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On-line quantitative monitoring of liquid-liquid extraction of *Lonicera japonica* and *Artemisia annua* using near-infrared spectroscopy and chemometrics

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ABSTRACT

Background: Liquid-liquid extraction of Lonicera japonica and Artemisia annua (JQ) plays a significant role in manufacturing Reduning injection. Many process parameters may influence liquid-liquid extraction and cause fluctuations in product quality. Objective: To develop a near-infrared (NIR) spectroscopy method for on-line monitoring of liquid-liquid extraction of JQ. Materials and Methods: Eleven batches of JQ extraction solution were obtained, ten for building quantitative models and one for assessing the predictive accuracy of established models. Neochlorogenic acid (NCA), chlorogenic acid (CA), cryptochlorogenic acid (CCA), isochlorogenic acid B (ICAB), isochlorogenic acid A (ICAA), isochlorogenic acid C (ICAC) and soluble solid content (SSC) were selected as guality control indicators, and measured by reference methods. NIR spectra were collected in transmittance mode. After selecting the spectral sub-ranges, optimizing the spectral pretreatment and neglecting outliers, partial least squares regression models were built to predict the content of indicators. The model performance was evaluated by the coefficients of determination (R^2), the root mean square errors of prediction (RMSEP) and the relative standard error of prediction (RSEP). Results: For NCA, CA, CCA, ICAB, ICAA, ICAC and SSC, R² was 0.9674, 0.9704, 0.9641, 0.9514, 0.9436, 0.9640, 0.9809, RMSEP was 0.0280, 0.2913, 0.0710, 0.0590, 0.0815, 0.1506, 1.167, and RSEP was 2.32%, 4.14%, 3.86%, 5.65%, 7.29%, 6.95% and 4.18%, respectively. Conclusion: This study demonstrated that NIR spectroscopy could provide good predictive ability in monitoring of the content of quality control indicators in liquid-liquid extraction of JQ.

Key words: Liquid-liquid extraction, near-infrared spectroscopy, on-line monitoring, partial least squares

INTRODUCTION

Lonicera japonica (Jinyinhua in Chinese) and Artemisia annua (Qinghao in Chinese) are two fundamental herbs in Reduning injection, which has long been used for the treatment of upper respiratory tract infections in China. Previous studies have shown that primary bioactive compounds of Jinyinhua are caffeoylquinic acids,^[1] including neochlorogenic acid (NCA), chlorogenic acid (CA), cryptochlorogenic acid (CCA),

Address for correspondence: Prof. Wei Xiao, National Key Laboratory of Pharmaceutical New Technology for Chinese Medicine, Kanion Pharmaceutical Corporation, Lianyungang, 222000, China. E-mail: xw_kanion@163.com isochlorogenic acid B (ICAB), isochlorogenic acid A (ICAA) and isochlorogenic acid C (ICAC), which possess anti-inflammatory properties.^[2] The primary compounds of Qinghao are artemisinin and essential oil,^[3] which show antimalarial and antibacterial activities.^[4]

Liquid-liquid extraction separates one or several substances present in a liquid phase by the addition of another liquid phase in which these substances transferred preferentially.^[5] It is a widely used purification technique in manufacturing traditional Chinese medicine (TCM), with its ability to remove impurities such as proteins and polysaccharides because of their low solubility in organic solvent phase.^[6] Nevertheless, many process parameters may influence extraction process, such as temperature, initial density of starting material, volume of solvent, flow rate of



solvent and pH of the solution. These factors can have a large impact on the downstream unit operations as well as on the final drug quality.

It is reported that the national adverse drug reaction monitoring database collected 12,000 adverse drug events about Chinese herbal injection in 2013. The safety evaluation of Chinese herbal injection has been concerned dramatically, and batch-to-batch variations are a major issue on safety. It is necessary to adopt effective analytical techniques for timely assessment of critical quality attributes in Chinese herbal injection.^[7] Various methods have been applied for quality control, such as ultraviolet spectroscopy, high-performance liquid chromatography (HPLC), liquid chromatography-mass spectrometry (LC-MS), gas chromatography (GC) and GC-MS.^[8-10] All these conventional methods are time-consuming and reagent-consuming. However, the near-infrared (NIR) spectroscopy technique has proven its effectiveness in different fields, such as agriculture,[11] food,^[12,13] oil,^[14] chemical^[15] and pharmaceutical industries^[16,17] relying on its speed, accuracy, low cost and non-destructive characteristic toward the analyzed sample.^[18] There are many comprehensive reviews demonstrating the successful use of NIR spectroscopy for quantitative and qualitative analysis in TCM research.^[19-22] Therefore, NIR technique is applied to study liquid-liquid extraction of JQ in this research.

