# The study of functional ingredients from corn silk

Li Peng Ph.D.

Doctoral Thesis Summary



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## Studium funkčních přísad z kukuřičného hedvábí

### The study of functional ingredients from corn silk

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Degree programme: P2901 Chemistry and Food Technology

Degree course: 2901V013 Food Technology

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Zl ń, November, 2020

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Published by <b>Tomas Bata University in Zl ń</b> in the Edition <b>Doctoral Thesis.</b> The publication was issued in the year 2020
Key words in Czech: kukuřičné hedvábí, tepelné vlastnosti, odstraňování radikálů, fáze dospělosti, flavonoidy, steroidy, polysacharidy, kinetick émodely
Key words: corn silk, thermal properties, radical scavenging, maturity stage, flavonoids, steroids, polysaccharides, kinetic models
Full text of the doctoral thesis is available in the Library of TBU in Zl ń.

#### **SUMMARY**

The thesis studies the physicochemical and biochemical properties of the bioactive functional substances from the extracts of corn silk including flavonoids, polysaccharides and steroids, which are widely applied in the areas of food production, pharmacy, veterinary, animal feed and healthcare.

There are three main parts of the research containing 1, corn silk flavonoids, polysaccharides and steroids extraction, determination (related to the technologies of UV-VIS, HPLC, NMR, FTIR and Fluorescence excitation—emission mapping) and extraction methods optimization. 2, corn silk physicochemical property analysis including thermal analysis (related to the technology of TG/DTA), microstructure research (related to the technology of SEM). 3, corn silk flavonoids, polysaccharides and steroids biochemical property analysis including antiradical (related to the free-radicals of DPPH, ABTS and technology of EPR), anti-oxident capability (to the ferricion and copper ion), vitro enzyme inhibition activity including the enzymes of  $\alpha$ -glucosidase and  $\alpha$ -amylase (anti-diabetes), thrombin (anti-coagulation), angiotensin converting enzyme(ACE) (anti-hypertension) and xanthine oxidase(XOD) (antigout).

The whole research is based on the three different maturity stages of corn slik for the different content and proportion of flavonoids, polysaccharides and steroids in each maturity stage, which are silking stage (CS - S), milky stage (CS - M) and mature stage (CS - MS).

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#### INTRODUCTION

Corn has a widespread application as a domastic animals feed, food additives and material of alcohol through fermentated or unfermentated technology (Ivanišová et al., 2017; Michalová & Tancinov á 2017). Corn silk (CS) is the dried thrum and stigma of Zea mays L. (corn), cheap, high yielding and usually considered as a by-product to be abandoned, burned or used as fodder (Zhang, 1998). It is one of the Chinese traditional medicine recorded in many classics. According to the Southern Yunnan Material Medicine and Chinese Medicine Dictionary, corn silk is non-poisonous, also diuretic, cholagogic and resolutive. Corn silk could be used to cure many kinds of diseases clinically, such as diabetes, nephritis and hypertension etc (Jin, 1980). In addition, in terms of the results from the worldwide scientists, corn silk also has the effects of anti-fatigue (Hu et al., 2010), anti-depression (Mahmoudi & Ehteshami, 2010), anti-free radical, anti-cancer (Ebrahimzadeh, Pourmorad, & Hafezi, 2008; Z. A. Maksimović & Kovačević, 2003) and anti-radiation (Bai et al., 2010). Native American Indians usually use corn silk to cure urinary tract infection, malaria and heart disease (Hasanudin et al., 2012). In many countries, corn silk is applied to sell in markets as tea and weight-losing products for its good effect of cooling blood, purging heat and removing the damp and heat in human body.

The previous researches have successfully applied the fermentated corn fodder to improve the nutrition quality of chicken meat (Angelovicová & Semivanová 2013; Macanga et al., 2017; Štenclová et al., 2016). In addition, corn was also used to improve the sensory quality of crackers (Kuchtová, Minarovicová, Kohajdová, & Karovicová, 2016). The corn fermentated alcohol has a broad usage in food and chemistry industry, veterinary, pharmaceutical and manufacturing industry for its nutritional value and anti-oxidant properties (Krejzová et al., 2017; Süli, Hamarová, & Sobeková, 2017).

Corn silk contains many sorts of nutritional and functional ingredients, including sterols, polysaccharides, alkaloids, flavones, cryptoxanthins, polyphenols, organic acids, vitamins and allantoins etc. (Li & Lapcik, 2018) In this thesis, it will mainly research about corn silk flavonoids, polysaccharides and steroids based on the three different maturity stages which are silking stage (CSS), milky stage (CSM) and mature stage (CSMS).

#### **OBJECTIVES OF THE THESIS**

Through the research of corn silk flavonoids, polysaccharides and steroids extraction, determination (related to the technologies of UV-VIS, HPLC, NMR, FTIR and Fluorescence excitation-emission mapping) and extraction methods optimization, the basic chemical structure, content of corn silk bioactive substances can be confirmed and the optimal method of bioactive substances extraction can be built. According to the research of corn silk physicochemical property analysis including thermal analysis (related to the technology of TG/DTA), microstructure research (related to the technology of SEM), the microstructure and thermal property of corn silk powder/fiber can be known. In accordance with the research of corn silk flavonoids, polysaccharides and steroids biochemical property analysis including antiradical (related to the free-radicals of DPPH, ABTS and technology of EPR), anti-oxident capability (related to the ferricion and copper ion), Vitro enzyme inhibition activity including the enzymes of  $\alpha$ -glucosidase and  $\alpha$ -amylase (anti-diabetes), thrombin (anti-coagulation), angiotensin converting enzyme (ACE) (anti - hypertension) and xanthine oxidase (XOD) (antigout), the pharmaceutical and healthcare function of corn silk extracts can be learnt for the further research of curing diseases such as diabetes, aging, hypertension, heart disease and gout. Also based on the research of different maturity stages of corn silk, the best picking time of corn silk for pharmaceutical and nutritional use can be certain.

#### MATERIALS AND METHODS

#### 1. CS SAMPLE PHYSICOCHEMICAL PROPERTY ANALYSIS

#### 1.1 SEM analysis of the CS powder samples

In this study, the microscopic shape and size of the CS powder and fiber samples were measured by the scanning electron microscopy (SEM), the images were captured by a Hitachi 6600 FEG microscope (Japan) operating in the secondary electron mode with an accelerating voltage of 1 keV. (Will, 2020)

#### 1.2 TG/DTA analysis of the CS powder samples

In this study, thermogravimetry (TG) and differential thermal analysis (DTA) experiments were performed on simultaneous DTA–TG apparatus (Shimadzu DTG 60, Japan). Measurements were performed at heat flow rate of 5 °C/min in the static nitrogen atmosphere (gas flow of 50 ml/min) at the temperature range from 30 °C to 550 °C. The apparat was calibrated using Indium as a standard (Q. Liu, Lv, Yang, He, & Ling, 2005; Wu, Pan, Deng, & Pan, 2008).

# 2. CS functional ingredients extraction, content determination and extraction methods optimization

#### **2.1 UV-VIS**

In this study, UV/VIS spectrophotometer used was Lambda 25 (Perkin Elmer, MA, USA). Measurements were performed in the wavelength range from 200 to 700 nm in 1 cm quartz cells (Marques et al., 2013).

#### 2.2 Fluorescence Excitation-Emission Mapping

In this study, fluorescence excitation—emission maps of the different maturity stages corn silk extracts were measured on a FLS980 fluorescence spectrometer (Edinburgh Instruments, UK). Each experiment was repeated 10 times (Dankowska, 2016).

#### 2.3 CS flavonoids extraction and content determination

#### 2.3.1 Materials

All reagents and chemicals used in this research such as rutin, ethanol, sodium nitrite, aluminium nitrate and sodium hydroxide were purchased from Sigma - Aldrich (USA) in an analytical reagents purity grade. As a solvent distilled water was used. Distilled water conductivity was about  $0.6~\mu\text{S/cm}$ ).

