
</tr>
</tbody>
</table>
Screening Results of Tablet Formulation

The results showed that the single lactose excipient had some phenomena, such as over adhesion of soft material, difficulty in granulation and low molding rate. Accordingly, match different component formula, detect each index. Dextrin can enhance the adhesion of raw materials and improve the tablet forming rate. MCC can enhance the hardness of tablets and the viscosity of neutralizing adhesives, and starch can alleviate the phenomenon that soft materials are not easy to be granulated due to the excessive adhesiveness of soft materials. The results are shown in Picture 6 & 7 and Table 4. Picture 8, Table 4, 1 is lactose: dextrin=1:1; 2 is lactose: dextrin=1:2; 3 is lactose: dextrin=2:1; 4 is lactose: MCC=2:1; 5 is lactose: starch=2:1.

![Picture 6](image6.png)  
**Picture 6: Test results of hardness and disintegration time of tablet formulation**

![Picture 7](image7.png)  
**Picture 7: Yield test results of tablet formulation**

<table>
<thead>
<tr>
<th>Tablet formulation</th>
<th>Weight difference</th>
<th>Tablet appearance</th>
<th>Comprehensive score</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Qualified</td>
<td>Lustrous tablets, particles easy to form, incomplete and complete smooth</td>
<td>98.9</td>
</tr>
<tr>
<td>2</td>
<td>Qualified</td>
<td>Glossy tablets, particles easy to shape, soft materials micro-adhesive</td>
<td>90.2</td>
</tr>
<tr>
<td>3</td>
<td>Qualified</td>
<td>Powder tablets, particles are easier to shape, soft materials too sticky</td>
<td>77.9</td>
</tr>
<tr>
<td>4</td>
<td>Qualified</td>
<td>Powder tablets, particles are not easy to form, too much fine powder</td>
<td>66.7</td>
</tr>
<tr>
<td>5</td>
<td>Disqualified</td>
<td>Powder tablets, particles are not easy to form, too much fine powder</td>
<td>60.3</td>
</tr>
</tbody>
</table>

**Table 4: Tablet formulation of the tablet weight difference and tablet appearance test results (n=20)**

Determination of tablet content

The rutin standard curve measured at 475 nm is shown in Picture 8.

![Picture 8](image8.png)  
**Picture 8: Rutin standard curve**
DISCUSSION

During the preparation of tablets, the moisture content affects the stability of drugs, so the wet-inducing characteristics of raw materials should be considered when choosing the excipients. Moreover, the raw material of flavonoids of purslane is acidic drugs, which have poor pressability. Therefore, excipients that do not react with acidic drugs, but also have low moisture absorption, high viscosity and do not delay drug disintegration should be selected. When making ordinary tablets, commonly used fillers are lactose, sucrose, starch, dextrin, pre-gelatinized starch, microcrystalline cellulose, inorganic salt, etc. [8]. Lactose has good mobility and compressibility, and there is no chemical reaction to most drugs, and the pressure of the pill is fine, beautiful and hard, and the release of the drug is faster, and the content of the main drug is less determined, and the time limit of the tablet is not prolonged [9]. Dextrin has strong viscosity and can be used in combination with ethanol to enhance the hardness of the tablet, but excessive dosage will lead to unilateral patina and slow disintegration and dissolution. At the same time, the dextrin is not soluble in cold water and ethanol, and if the content is determined, it is easy to influence the accuracy of the results. Sucrose is easily absorbent and is used for loose texture, but acidic and alkaline drugs can increase the effect of this line, so it is not suitable for the distribution. Microcrystalline cellulose is not soluble in water, ethanol and other organic solvents, soluble in alkali solution, can enhance the hardness, liquidity and disintegration of the tablet, and the amount of the tablet is better when the content is 30 percent. Starch is not soluble in cold water and ethanol, the property is very stable, hygroscopicity is small, but the compressibility is poor, often with dextrin, sugar powder with the use of tablets alone is too loose. To sum up, the use of lactose and microcrystalline cellulose alone, with the use of starch, dextrin, is more in line with the requirements of olaracea flavone tablet filler. Disintegrating agents commonly used in tablet pressing include cross-linked polyvidone, carboxymethyl cellulose and derivatives of carboxymethyl cellulose salt, etc. [10], and the amount of disintegrating agent and the way of adding have a great impact on the disintegrating rate and dissolution rate. The amount of disintegrating agent is generally 5-20% of the total tablet weight. The higher the dosage, the faster the disintegration efficiency. In this paper, we only need to make the flavone tablets of purslane as ordinary tablets, so we only need to comply with the provisions of Chinese pharmacopoeia. Therefore, we choose to add disintegrating agent inside, and the dosage is 6% of the total tablets. Because most natural drug extracts have poor compressibility, the adhesive with different viscosity is very important to the production process. The viscosity of commonly used adhesives is starch paste < dextrin paste < syrup < glue paste. In this paper, dextrin (83 g/L), glucose syrup (500 g/L), sucrose syrup (500 g/L), povidone dry powder (4%), povidone solution (20 g/L) were investigated. Therefore, lactose, dextrin, microcrystalline cellulose and starch were selected as fillers, cross-linked polyvidone was used as disintegrating agent, and 4% polyvidone dry powder was used as adhesive to prepare the flavonoids tablets of purslane.

CONCLUSION

In this paper, the preparation pressure, type and ratio of fillers, and type of adhesives of tablets were investigated by single factor test, and the optimal process was determined according to the indexes of tablets in 2015 edition of Chinese pharmacopoeia. The results showed that the optimal process of olaracea flavone tablets was: the ratio of raw and excipient was 1:1.5, and the excipient was lactose: dextrin 1:1. Adding 6% crosslinked povidone, 4% povidone dry powder, 95% ethanol wetting to prepare soft. After granulation, the tablets were dried and whole, 0.05% magnesium stearate was added, and the pressure was 7.5 to prepare the tablets.

REFERENCES