Neochlorogenic acid, CA, CCA, ICAB, ICAA and ICAC are the principal bioactive components in Reduning injection^[23] and have relatively high concentration in the JQ extraction solution. The soluble solid content (SSC) is the percentage of the total content of substances dissolved in the solution and indicates the purification ability of liquid-liquid extraction. Therefore, NCA, CA, CCA, ICAB, ICAA, ICAC and SSC were selected as quality control indicators in liquid-liquid extraction of JQ. Partial least squares (PLS) algorithm was applied to construct quantitative models to predict the content of above indicators. The prediction accuracy of the established models was evaluated by statistical parameters.

MATERIALS AND METHODS

Materials

Reference substance of CA was purchased from National Institutes for Food and Drug Control (Beijing, China). Reference substances of NCA, CCA, ICAB, ICAA and ICAC were purchased from Chengdu Must Bio-technology Co., LTD. HPLC grade methanol was obtained from Tedia Company (Fairfield, USA). Deionized water was obtained from a Mill-Q water purifier system (Millipore, Bedford, MA, USA). All the other reagents utilized in the article were of analytical grade.

Liquid-liquid extraction process

The process flow diagram of liquid-liquid extraction of JQ is shown in Figure 1. The countercurrent extraction column was made of glass pipe and packed with stainless-steel Raschig rings.^[24] Twelve separate columns comprised a group and their pipelines converged into a main pipeline down to a storage tank. Firstly, the pH value of starting materials was adjusted to 2.0 through adding hydrochloric acid. Then starting materials were divided into twelve parts evenly, and each pumped into the extraction column. Next, water-saturated ethyl acetate used as extraction solvent was continuously pumped into the column from the bottom. The extraction solution began to flow out from the column after adding ethyl acetate for 2 h. The extraction usually lasted for 10-12 h, which depended on the volume of starting materials. A sampling valve was installed into the main pipeline to collect extraction solution. NIR transmission probes were immersed into the solution to collect spectra.

Sample collection

The sample was collected with a volume of 80 mL at 20 min interval at the same time the sampling time was recoded. The number of samples may vary among batches because the volume of starting materials was different. Spectra were automatically collected every 20 s. According to the recorded sampling time, the corresponding spectrum was selected from the continuous spectra. Eleven batches of samples were collected in total, ten batches (250 samples) as a calibration set for model development and one (27 samples) as a prediction set



Figure 1: The process flow diagram of liquid-liquid extraction of JQ

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for model evaluation. Each sample was marked as batch number-sampling order, e. g, 140110-1, meaning the first sample collected from batch 140110.

Reference assays

A newultra-high performance liquid chromatography (UPLC) method was developed to measure the content of NCA, CA, CCA, ICAB, ICAA and ICAC. The chromatographic analysis was performed on an Agilent 1290 UPLC system (Agilent Technologies, USA), which was comprised of a quaternary bump, an online vacuum degasser, an auto-sampler, a thermostatic column compartment and a diode-array detector. The chromatographic separations were undertaken on an Agilent ZORBAX SB-C18 column (100 mm \times 3.0 mm, 1.8 µm). Solvent A [methanol] and solvent B (aqueous phosphoric acid solution [0.1%, v/v]) were used as mobile phase for the gradient elution [Table 1]. The detection wavelength was set at 324 nm. The flow rate was 0.4 mL/min and the injection volume was 2 µL. The column temperature was maintained at 30°C.

The extraction solution was centrifuged at 10,000 rpm for 10 min and then diluted to 50 mL with methanol:water (50:50, v/v). The supernatant fluid was passed through a 0.22 μ m syringe filter, and 2 μ L filtrate was injected into the UPLC system.