#### 2.3.2 Rutin standard curve

Disolve 20 mg lutin into 70v.% ethanol to 50 ml (0.4 mg/ml lutin solution); separately were brought 0, 1, 2, 4, 6, 8, 10 ml 0.4 mg/ml lutin standard solutions into 50 ml volumetric flasks, added 70% ethanol 12 ml, then added 2 ml 5w.% NaNO<sub>2</sub>, shaked up and placed for 10 min to react. Then into the solutions were added 2 ml 10w.% Al(NO3)<sub>3</sub>, shaked up and placed for 10min to react, then diluted with 20 ml 10w.% NaOH to the scale of volumetric flask, placed for 5min. Each experiment was repeated 5 times. There was used 510 nm UV spectrometry to measure the absorbance of the solutions. Obtained absorbance vs. concentration dependency data were used to build up the standard curve. The numerical linear regression analysis was performed to obtain standard curve linear regression parameters. Each experiment was repeated 3 times (Peng et al., 2016).

#### 2.3.3 Determination of the CS flavonoids content

Use the 1/10 solid-liquid ratio of cornsilk powder and 70v.% ethanol to extract the flavonoids in temperatures of 40 °C and 80 °C for 20, 30, 40, 50, 60 minutes extraction time intervals. Then the flavonoids extract solutions were centrifuged on Hettich EBA 21 centrifuge (Germany) at 3000 rpm for 10 min to get the supernatant. Then there was used the same methodology as lutin standard curve to measure the flavonoids absorbance and there was used the lutin standard curve to count the given content of flavonoids. Each experiment was repeated 5 times (Peng et al., 2016).

#### 2.4 CS polysaccharides extraction and content determination

#### 2.4.1 Materials

All reagents and chemicals used in this research such as glucose, diethyl ether phenol, sulfuric acid ( $H_2SO_4$ ), ethanol, anhydrous ethanol, acetone were purchased from Sigma - Aldrich (USA) in an analytical reagents purity grade. As a solvent distilled water was used. Distilled water conductivity was about 0.6  $\mu$ S/cm).

#### 2.4.2 Glucose standard curve

Precisely weigh 100.5mg glucose which was dried to constant weight. Dilute the glucose with distilled water to 100ml to get the standard glucose solution. Take standard solution 0, 2, 4, 6, 8, 10, 12ml and dilute them with distilled water to 100ml. Suck up the above solutions 0.3ml into 10ml test tubes, then add 6% phenol solution 0.6ml, then instantly add 3ml concentrated sulfuric acid, shake up. Then place them into 40°C water bath for 30min, then put them into refrigerator and cool down for 10min. Then use UV spectrometry to measure their absorbance in 490nm. Use absorbance (A) as Y axil, mass concentration (C) ( $\mu$ g/ml) as X axil to get the regression equation (Hossain et al., 2014).

#### 2.4.3 Determination of the CS polysaccharides content

Use diethyl ether to degrease the CS powder sample in room temperature for 12 hours, then use 36°C constant temperature drying box to dry to the degreased corn silk powder, then weigh 50g degreased power into round-bottom flask and add 750ml distilled water, then put the round-bottom flask into 100°C water bath for 2 hours. Then centrifuge the sample in the speed of 10000 r/min for 10min to get the supernatant and sediment, then use the sediment to repeat the above hot water extraction and centrifugation process and merge twice centrifugal supernatant,

then vacuum concentrate, then add 70% ethanol into concentrated solution to precipitate, then use anhydrous ethanol, acetone and ether to ordinally wash the sediment, then dry the sediment with 50°C hot air, then get the corn silk polysaccharides. Use the same method as the standard glucose curve to measure the UV absorbance, then use the standard glucose curve to calculate the content of polysaccharides. Each experiment was repeated 5 times (Hossain et al., 2014).

#### 2.5 CS steroids extraction and content determination

#### 2.5.1 Materials

All reagents and chemicals used in this research such as  $\beta$ - sitosterol, ethanol, phosphoric acid, sulfuric acid and ferric chloride were purchased from Sigma-Aldrich (USA) in an analytical reagents purity grade. As a solvent distilled water was used. Distilled water conductivity was about 0.6  $\mu$ S/cm).

#### 2.5.2 β - sitosterol standard curve

Precisely weigh  $10 \text{ mg }\beta$  – sitosterol into 10 ml volumetric flask, use absolute ethyl alcohol dilute to scale. Fetch 1ml above solution into 10ml volumetric flask, use absolute ethyl alcohol to dilute to scale as the standard sample solution. Precisely move the standard sample solution 0, 1, 2, 3, 4, 5 ml into 50 ml conical flask, separately add 5, 4, 3, 2, 1, 0 ml absolute ethyl alcohol, then slowly pour the pulfate – phosphate - ferric reagent 5 ml long the cup wall into every conical flask separately, shake up, cool down in room temperature for 20 min. Then was measured the 530 nm absorbance by UV spectrometry. Each experiment was repeated  $5 \times 0.0 \text{ ml}$ . Obtained absorbance vs. concentration dependency data were used to build up the standard curve. The numerical linear regression analysis was performed to obtain standard curve lineasr regression parameters. Each experiment was repeated  $3 \times (\text{Hossain et al.}, 2014)$ .

#### 2.5.3 Determination of the CS steroids content

Weigh 5 samples 3 g corn silk powder, add 70 % ethanol (material: liquid = 1:20), then use 200 W ultrasonic extract for 15 min, 30 min, 45 min, 60 min, 75 min for the silking, milky and mature stages. Then was used 119 W microwave extraction apparatus for 8 min, followed by addition of pulfate- phosphate- ferric reagent to process for 20 min (the same procedure as for standard curve determination), then measure the 530 nm absorbance by UV spectrometry. Then use the standard curve to calculate the content of steroids from the CS samples. Each experiment was repeated  $5 \times$  (Hossain et al., 2014).

#### 3. BIOCHEMICAL PROPERTY ANALYSIS OF CS EXTRACTS

#### 3.1 Radical scavenging effect of CS extracts determination

Free radical reaction, also known as free radical reaction, is a variety of chemical reactions involving free radicals. The outer layer of the radical electron shell has an unpaired electron, which has a strong affinity for adding a second electron, so it can act as a strong oxidant. The more important one in the atmosphere is the OH-radical, which can react with various trace gases. In the chemical reaction formed by photochemical smog, there are many free radical reactions, which play an important role in the initiation, transmission and termination processes in the chain reaction. Many free radicals are intermediate products, such as hydrogen peroxide radicals (HO2-), alkoxy radicals (RO-), peroxyalkyl radicals (RO2-), acyl radicals (RCO-), etc. (Reddy, 2020)

#### 3.1.1 Materials

All reagents and chemicals used in this research such as DPPH, ABTS, potassium persulfate  $(K_2S_2O_8)$ , ethanol, methanol, copper chloride, hydrogen peroxide  $(H_2O_2)$ , 5,5-dimethyl-1-pyrroline N-oxide (DMPO), sodium phosphate buffer, potassium ferricyanide  $[K_3Fe(CN)_6]$ , trichloroacetic acid, ferric chloride  $(FeCl_3)$ , neokuproiny, ammonium acetate  $(CH_3COONH_4)$  were purchased from Sigma-Aldrich (USA) in an analytical reagents purity grade. As a solvent distilled water was used. Distilled water conductivity was about  $0.6~\mu S/cm$ ).

#### 3.1.2 EPR spin - trapping measurement of the CS extracts

The thermal decomposition of copper chloride (CuCl<sub>2</sub>) in H<sub>2</sub>O<sub>2</sub> at 333 K was used as reactive radicals to test the radical scavenging capability (RSC) of cornsilk extracts, the EPR spintrapping technique was used 5,5-dimethyl-1-pyrroline N-oxide as a spin trap. Chloride radical anions (Cl•) generated upon thermal decomposition of CuCl<sub>2</sub> represent reactive species with high reduction potential, capable to react with organic compounds, the paramagnetic species in H<sub>2</sub>O<sub>2</sub> solvent were added to the double bond of DMPO spin trapping agent to produce the corresponding spin adducts. All EPR measurements were operated in a 4-mm flat quartz cell in a Bruker TE102 (ER 4102 ST) cavity, using the EMX EPR spectrometer (Bruker, Germany) working in the X-band. The ER 4111 VT temperature unit (Bruker, Germany) was used for temperature regulation. The reaction mixture contained 100 µL H<sub>2</sub>O<sub>2</sub> extracts (pure H<sub>2</sub>O<sub>2</sub> as reference), 100 µL H<sub>2</sub>O<sub>2</sub>, 25 µL of 0.2 M DMPO dissolved in H<sub>2</sub>O<sub>2</sub> and 25 µL of 0.01 M CuCl<sub>2</sub> (H<sub>2</sub>O<sub>2</sub>). The time course of EPR spectra of the DMPO spin adducts was recorded in 66-s intervals for 22 min at 333 K (each spectrum represents an accumulation of three scans). The integral EPR intensity (double integral) found after 22 min of thermal treatment for the sample solution was compared with the reference measurement. The difference between the integral EPR intensities of the reference and the samples in 22nd min characterises the amount of radicals scavenged by the various components present in the sample acting as radical scavengers. The RSC values were calculated as a percentage of scavenged radicals relative to the reference sample (H<sub>2</sub>O<sub>2</sub>). These values were recalculated to trolox equivalent antioxidant capacity (TEAC) using calibration curve measured analogously for trolox solutions in CuCl<sub>2</sub>/DMPO/H<sub>2</sub>O<sub>2</sub> systems, then the obtained radical scavenging characteristics of investigated samples were evaluated in 1 mol of trolox/1 g of extract (Yamasaki & Grace, 1998).

#### 3.1.3 DPPH radical scavenging activity measurement

Separately put 1 mL CS-S, CS-M, CS-MS corn silk extracts into 3 mL  $1.44 \times 10^{-4}$  mol/L DPPH radical, use same concentration vitamin C and lutin as comparison groups, 4 mL  $1.44 \times 10^{-4}$  mol/L DPPH without scavenger as the control group to determine the 516 nm absorbance in different reaction time. Determine the dynamic relation between absorbance and time along with the reaction between the different scavengers and DPPH radical and use absorbance as the Y - axis, time as X - axis to draw the dynamic relation curves, the inhibition level of DPPH was calculated by the following equition:

Inhibition = 
$$[(A_0 - A_1)/A_0] \times 100\%$$

where  $A_0$  is the 516 nm absorbance of the control group at the initial time (0 min),  $A_1$  is the 516 nm absorbance of the scavenger sample at the ending time (Masek, Chrzescijanska, Latos, Zaborski, & Podsedek, 2017).

#### 3.1.4 ABTS radical scavenging activity measurement

Mix 17,2 mg ABTS, 3,3 mg  $K_2S_2O_8$  and 5 ml  $H_2O$  as oxidation starter, radicals are formed and oxidized after 24 hours and then kept in freezer (-6 °C), then add 60 ml of water to 1 ml of oxidized solution of ABTS as the solution of ABTS to test the CS extracts radical scavenging activity. Then mix 2.5ml ABTS solution with 0.5ml CS-S, CS-M, CS-MS CS extracts solution respectively (each maturity stage CS sample was extracted for 60 min and 90 min), use VC (the same concentration as the mean concentration of all of the CS extracts samples) as comparison, water as reference and determine the 734 nm absorbance in different time. Determine the dynamic relation between absorbance and time along with the reaction between the different scavengers and ABTS radical and use absorbance as the Y - axis, time as X - axis to draw the dynamic relation curves , the inhibition level of ABTS was calculated by the following equition:

Inhibition = 
$$[(A_0 - A_1)/A_0] \times 100\%$$

where  $A_0$  is the 734 nm absorbance of the control group at the initial time (0 min),  $A_1$  is the 734 nm absorbance of the scavenger sample at the ending time (Masek et al., 2017).

#### 3.1.5 Ferricion reducing anti-oxidant power

Separately put 1 mL CS-S, CS-M, CS-MS CS extracts solutions into 2.5 mL, 0.2 M, pH 6.6 phosphate buffer and 2.5 mL, 1% K<sub>3</sub>Fe(CN)<sub>6</sub>. Then incubate the mixture at 50°C for 20 minutes. Then terminate the reaction by adding 2.5 mL, 10% trichloroacetic acid. Then mix the upper layer of 2.5 mL solution with 2.5 mL distilled water and 0.5 mL, 0.1% FeCl<sub>3</sub>, then measure the 700 nm absorbance of the reaction mixture against a blank sample. The increased absorbance of the reaction mixture represents the reducing power (Masek et al., 2017).

#### 3.1.6 Copper ion reductive capability

The base of this assay is Cu(II) to Cu(I) reduction. 0.25 mL,  $0.01 \text{ M CuCl}_2$  was mixed with 0.25 mL,  $7.5 \times 10^{-3} \text{ M}$  neokuproiny ethanol solution and 0.25 mL,  $1 \text{ M CH}_3\text{COONH}_4$  buffer solution. Then add CS-S, CS-M, CS-MS CS extracts solution 0.25 mL respectively, then volume the mixture to 2 mL by distilled water and maintain at the room temperature for 30 minutes. Then measure the 450 nm absorbance against the blank sample (0.25 mL water). The increased absorbance represents the copper ion reductive capability (Masek et al., 2017).

# 3.2 CS extracts bioactive ingredients, functional groups and molecuar chemical structures analysis and determination

#### 3.2.1 Materials

All reagents and chemicals used in this research such as methanol, glacial acetic acid, acetonitrile were purchased from Sigma-Aldrich (USA) in an analytical reagents purity grade. As a solvent distilled water was used. Distilled water conductivity was about  $0.6~\mu S/cm$ ).

#### 3.2.2 NMR analysis

In this study, the methanolic CS-S, CS-M, CS-MS CS extracts were put into 5 mm NMR (Bruker, Germany) tubes and measured on a 600 MHz NMR spectrometer equipped with a VNMRS console and PFG Dual Broadband DBG600-5F probe (Varian/Agilent). After initial tuning, matching, locking, and shimming, the 90-degree pulse was calibrated and transmitter offset was set to residual water signal frequency. The tnnoesy pulse sequence was used with 1 s relaxation delay (including 0.99 s water presaturation), 100 ms mixing time (with water presaturation) and 4 s acquisition time (Larina, 2019).

#### 3.2.3 FTIR analysis

In this study, the FTIR spectra were recorded on a FTIR spectrometer (Thermo Electron Scientific Instruments Corporation, USA) in the region of 4000 - 400 cm<sup>-1</sup> (Mohamed, Jaafar, Ismail, Othman, & Rahman, 2017).