The SSC was determined by drying 5 mL centrifuged solution at 105°C to a constant weight. The results are expressed in weight mg/mL.

Spectral collection

A Luminar 3060 acousto-optic tunable filter-NIR spectrometer (Brimrose Co., USA) with an InGaAs detector was used to collect NIR spectra in transmittance mode. Two fiber optic probes with 2 mm optical path length were connected to the spectrometer by an optical fiber. The spectral range was from 1,100 nm to 2,300 nm. Each spectrum was the average of 200 scans with a wavelength increment of 2 nm.

Spectral preprocessing

Spectral preprocessing is the mathematical correction of spectra that reduce, eliminate or standardize the effect of variable physical properties or instrumental effects.^[25] A variety of preprocessing methods were

Table 1: Gradient elution schedule				
Time (min)	A (%)	B (%)		
0	15	85		
5	25	75		
15	35	65		
30	50	50		

used to separate the useful information from noise, such as multiplicative signal correction (MSC), standard normal variate, first derivative, the second derivative and Savitzky-Golay smoothing. The optimum method was selected based on the lowest root mean square errors of calibration (RMSEC) and root mean square errors of cross-validation (RMSECV).^[26]

Model building

A PLS model was constructed by spectral data matrix X and the reference content matrix Y. Model was validated by cross-validation with the leave-one-out method. The leaving-one-out method is leaving one sample out, using the rest of the samples in the calibration set to predict the sample being left out.^[27]

RESULTS

Method validation for ultra-high performance liquid chromatography

The UPLC chromatograms of mixed standards [Figure 2a] and sample solution [Figure 2b] were baseline separated. The calibration curves were created by plotting the concentration (x, $\mu g/mL$) versus the peak area (y) of each analyte. The linear regression equations and the coefficients of determination (R^2) were as follows: Y = 14.48x + 0.214 ($R^2 = 0.9999$), for NCA; y = 14.108x - 5.757 ($R^2 = 0.9999$), for CA; y = 12.657x - 0.331 ($R^2 = 0.9999$), for CCA; y = 14.394x - 0.909 ($R^2 = 0.9999$), for ICAB; y = 16.956x - 3.364 ($R^2 = 0.9999$), for ICAA; y = 16.066x - 8.299 ($R^2 = 0.9999$), for ICAC. Good linearity was achieved in the investigated ranges. The linear ranges were 4.28-17.12, 36.72-146.88, 7.75-31.01, 7.28–29.13, 8.39–33.57 and 15.66–62.64 µg/mL for NCA, CA, CCA, ICAB, ICAA and ICAC, respectively.

The repeatability of the method was determined by analyzing six independently prepared solution of sample 140110–11. The relative standard derivation (RSD) values of the concentration for six analytes were found to be within 0.94–1.08%.

The stability was tested at room temperature by analyzing the same solution (140110–11) at 0, 2, 4, 8, 16 and 24 h, respectively. The RSD values of the peak areas for six analytes were all <1.11%, indicating sample solution was stable within 24 h.

The percent recovery was detected by adding known amounts of the corresponding analytes to previously analyzed sample 140110–11 and reanalyzing the sample. The percent recovery was between 97.32% and 103.40%,



Figure 2: The ultra-high performance liquid chromatography chromatograms of mixed standards (a) and sample solution (b). (1: Neochlorogenic acid; 2: Chlorogenic acid; 3: Cryptochlorogenic acid; 4: Isochlorogenic acid B; 5: Isochlorogenic acid A; 6: Isochlorogenic acid C)

with RSD values <1.73% for six analytes, indicating the analytical method used for quantitative determination was acceptable.

Dynamic curves during liquid-liquid extraction of JQ The dynamic curves for NCA, CA, CCA, ICAB, ICAA, ICAC and SSC of batch 140110 during liquid-liquid extraction of JQ are shown in Figure 3. These indicators had similar changing trends, and their concentration constantly decreased with the passage of time. It is easy to notice that CA and ICAC decreased dramatically, but NCA, CCA, ICAB and ICAA declined slowly. Their content tended to be stable in the 600th min. SSC dropped from 23 mg/mL to 4.4 mg/mL, indicating the effective ingredients were transferred into the ethyl acetate solution from the starting material.