#### 3.2.4 HPLC analysis

In this study, absolute methanol extracted corn silk flavonoids (extraction temperatures were 60°C, 70°C and 80°C) were filtrate by 0.2μm filter membrane, stored in 4°C, then analyzed by UHPLC Dionex Ultimate 3000, UV / VIS detector, DAD (Thermo Fisher Scientific, USA), conditions of HPLC are Phenomenex Kinetex C18 chromatographic column (150mm x 4.6 mm; 5 μm); mobile phase A : 99/1: water / glacial acetic acid 99.8%; mobile phase B: 67/32/1: water / acetonitrile / glacial acetic acid 99.8%; flow rate: 1ml/min; determine wavelength: 275nm; column temperature: 30°C; injection volume: 10μl; external standard method; processing time: 45min; mobile phase gradient: gradient elution: 0 - 10 min 10 - 20% B,10 - 16 min 20 - 40% B, 16 - 20 min 40 - 50% B, 25 - 25 min 50 - 70% B, 25-30 min 70% B, 30 - 40 min 70 - 10% B, 40 - 45 min 10% B. (Lozano-S ánchez, Borr ás-Linares, Sass-Kiss, & Segura-Carretero, 2018)

#### 3.3 Vitro Enzyme Inhibition Test

The vitro enzyme inhibition test refers to the simulation of the external animal and plant environment in vitro, which is based on the enzyme inhibitor and the enzyme reaction kinetics. It has a wide range of uses in biochemistry, pharmaceuticals and medicine. (Deng et al., 2019) **3.3.1 Materials** 

All reagents and chemicals used in this research such as 10% fetal bovine serum, dimethyl sulfoxide (DMSO), 5% thiazolyl blue tetrazolium bromide (MTT), phosphate buffer (PBS), Tris-HCl, hydrochloric acid (HCl), Sodium carbonate (Na<sub>2</sub>CO<sub>3</sub>) boric acid buffer, P – nitroglucoside (pNPG) thrombin, fibrin, angiotensin converting enzyme, hippuryl histidine leucine,  $\alpha$ - glucosidase,  $\alpha$ - amylase, xanthine, xanthine oxidase (XOD), uric acid and captopril were purchased from Sigma-Aldrich (USA) in an analytical reagents purity grade. As a solvent distilled water was used. Distilled water conductivity was about 0.6  $\mu$ S/cm).

#### 3.3.2 Thrombin inhibition activity test

Put 40  $\mu$ L CS flavonoids, polysaccharides and steroids extracts into 140  $\mu$ L 0.1% fibrinogen, place for 10 min, then add 10  $\mu$ L thrombin solution, shake up, then place in 37°C for 2 min, then measure the 405 nm absorbance (A<sub>405</sub>). Use 0.05M Tris - HCl (pH 7.2) buffer instead of thrombin as the blank group. The thrombin inhibition rate equation is as follows:

$$Y=(A_1-A_2)/(A_1-A_0) \times 100\%$$

where Y is the thrombin inhibition rate,  $A_1$  is the  $A_{405}$  without inhibitor,  $A_2$  is the  $A_{405}$  with inhibitor and  $A_0$  is the  $A_{405}$  of the blank group (Whelihan, Kiankhooy, & Brummel-Ziedins, 2014).

#### 3.3.3 ACE inhibition activity test

Put  $10\mu L$  CS flavonoids, polysaccharides and steroids extracts into  $30\mu L$  hippuryl histidine leucine as the substrate, then place the mixed solution  $37^{\circ}C$  thermostatic waterbath for 5 min, then add  $20\mu L$  ACE in  $37^{\circ}C$  temperature for 30 min, then add  $60\mu L$  1mol/L HCl to terminate the reaction. Use 228nm absorbance (A<sub>228</sub>) to calculate the ACE inhibition rate. Use captopril with pH 8.3 boric acid buffer plus CS extracts sample solutions as the blank group. The ACE inhibition rate equation is as follows:

$$Y = [(M-N)/M]X100\%$$

where Y is the ACE inhibition rate, M is the blank group  $A_{228}$ , N is the  $A_{228}$  of the CS sample extracts groups (Sawicki et al., 2019).

#### 3.3.4 α- glucosidase inhibition activity test

Put  $5\mu L$   $\alpha$ - glucosidase (10 unit/mL 0.1mol/L pH 6.8 PBS) solution and  $10\mu L$  CS extracts samples into  $620\mu L$  0.1mol/L PBS, place the mixed solution in  $37.5^{\circ}C$  for 20 min, then add  $10\mu L$  10mmol/L pNPG as the substrate to initiate the reaction. After 30 min reaction in  $37.5^{\circ}C$ , add  $650\mu L$  1mol/L  $Na_2CO_3$  to terminate the reaction. Use  $410_{nm}$  light to measure the absorbance (A<sub>410</sub>). Use the above mixed solution without CS extracts as the blank group. The  $\alpha$ - glucosidase inhibition rate calculation equation is as follows:

$$Y = [(A_0 - A_1)/A_0] \times 100\%$$

where Y is the  $\alpha$ - glucosidase inhibition rate,  $A_0$  is the blank group  $A_{410}$  and  $A_1$  is the CS extracts groups  $A_{410}$ . (Kato-Schwartz et al., 2020)

#### 3.3.5 $\alpha$ - amylase inhibition activity test

Put  $10\mu L$   $\alpha$ - amylase (1 unit/L) into  $10\mu L$  CS extracts solutions, place for 15 min, then add  $500\mu L$  1% starch (PBS) solution to initiate the reaction. After 5 min reaction in 37.5°C, add  $600\mu L$  DNS reagent to terminate the reaction. Then put the above reacted solution in boiling water bath for 15 min, then cool down to the room temperature. Use 540 nm light to measure the absorbance (A<sub>540</sub>) and use the group without CS extracts as the blank. The  $\alpha$ - amylase inhibition rate calculation equation is as follows:

$$Y = [(A_0 - A_1)/A_0] \times 100\%$$

where Y is the  $\alpha$ - amylase inhibition rate,  $A_0$  is the blank group  $A_{540}$  and  $A_1$  is the CS extracts groups  $A_{540}$ . (Kato-Schwartz et al., 2020)

#### 3.3.6 XOD inhibition activity test

Mix  $300\mu L$  CS extracts solutions,  $210\mu L$  pH 7.5 PBS and  $180\mu L$  0.01unit/mL XOD solution (pH 7.5 PBS as the solvent) in  $25\,^{\circ}$ C for 15 min, then add  $360\mu L$  1.5M/mL xanthine solution (pH 7.5 PBS as the solvent) to react in  $25\,^{\circ}$ C for 30 min. Use the 150mL 1mol/L HCl as the

terminator, measure the 290nm absorbance ( $A_{290}$ ) to calculate the XOD inhibition rate. Use the group adding HCl before XOD as the blank. The XOD inhibition rate calculation equation is as follows:

$$Y = [(A_0 - A_1)/A_0] \times 100\%$$

where Y is the XOD inhibition rate,  $A_0$  is the blank group  $A_{290}$  and  $A_1$  is the CS extracts groups  $A_{290}$ . (Li et al., 2018)

#### RESULT AND DISCUSSION

#### 1. CS SAMPLE PHYSICOCHEMICAL PROPERTY ANALYSIS

#### 1.1 SEM analysis of the CS powder samples

SEM images of the tested corn silk powders and fibers are characteristic with the rectangular shape of individual particles exhibiting complex microporous structure on the intersection. Such structures are typical for plants cellulose based materials.

#### 1.2 TG/DTA analysis of the CS powder samples

The moisture content and thermoal analysis by TG and DTA of the samples were resulted as the typical multistep decomposition process for all CS- S, CS- M and CS- MS samples. The first step decomposition of CS-S was in the temperature range from 30 to 120 °C with observed weight loss 8.3 % attributed to the moisture content. Total decomposition step was about 77.45 % in the temperature range of 30 to 550 °C. Similarly, for the sample CS-M and CS-MS, TG data exhibited the first step decomposition of 5.9 % and 10.04 % in the same temperature range as CS-S followed by the total weight loss of 65.5 % and 83.88% in the temperature range of 30 to 550 °C, which indicated the CS-MS contains the most thermally labile substances compared with CS-S and CS-M. There are two endothermic peaks. The first one was located in the temperature of 54.3 °C for CS-S and CS-MS, 60.2 °C for CS-M attributed to the melting point of flavonoids (Miziara et al. 2017). The second one was observed at 397 °C (CS-MS), 415.1 °C (CS-M) and 419.7 °C (CS-S) attributed to the total thermal decomposition with the formation of a low quantity carbonaceous residues respectively. Observed exotheric process at 524 °C corresponds to the decomposition CS - MS sample.