Selecting spectral sub-range

In general, peak assignment to specific chemical groups with accuracy is difficult since NIR spectra are combinations of stretching and deformation vibration of hydrogen bonds,^[28] but the bands of interest for analytes could be identified in NIR spectroscopy. The collected original spectra are depicted in Figure 4. The absorbance values at 1950-2300 nm were higher than 1.5, belonging to saturated absorbance, so this region was not recommended for building NIR models.^[29] The basic functional groups of caffeoylquinic acids are carbonyl, hydroxyl and carboxyl groups. The band around 1167 nm is caused by the C-H stretching 2nd overtone of CH₂ and -CH₂- groups, and those around 1391 nm and 1413 nm are caused by the 2C-H stretching and C-H deformation vibration of CH, and -CH₂- groups, respectively.^[30] Besides, the peak at 1720 nm is attributed to the asymmetric overtone of the C-H bond.^[31] Furthermore, the peak at 1900 nm is assigned to the 2nd overtone of the CO bond.^[32] After comparing the combinations of different wavelength ranges, it was found that 1178-1918 nm was optimum



Figure 3: Time evolution curves of phenolic acids (a) and soluble solid content (b) of batch 140110 during liquid-liquid extraction of JQ

with the lowest RMSECV and RMSEC. The same mother nucleus structure of caffeoylquinic acids (NCA, CA, CCA, ICAB, ICAA and ICAC) made their NIR absorption peaks close, so their spectral sub-ranges were identical.



Figure 4: Original near-infrared spectra collected in the liquid-liquid extraction of JQ

Establishing calibration models

By comparing various spectral pretreatments, the MSC combined with first derivative and Savitzky-Golay smoothing revealed low RMSRC and RMSECV for NCA, CA, CCA, ICAB, ICAA, ICAC and SSC. Before building calibration models, spectral data sets were analyzed through principal component analysis (PCA). The matrix X is decomposed into a few new variables, called principal components (PCs), with the aim to maximize the explained variance in the matrix X.^[33,34] Taking NCA as example, X variance was explained by PC1 83% and PC2 8%, matrix Y was explained by PC1 54% and PC2 18%, and six PCs were selected for building the model. Outliers are detected based on the leverage level and the residual of X variance. Taking ICAB as representative, observation 57 had a large residual and a relative small leverage level, indicating it was an outlying observation [Figure 5] and should be removed from the calibration model. After PCA and detecting outliers, the PLS models for NCA, CA, CCA, ICAB, ICAA, ICAC and SSC were established, and the model quality parameters were listed in Table 2.

On-line quantitative monitoring

The predictive ability of established PLS models was evaluated on batch 140120. The measured values were



Figure 5: Partial least squares prediction regression equations of neochlorogenic acid (a), chlorogenic acid (b), cryptochlorogenic acid (c), isochlorogenic acid B (d), isochlorogenic acid A (e), isochlorogenic acid C (f) and soluble solid content (g) of batch 140120, scanned by on-line near-infrared and measured by reference assays

Table 2	2: Stati	stics o	of the	optimal	PLS	models	for
liquid-	liquid e	extract	ion o	f JQ			

PCs	Calibra	Calibration set		Cross-validation set		
	R ²	RMSEC	R ²	RMSECV		
6	0.9672	0.0263	0.9634	0.0278		
6	0.9742	0.2528	0.9712	0.2669		
6	0.9734	0.0700	0.9705	0.0737		
6	0.9832	0.0627	0.9777	0.0652		
6	0.9748	0.0725	0.9722	0.0761		
6	0.9771	0.1316	0.9749	0.1379		
5	0.9781	0.8911	0.9759	0.9339		
	PCs 6 6 6 6 6 6 5	PCs Calibra R ² 6 6 0.9672 6 0.9742 6 0.9734 6 0.9832 6 0.9748 6 0.9771 5 0.9781	PCs Calibration set R² RMSEC 6 0.9672 0.0263 6 0.9742 0.2528 6 0.9734 0.0700 6 0.9832 0.0627 6 0.9748 0.0725 6 0.9771 0.1316 5 0.9781 0.8911	PCs Calibration set R ² Cross-val RMSEC 6 0.9672 0.0263 0.9634 6 0.9742 0.2528 0.9712 6 0.9734 0.0700 0.9705 6 0.9832 0.0627 0.9777 6 0.9748 0.0725 0.9722 6 0.9771 0.1316 0.9749 5 0.9781 0.8911 0.9759		