## 2. CS FUNCTIONAL INGREDIENTS EXTRACTION, CONTENT DETERMINATION AND EXTRACTION METHODS OPTIMIZATION

#### 2.1 CS flavonoids extraction and content determination

#### 2.1.1 Rutin standard curve

Typical lutin standard curve as obtained according to the standard procedure described in detail in the materials and methodology section. Obtained data were of a high correlation as indicated by the correlation coefficient being 0.995.

#### 2.1.2 Determination of the CS flavonoids content and extraction methods optimization

Effects of the extraction time and of the extraction temperature are summarized in Figure 1, where kinetics data of the extraction of CS-S, CS-M and CS-MS samples are plotted in flavonoids concentration vs. extraction time coordinates. Each extraction was performed at two different temperatures, the first one was of 40 °C and the second one at 80 °C. All of the CS-S, CS-M and CS-MS dependencies were of a complex non-linear character, modeled as a third order polynomial dependency. However, in the case of CS-S, for both extraction temperatures the same characteristic sinusoidal pattern was found exhibiting the maximum extraction efficiency at 50 °C. Observed concentrations were of about  $6.5 \times 10^{-3}$  mg/ml for CS-S extracted at 40 °C, about  $5 \times 10^{-3}$  mg/ml for CS-MS extracted at 80 °C indicates that the extracted flavonoids substances are sensitive on thermal history and are undergoing thermal decomposition similarly as reported by Chaaban et al.

(Chaaban et al., 2017). In the latter paper, the linear degradation pattern was found for rutin at 70 °C degradation temperature. With increasing degradation temperature up to 130 °C the exponential decay pattern was found. In the case of CS-M, the maximum extraction efficiency was found at 30 min extraction time being of  $5.8 \times 10^{-3}$  mg/ml at 40 °C extraction temperature. However, at higher temperature, the degradation of the flavonoids content was observed, and only the exponential decay pattern was found as indicated in Figure 4B. That is why, the maximum extraction was found at 20 min extraction time at 80 °C extraction temperature. It was found, that the highest extracted flavonoids content was  $(7.2 \pm 0.3) \times 10^{-3}$  mg/ml for CS-M 80 °C sample and  $(12.2 \pm 0.4) \times 10^{-3}$  mg/ml for CS - MS 80 °C sample. However, for the CS-S the highest content was found of  $(6.8 \pm 2.1) \times 10^{-3}$  mg/ml for CS-S 40 °C sample. To characterize obtained extracts of flavonoids, the UV VIS as well as fluorescence spectra were measured, prior to the fluorescence mapping analysis of the studied corn silk extracts, the UV VIS spectra were recorded.

These were typical with three major absorption regions at 260nm and 360 nm (near ultraviolet region), and at visible light region of 490 nm. The absorption of electromagnetic radiation in the near ultraviolet region is typical for poly-unsaturated and aromatic compounds such as flavonoids. All of CS-S, CS-M and CS-MS exhibited similar UV VIS spectra except the visible range region, where a 480 nm shoulder peak occurred for CS-M sample.

Results of the fluorescence excitation-emission mapping of the studied extracts are characteristic similarly as the UV VIS absorption spectra with the three distinct fluorescence emission regions at 320 nm, 450 nm and 680 nm. Emission region located at 450 nm was ascribed to the flavonoids compounds similarly as observed by Shan et al. (Shan et al. 2017), who found, that the flavonois characteristic excitation/emission spectral range is  $365 - 390 \, \text{nm}/450 - 470 \, \text{nm}$ . The excitation/emission spectral range of  $480 - 500 \, \text{nm}/510 - 520 \, \text{nm}$  was ascribed to flavanols.

There were found three distinct excitation wavelengths regions at about 275nm, 350 nm and 420 nm. Obtained results indicate, that the major difference among CS-S, CS-M and CS-MS is in the fluorescence emission centered at the 450 nm region, where the intensity of the fluorescence emission was highest for CS-S extracted at 80  $^{\circ}$ C for 50 minutes. Furthermore, there was found that the fluorescence emission intensity region located at 450 nm region was of higher intensity for CS-S in comparison to CS-M and CS-MS at 80  $^{\circ}$ C extraction temperature. Observed results were considered as statistically significant (p $\leq$ 0.05). These results are in an excellent correspondence with the TG analysis, where the first step decomposition weight loss was found higher for CS-S in comparison to CS-M and CS-MS. However, there was not found any major difference between fluorescence emission intensities located at 320 nm region for all studied materials.

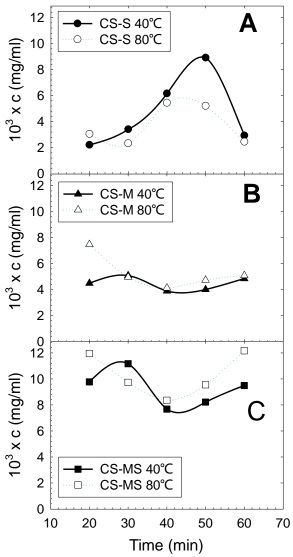


Figure 1. Flavonoids extraction kinetics: A – corn silk silking stage, B – corn silk milky stage, C – corn silk mature stage.

#### 2.2 CS polysaccharides extraction and content determination

#### 2.2.1 Glucose standard curve

There is shown a typical lutin standard curve as obtained according to the standard procedure described in detail in the materials and methodology section. Obtained regression parameters are given. Obtained data were of a high correlation as indicated by the correlation coefficient being 0.995.

# 2.2.2 Determination of the CS polysaccharides content and extraction methods optimization

As can be seen in Figure 2, within the same extraction temperature of  $100^{\circ}$ C, the all of the CS-S, CS-M and CS-MS polysaccharides concentrations have the positive correlation with extraction time and all of the 3 maturity stages showed the similar trends of the kinetics, which is from 80-100 min, the polysaccharides content increase trends are sheer and from 100-120 min, the trends became subdued. The maximum polysaccharides extraction content for CS-S sample was  $0.12~\mu g/ml$  in the 120~min as well as  $0.16~\mu g/ml$  for CS-M and  $0.33~\mu g/ml$  for CS-

MS. It is notable that the CS-MS has a much higher polysaccharides yield than CS-S and CS-M. On the contrary, the CS-S has the lowest yield of polysaccharides.

Figure 3 shows, within the same extraction time 2 hours, all of the CS-S, CS-M and CS-MS polysaccharides concentrations have the positive correlation with the extraction temperature. CS-M and CS-MS have the similar trend of the temperature vs. polysaccharides concentration kinetics, which is 60-80 °C, the increasing was sheer and from 80-100 °C, the trend became subdued. The CS-S polysaccharides extraction kinetic showed a potential of increase even in the temperature higher than 100 °C, however the totally trend of CS-S polysaccharides yield increase from 60-100 °C was subdued compared with the trend of CS-M and CS-MS. It is remarkable that the CS-MS yield was much higher than the CS - S and CS-M, which is in accord with the result of Figure 2.

The CS-S, CS-M and CS-MS polysaccharides extracts 200-700 nm wavelength absorbance was also determined by UV VIS and they all showed the similar curves of the light absorbance. The apparent peaks are in the wavelength range of 470-490 nm, which is the typical absorption wavelength of polysaccharides.

Therefore, in accordance with the Figure 2 – Figure 3 results, the optimal extraction method of CS polysaccharides extraction conditions are 100 °Cextraction temperature, 2 hours extraction time with the CS-MS sample.

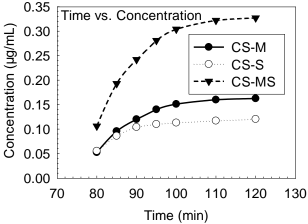


Figure 2. Within 100°C extraction temperature, the CS-S, CS-M and CS-MS polysaccharides extraction time vs. concentration kinetics

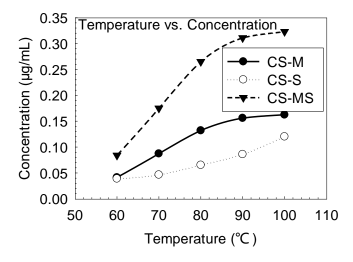


Figure 3. Within 2 hours extraction time, the CS-S, CS-M and CS-MS polysaccharides extraction temperature vs. concentration kinetics

#### 2.3 CS steroids extraction and content determination

#### 2.3.1 \( \beta \)- sitosterol standard curve

The  $\beta$ - sitosterol standard curve is shown the regression parameters. The Obtained data were highly correlated as the correlation coefficient 0.999.

#### 2.3.2 Determination of the CS steroids content and extraction methods optimization

Figure 4 shows the effects of the Ultrasonic time and different maturity stages by the steroids concentration vs. ultrasonic time correlations. All of the CS-S, CS - M and CS-MS were a non – linear character, modeled as a third order polynomial dependencies. However, all of those three stages extraction contrations had a direct proportionality trend with the increase of the ultrasonic processing time. All of the three stages samples had an obvious increasing range of the extraction concentration from ultrasonic extraction time 15 min to 60 min. From 60 min to 75 min, three samples showed a similar steady tendency which means from 60 min to 75 min the increase of the concentration is not conspicuous anymore. Therefore, the 75 min ultrasonic extraction time can be marked as the optimum extraction time to obtain maximum extracted content for all of the three stages. Obtained maximum concentrations were as follows: CS-S 0.09 mg/ml, CS-M 0.12 mg/ml, CS-MS 0.37 mg/ml. It is noteworthy, that the CS-MS had a much higher maximum extraction concentration as well as the increasing rate in comparison to CS-S and CS-M samples. This imply that the CS-MS has much higher content of steroids than CS-S and CS-M. Simultaneously, the ultrasonic assisted technique can be sonsidered to be much more effective to CS-MS sample extraction rather than for CS-S and CS-M., which is in agreement with the results of UV-VIS and fluorescence excitation – emmision mapping.

The UV VIS and fluorescence spectra were the typical three major light absorption regions at near ultraviolet region of 350 nm and visible light region of 500 nm and 650 nm. The electromagnetic radiation absorption in the visible light region is typical for anthraquinone and phenanthrene compounds such as steroids. All of the CS-S, CS-M and CS-MS exhibited similar UV VIS spectra.

The fluorescence excitation-emmision mapping result of the CS extracts are similar as the UV VIS measurement, there were three distinct fluorescence enission regions at 300 nm, 430 nm and 680 nm respectively.

There were three distinct excitation wavelengths regions at about 275nm, 350 nm and 380 nm, which indicate the difference among CS-S, CS-M and CS-MS is in the fluorescence emission centered at the 300 nm and 430 nm regions. The highest intensity of the fluorescence emission at 300 nm was found for CS-MS extracted at 40  $^{\circ}$ C for 15 minutes in the ultrasonic extraction bath. Moreover, the fluorescence emmision intensity region at 430 nm region was the highest intensity for CS-MS as well. Observed results were considered as statistically significant (p $\leq$ 0.05). However, there was no any major difference for all studied materials at the fluorescence emmission intensity of 670 nm region.

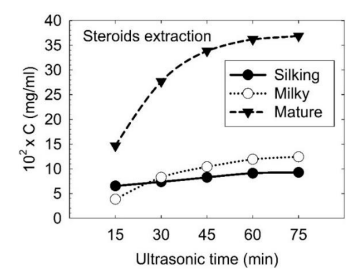


Figure 4. Ultrasonic assisted CS steroids extraction kinetics

#### 3. BIOCHEMICAL PROPERTY ANALYSIS OF CS EXTRACTS

#### 3.1 Radical scavenging effect of CS extracts determination

#### 3.1.1 EPR spin - trapping measurement of the CS extracts

As can be seen in Figure 5, the extracts of CS with 70% ethanol has much stronger radical scavenging capability than water extracts and both water and 70% ethanol extracts show the similar difference according to the maturity stages of CS, which is the CS-MS extracts have the strongest radical scavenging capability, then CS-S medium, CS-M the weakest. Compared with water, 70% ethanol is a more effective solvent to extract more radical scavengers from CS but also more influential to the DMPO radical since the reducibility of the ethanol itself. Therefore, the afterwards experiments applied the 70% ethanol as the solvent.

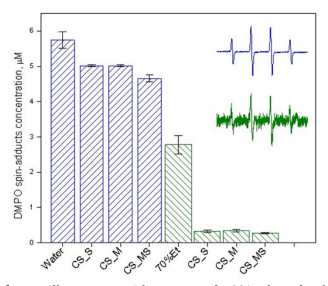


Figure 5. EPR test of cornsilk exrtacts with water and 70% ethanol solvent in CS-S, CS-M, CS-MS

#### 3.1.2 DPPH radical scavenging activity measurement

Figure 6 - 8 illustrate the effect of DPPH scavenging from the strongest to the lightest is vitamin C, rutin and CS extracts according to the time and A<sub>516</sub>, all of vitamin C, rutin and CS extracts have the prominent scavenging effect from 0 to 5 min and from 5-60 min the effect became mitigatory gradually. The scavenging effect of vitamin C stopped after 5 min means vitamin C had consumed all of the DPPH radical in the mixed solution. The same pattern of CS extracts and rutin also can be seen in Figure 6 - 8, and table 1, along with the higher

concentration of the anti-oxidant materials, comes the stronger effect of DPPH scavenging. Table 1 shows the DPPH radical scavenging capability of the CS-S, CS-M, CS-MS extracts and the corresponding same concentration VC and rutin. CS-S extracts had a higher capability of DPPH radical scavenging than the CS-M but almost the same. Comparatively, CS-MS extracts had a much higher capability of DPPH radical scavenging than CS-S and CS-M. The result is also matched the result of EPR measurement.

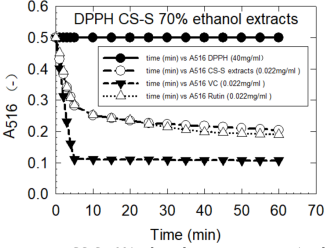


Figure 6. DPPH CS-S 70% ethanol extracts scavenging kinetics

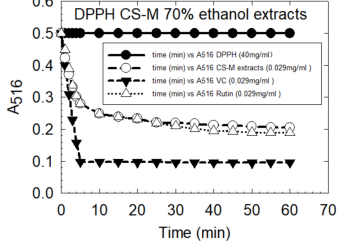


Figure 7. DPPH CS-M 70% ethanol extracts scavenging kinetics

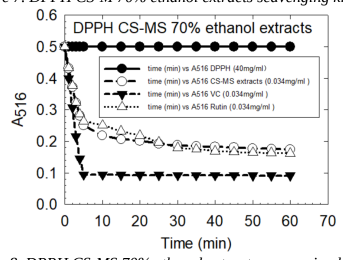


Figure 8. DPPH CS-MS 70% ethanol extracts scavenging kinetics

Table 1. The inhibition of the DPPH radical

Scavenger	Inhibition (%)
CS-S (0.22 mg/mL)	59.33 ±0.61
VC (0.22 mg/mL)	$78.63 \pm 0.45$
Rutin (0.22 mg/mL)	$62.03 \pm 0.35$
CS-M (0.29 mg/mL)	59.20 ±0.92
VC (0.29 mg/mL)	$80.63 \pm 0.96$
Rutin (0.29 mg/mL)	$62.30 \pm 0.76$
CS-MS (0.34 mg/mL)	$65.20 \pm 0.90$
VC (0.34 mg/mL)	$81.63 \pm 0.67$
Rutin (0.34 mg/mL)	$67.30 \pm 1.08$

#### 3.1.3 ABTS radical scavenging activity measurement

As can be seen from Table 2, along with the longer extraction time 90 min it showed a stronger capability of ABTS radical scavenging in each maturity stage CS extracts sample. The same pattern as in DPPH scavenging, the capability of different maturity stages CS ABTS scavenging rank from the strongest to the weakest was also CS-MS, CS-S and CS-M. Also all of the samples had an apparent decreasing trend in the beginning 5 min and then the trends became placid. It is noticeable that the ABTS radical scavenging capability was getting remarkably discriminatory according to the maturity stages especially the difference between the CS-S and CS-M compared with the DPPH radical scavenging measurement and analysis. Additionally, the vitamin C as the comparison group, showed a much stronger radical scavenging capability than the CS extracts samples.