PLS: Partial least squares; PCs: Principal components; RMSEC: Root mean square errors of calibration; RMSECV: Root mean square errors of cross-validation; NCA: Neochlorogenic acid; CA: Chlorogenic acid; CCA: Cryptochlorogenic acid; ICAB: Isochlorogenic acid B; ICAA: Isochlorogenic acid A; ICAC: Isochlorogenic acid C; SSC: Soluble solid content

compared with the predicted values. The root mean square errors of prediction (RMSEP) and relative standard error of prediction (RSEP) were used to assess the predictive ability [Table 3].

DISCUSSION

As shown in Table 2, the R^2 of calibration and cross-validation sets was higher than 0.96 and RMSECV was close to RMSEC. NCA model had the minimum RMSEC and RMSECV, whereas SSC model had the maximum RMSEC and RMSECV. It could be caused by the low content of NCA between 0.10 mg/ml and 0.53 mg/ml, and the high content of SSC ranged from 2.18 mg/ml to 20.76 mg/ml.

In this study, when RSEP is lower than 8%, the predictive accuracy is acceptable for quality control requirements. As seen in Table 3, all the RSEP satisfied the standard, indicating all the models were suitable for industrial application. The RSEP values were sorted in ascending order as follows: NCA, CCA, CA, SSC, ICAB, ICAC and ICAA. Obviously, NCA model with an RSEP value of 2.32% displayed the best predictive ability, although its content was lower than 0.6 mg/ml in the extract solution. However, ICAA model with an RSEP value of 7.29% showed maximum predictive errors, although its high content ranged from 0.1 mg/ml to 1.5 mg/ml. The possible reason is the NIR absorption peaks are more representative of NCA than ICAA, although they have similar chemical structures. The predictive accuracy may be not only associated with the content of analyte, but also depend on the linkage between the chemical structure and the characteristic absorption. And the similar chemical structures probably cause multicollinearity,^[35] which brings about the decline of predictive ability.

Figure 6 shows the concentration (content) regression trends predicted by on-line NIR and measured by reference assays. It has to be mentioned that the curves of NCA, CA, CCA and SSC showed good linear relationship and small deviations between measured values and predicted values. However, the curves of ICAB, ICAA and ICAC showed high predictive errors, which were consistent with the high RSE*P* values in Table 3. Overall, the predictive ability of NCA, CA, CCA models were slightly better than that of ICAB, ICAA and ICAC models.

CONCLUSION

A NIR spectroscopy method was developed to monitor liquid-liquid extraction of JQ by predicting the content of NCA, CA, CCA, ICAB, ICAA, ICAC and SSC.



Figure 6: Partial least squares prediction regression equations of neochlorogenic acid (a), chlorogenic acid (b), cryptochlorogenic acid (c), isochlorogenic acid A (e), isochlorogenic acid C (f) and soluble solid content (g) of batch 140120, scanned by in-line near-infrared and measured by reference assays

Table 3: Statistics of the prediction set of the quantitative models

		Prediction set			
	R ²	RMSEP	RSEP %		
NCA	0.9674	0.0280	2.32		
CA	0.9704	0.2913	4.14		
CCA	0.9641	0.0710	3.86		
ICAB	0.9514	0.0590	5.65		
ICAA	0.9436	0.0815	7.29		
ICAC	0.9640	0.1506	6.95		
SSC	0.9809	1.167	4.18		

RMSEP: Root mean square errors of prediction; RSEP: Relative standard error of prediction; NCA: Neochlorogenic acid; CA: Chlorogenic acid; CA: Cryptochlorogenic acid; ICAB: Isochlorogenic acid B; ICAA: Isochlorogenic acid A; ICAC: Isochlorogenic acid C; SSC: Soluble solid content

The low RMSEP, RSEP and high R² confirmed the established models had good predictive performance in real-time quality control. Based on this study, NIR spectroscopy could be extended to be applied in liquid-liquid extraction of TCM, ensuring rapid monitoring of batch production process and minimizing the batch-to-batch fluctuations.

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