Table 2. The inhibition of the ABTS radical

Scavenger	Inhibition (%)
CS-S (60 min)	$19.21 \pm 0.55$
CS-S (90 min)	$25.68 \pm 0.60$
CS-M (60 min)	$14.52 \pm 0.60$
CS-M (90 min)	$16.41 \pm 0.65$
CS-MS (60 min)	$65.46 \pm 0.72$
CS-MS (90 min)	$69.17 \pm 0.56$
VC	$93.28 \pm 0.45$

#### 3.1.4 Ferricion reducing anti-oxidant power and copper ion reductive capability

Table 3 and table 4 show the anti-oxidant capability of different maturity stages CS extracts in the reduction reaction with Fe<sup>3+</sup> and Cu<sup>2+</sup>. For both Fe<sup>3+</sup> and Cu<sup>2+</sup>, the different maturity stages CS extracts were capable to reduce the Fe<sup>3+</sup> and Cu<sup>2+</sup> by it anti-oxidant property. The among the three stages of CS, CS-MS extracts had a much stronger anti-oxidant effect than CS-S and CS-M, CS-M had the weakest anti-oxidant capability, which is also consist with the result of EPR, DPPH and ABTS measurement. However, CS-S extracts showed a remarkably higher anti-oxidant effect than the CS-M to the Fe<sup>3+</sup> compared with the similarities between CS-M and CS-S in the previous EPR, DPPH and ABTS assays.

Table 3. Ferricion reducing anti-oxidant power

CS extracts samples	Ferricion reducing power
CS-S	$1.33 \pm 0.32$
CS-M	$0.53 \pm 0.11$
CS-MS	$2.63 \pm 0.15$

Table 4. Copper ion reductive capability

CS extracts samples	Copper ion reductive capability
CS-S	$0.89 \pm 0.09$
CS-M	$0.78 \pm 0.09$
CS-MS	$1.21 \pm 0.13$

# 3.2 CS extracts bioactive ingredients, functional groups and molecuar chemical structures analysis and determination

#### 3.2.2 HPLC analysis

The similar elution time vs. intesity curves have shown for CS-S, CS-M and CS-MS methanol extract samples in HPLC analysis in the 60°C, 70°C and 80°C extraction temperatures respectively, indicating the kinds of the determined functional ingredients from CS-S, CS-M and CS-MS in 60°C, 70°C and 80°C extraction temperatures are similar. According to the intesity peaks along with the elution time, the caffeic acid, chlorogenic acid, elagic acid, epicatechin, epigallocatechin, ferrulic acid, hydroxybenzoic\_acid, kaempferol, protocatechuic etylester, protocatechuic acid, rutin, sinapic acid, t-2-hydroxycin.acid o-coum, trans - cinnamic acid, trans - p - coum acid, vanillic acid and their concentrations were determined, the results and the extraction temperature vs. concentration kinetics and the comparison among CS-S, CS-M and CS-MS samples were studied. It is illustrated from that caffeic acid, epicatechin, ferrulic acid, protocatechuic etylester, protocatechuic\_acid, rutin, trans - p - coum acid have the higheat extraction rate in the temperature of 70°C, the rest of the phenolic acids have the higheat extraction yield rate in the temperature of 80°C for the various heat sensitivities of the different phenolic acids. Almost all of the determined phenolic acids contents are significantly coralated to the maturity stages, which is the CS-MS has the highest extracted phenolic acids content, the CS-S has the lowest, except for rutin, which has the highest phenolic acids content for CS-M samples. Therefore, the extraction temperature has the significant influences to the menthol extraction rate of the CS phenolic acids and the maturity stage has the significant influences to the CS phenolic acids content.

#### 3.3.1 Thrombin inhibition activity test

Figure 9 shows the similar trends of thrombin inhibition activity kinetics for CS flavonoids, polysaccharides and steroids, the inhibition rates are all positively coorelated to the reaction time, where the polysaccharides have the highest thrombin inhibition activity than the CS polysaccharides and steroids extracts, meanwhile, the steroids have the lowest. It is notable that from 0-1 min, the inhibition rate increased sheerly and after 1 min, the trend became subdued for all of the CS flavonoids, steroids and polysaccharides samples. The result indicates the CS polysaccharides have the best effect of anti-coagulation among the three CS extracts and CS steroids have the worest.

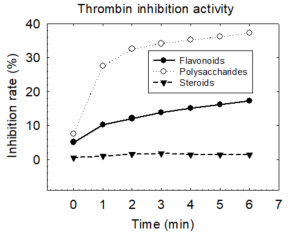


Figure 9. CS flavonoids, polysaccharides and steroids thrombin inhibition activity kinetics 3.3.2 ACE inhibition activity test

Figure 10 shows the similar trends of ACE inhibition activity kinetics for CS flavonoids, polysaccharides and steroids, the inhibition rates are all positively coorelated to the CS extract concentrations, where the polysaccharides have the highest ACE inhibition activity than the CS polysaccharides and steroids extracts, meanwhile, the steroids have the lowest. It is notable that from 20 – 100 mg/mL CS extract concentration, the inhibition rate increased sheerly and after 100 mg/mL, the trend became subdued for all of the CS flavonoids, steroids and polysaccharides samples. The result indicates the CS polysaccharides have the best effect of anti- hypertension among the three CS extracts and CS steroids have the worest.

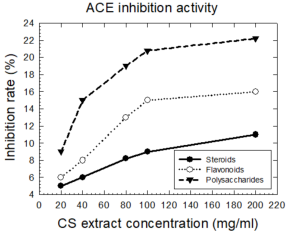


Figure 10. CS flavonoids, polysaccharides and steroids ACE inhibition activity kinetics 3.3.3 α- glucosidase inhibition activity test

Figure 11 shows the similar trends of  $\alpha$ - glucosidase inhibition activity kinetics for CS flavonoids, polysaccharides and steroids, the inhibition rates are all positively coorelated to the CS extract concentrations, where the polysaccharides have the highest  $\alpha$ - glucosidase inhibition activity than the CS polysaccharides and steroids extracts, meanwhile, the steroids have the lowest. It is notable that from 20-80 mg/mL CS extract concentration, the inhibition rate increased sheerly and after 80 mg/mL, the trend became subdued for the CS flavonoids sample. The result indicates the CS polysaccharides have the best effect of anti - diabetes among the three CS extracts and CS steroids have the worest.

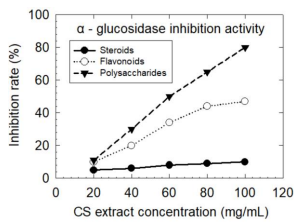


Figure 11. CS flavonoids, polysaccharides and steroids  $\alpha$ - glucosidase inhibition activity kinetics

#### 3.4.4 $\alpha$ - amylase inhibition activity test

Figure 12 shows the similar trends of  $\alpha$ - amylase inhibition activity kinetics for CS flavonoids, polysaccharides and steroids, the inhibition rates are all positively coorelated to the CS extract concentrations, where the polysaccharides have the highest  $\alpha$ - amylase inhibition activity than the CS polysaccharides and steroids extracts, meanwhile, the steroids have the lowest. The result indicates the CS polysaccharides have the best effect of anti - diabetes among the three CS extracts and CS steroids have the worest.

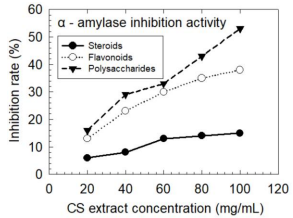


Figure 12. CS flavonoids, polysaccharides and steroids  $\alpha$ - amylase inhibition activity kinetics **3.4.5 XOD** inhibition activity test

Figure 13 shows the similar trends of XOD inhibition activity kinetics for CS flavonoids, polysaccharides and steroids, the inhibition rates are all positively coorelated to the CS extract concentrations, where the flavonoids have the highest XOD inhibition activity than the CS polysaccharides and steroids extracts, meanwhile, the steroids have the lowest. The result indicates the CS flavonoids have the best effect of anti - gout among the three CS extracts and CS steroids have the worest.

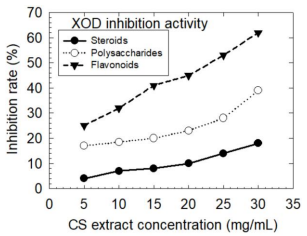


Figure 13. CS flavonoids, polysaccharides and steroids XOD inhibition activity kinetics **CONCLUSION** 

The current work researched about the three different maturity stages (silking stage, milky stage and mature stage) corn silk fiber and powder physicochemical properties and corn silk extracts biochemical properties, applied the technologies of UV-VIS, HPLC, NMR, FTIR and Fluorescence excitation-emission mapping to extract, determine the corn silk flavonoids, polysaccharides and steroids and optimized the extraction methods, the phenolic acids such as caffeic acid, chlorogenic acid, elagic acid, epicatechin, epigallocatechin, ferrulic acid, hydroxybenzoic\_acid, kaempferol, protocatechuic etylester, protocatechuic\_acid, rutin, sinapic acid, t-2-hydroxycin.acid\_o-coum, trans - cinnamic acid, trans - p - coum acid, vanillic acid and their concentrations were determined. It was applied the technologies of TG/DTA and SEM to analyze and research the corn silk fiber and powder thermol properties and microstructures, applied the technologies of EPR, DPPH and ABTS free - radicals scavenging capability test, ferricion and copper ion anti - oxident capability test and vitro enzyme inhibition activity test including the enzymes of  $\alpha$  - glucosidase and  $\alpha$  - amylase (anti - diabetes), thrombin (anti angiotensin converting enzyme (ACE) (anti hypertension) xanthine oxidase (XOD) (anti-gout) to analyze the corn silk extracts biochemical properties, the results show that the CS maturity is significantly corrected to the radical scavenging capability, where the CS-MS extracts have the strongest radical scavenging capability, then CS-S medium, CS - M the weakest, furthermore, the CS polysaccharides have the best anti-diabetes, anti-coagulation and anti-hypertension effect, CS flavonoids have the best anti-gout effect, meanwhile, the CS steroids showed the worest effect for all of the anti-diabetes, anticoagulation, anti-hypertension and anti-gout vitro enzymatic inhibition tests.

It can assume that the above research will play an irreplaceable role in analyzing the physicochemical and biochemical properties of botanic materials in the areas of food production, pharmacy, veterinary, animal feed and healthcare.

## BENEFITS OF THE OBTAINED RESULTS FOR THE STUDY SUBJECT FOOD TECHNOLOGY

Based on the above mentioned results of the study performed one can expect the following benefits for the study subject as follows:

1. There were obtained extraction kinetics data of medically valuable substances, such as flavonoids, polysaccharides and plant steroids suitable as the starting parameters for the optimization of the extraction procedures in semi-pilot scale production units.

- 2. There was confirmed high radical scavenger activity of all above mentioned extracted substances by means of Electron Paramagnetic Resonance and DPPA and ABTS radical inhibition experiments.
- 3. There were confirmed high inhibition activities of extracted substances from the corn silk material in vitro to the enzymes such as  $\alpha$  glucosidase and  $\alpha$  amylase (anti diabetes), thrombin (anti coagulation), angiotensin converting enzyme (ACE) (anti hypertension) and xanthine oxidase (XOD) (anti-gout) thus indicating its positive health effect.
- 4. There were determined 16 phenolic substances such as  $\alpha$  glucosidase and  $\alpha$  amylase (anti diabetes), thrombin (anti coagulation), angiotensin converting enzyme (ACE) (anti hypertension) and xanthine oxidase (XOD) (anti-gout) in extracts and their extraction kinetics were quantified.
- 5. Obtained data represent complex study of both nutrition as well as pharmaceutical industry interesting substances suitable for application in special food products.

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#### LIST OF SYMBOLS AND ABBREVIATIONS

%	percent
°C	centigrade
D	detector
ei	error

g gram

k the number of independent parameters

 $\begin{array}{ll} S & & \text{light source} \\ X_i & & \text{variable} \\ X_j & & \text{variable} \\ Y & & \text{response} \end{array}$ 

 $\beta_0$  model intercept coefficient  $\beta_{ij}$  the second-order terms

β<sub>i</sub> interaction coefficients of linear

 $\begin{array}{ll} \beta_{jj} & quadrantic \\ \mu M & millimole \\ ml & milliliter \end{array}$ 

ABTS 2, 2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid)

ACE angiotensin converting enzyme

ADP adenosine diphosphate BBD box-behnken design

CAT catalase from micrococcus lysodeiktic

CS corn silk

CSBIE corn silk bioactive ingredients extraction

**CSF** corn silk flavonoids **CSM** corn silk milky stage **CSMS** corn silk mature stage **CSP** corn silk polysaccharides **CSS** corn silk silking stage **DCS** dry mass of corn silk **DMSO** dimethyl sulfoxide deoxyribonucleic acid **DNA** 

DPPH 1,1-diphenyl-2-picrylhydrazyl
DTA differential thermal analysis

EPR electron paramagnetic resonance

FTIR fourier transform infrared spectrometer

GAE gallic acid equivalents HCl hydrochloric acid  $H_2O_2$  hydrogen peroxide

HO2- hydrogen peroxide radicals

HPLC high performance liquid chromatography

H<sub>2</sub>SO<sub>4</sub> sulfuric acid MPa mega pascal

MTT thiazolyl blue tetrazolium bromide

mRNA messenger ribonucleic acid NMR nuclear magnetic resonance PBS phosphate buffer
RCO- acyl radicals
RE rutin equivalents
RO- alkoxy radicals
RO2- peroxyalkyl radicals

Sa sample

SEM scanning electron microscope

SOD superoxide dismutase
TAC total anthocyanin
TFC total flavonoids

TG thermogravimetric analysis

TPC total phenolic

UV-VIS ultraviolet and visible spectrophotometry

XOD xanthine oxidase

#### LIST OF PUBLICATIONS

Year of publication 2019

Jimp

SALEK, R. N., VAŠINA, M., LAPČÍK, L., ČERNÍKOVÁ, M., LORENCOVÁ, E., LI, P., BUŇKA, F. Evaluation of various emulsifying salts addition on selected properties of processed cheese sauce with the use of mechanical vibration damping and rheological methods. LWT-Food Science and Technology. 2019, 107, 178-184.

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Year of publication 2018

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LI, P., LAPČÍK, L. Výzkum funkčních složek kukuřičného hedvábí. Chemické listy. 2018, 112 (2), 93-97.

Jscop

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VALENTA, T., LAPČÍKOVÁ, B., LAPČÍK, L., LI, P. The effect of conformational transition of gelatin-polysaccharide polyelectrolyte complex on its functional properties. Potravin árstvo Slovak Journal of Food Sciences. 2017, 11, 587-596.

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## The study of functional ingredients from corn silk

Studium funkčních přísad z kukuřičného hedvábí

**Doctoral Thesis Summary** 

Published by: Tomas Bata University in Zl ń, n ám. T. G. Masaryka 5555, 760 01 Zl ń.

Edition: published electronically

Typesetting by: Li Peng Ph.D.

This publication has not undergone any proofreading or editorial review.

Publication year: 2020

First Edition

ISBN 978-80-7454-964-9

