A peer-reviewed version of this preprint was published in PeerJ on 22 January 2019.

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Wang H, Chen Y, Zhang Z, Chen H, Li X, Wang M, Chai H. 2019. Quantitatively estimating main soil water-soluble salt ions content based on Visible-near infrared wavelength selected using GC, SR and VIP. PeerJ 7:e6310 <u>https://doi.org/10.7717/peerj.6310</u>

Quantitatively estimating main soil water-soluble salt ions content based on Visible-near infrared wavelength selected using GC, SR and VIP

Haifeng Wang $^{1,\,2}$, Yinwen Chen 3 , Zhitao Zhang $^{\rm Corresp.,~1,\,2}$, Haorui Chen 4 , Xianwen Li 2 , Mingxiu Wang 5 , Hongyang Chai 2

¹ Key Laboratory of Agricultural Soil and Water Engineering in Arid and Semiarid Areas, Ministry of Education, Northwest A&F University, Yangling, Shaanxi, China

² College of Water Resources and Architectural Engineering, Northwest A&F University, Yangling, Shaanxi, China

³ Department of Foreign Languages, Northwest A&F University, Yangling, Shaanxi, China

⁴ Department of Irrigation and Drainage, China Institute of Water Resources and Hydropower Research, Beijing, China

⁵ Department of Civil and Environmental Engineering, University of California, Irvine, CA, USA

Corresponding Author: Zhitao Zhang Email address: zhitaozhang@126.com

Soil salinization is the primary obstacle to the sustainable development of agriculture and eco-environment in arid regions. The accurate inversion of the major water-soluble salt ions in the soil using visible-near infrared (VIS-NIR) spectroscopy technique can enhance the effectiveness of saline soil management. However, the accuracy of spectral models of soil salt ions turns out to be affected by high dimensionality and noise information of spectral data. This study aims to improve the model accuracy by optimizing the spectral models based on the exploration of the sensitive spectral intervals of different salt ions. To this end, 120 soil samples were collected from Shahaogu Irrigation Area in Inner Mongolia, China. After determining the raw reflectance spectrum and content of salt ions in the lab, the spectral data were pre-treated by standard normal variable (SNV). Subsequently the sensitive spectral intervals of each ion were selected using methods of gray correlation (GC), stepwise regression (SR) and variable importance in projection (VIP). Finally, the performance of both models of partial least squares regression (PLSR) and support vector regression (SVR) was investigated on the basis of the sensitive spectral intervals. The results indicated that the model accuracy based on the sensitive spectral intervals selected using different analytical methods turned out to be different: VIP was the highest, SR came next and GC was the lowest. The optimal inversion models of different ions were different. In general, both PLSR and SVR had achieved satisfactory model accuracy, but PLSR outperformed SVR in the forecasting effects. Great difference existed among the optimal inversion accuracy of different ions: the predicative accuracy of Ca²⁺, Na⁺, Cl⁻, Mg²⁺ and SO_4^{2-} was very high, that of CO_3^{2-} was high and K⁺ was relatively lower, but HCO_3^{-} failed to have any predicative power. These findings provide a new approach for the optimization of the spectral model of water-soluble salt ions and improvement of its predicative precision.

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5 Haifeng Wang^{1,2*}, Yinwen Chen^{3*}, Zhitao Zhang^{1,2}, Haorui Chen⁴, Xianwen Li², Mingxiu

6 Wang⁵ and Hongyang Chai²

7

⁸ ¹ Key Laboratory of Agricultural Soil and Water Engineering in Arid and Semiarid Areas,

- 9 Ministry of Education, Northwest A&F University, Yangling, Shaanxi, China
- 10 ² College of Water Resources and Architectural Engineering, Northwest A&F University,
- 11 Yangling, Shaanxi, China
- ¹² ³ Department of Foreign Languages, Northwest A&F University, Yangling, Shaanxi, China
- ⁴ Department of Irrigation and Drainage, China Institute of Water Resources and Hydropower
- 14 Research, Beijing, China
- ⁵ Department of Civil and Environmental Engineering, University of California, Irvine, CA, USA
- 17 * These authors contributed equally to this work.
- 18 Corresponding Author:
- 19 Zhitao Zhang^{1,2}
- 20 No.23 Weihui Road, Yangling, Shaanxi, 712100, China
- 21 Email address: <u>zhitaozhang@126.com</u>
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33

34 **ABSTRACT**

Soil salinization is the primary obstacle to the sustainable development of agriculture and eco-35 environment in arid regions. The accurate inversion of the major water-soluble salt ions in the 36 soil using visible-near infrared (VIS-NIR) spectroscopy technique can enhance the effectiveness 37 38 of saline soil management. However, the accuracy of spectral models of soil salt ions turns out to be affected by high dimensionality and noise information of spectral data. This study aims to 39 improve the model accuracy by optimizing the spectral models based on the exploration of the 40 sensitive spectral intervals of different salt ions. To this end, 120 soil samples were collected 41 from Shahaoqu Irrigation Area in Inner Mongolia, China. After determining the raw reflectance 42 43 spectrum and content of salt ions in the lab, the spectral data were pre-treated by standard normal variable (SNV). Subsequently the sensitive spectral intervals of each ion were selected using 44 methods of gray correlation (GC), stepwise regression (SR) and variable importance in 45 projection (VIP). Finally, the performance of both models of partial least squares regression 46 (PLSR) and support vector regression (SVR) was investigated on the basis of the sensitive 47 spectral intervals. The results indicated that the model accuracy based on the sensitive spectral 48 intervals selected using different analytical methods turned out to be different: VIP was the 49 highest, SR came next and GC was the lowest. The optimal inversion models of different ions 50 were different. In general, both PLSR and SVR had achieved satisfactory model accuracy, but 51 PLSR outperformed SVR in the forecasting effects. Great difference existed among the optimal 52 inversion accuracy of different ions: the predicative accuracy of Ca²⁺, Na⁺, Cl⁻, Mg²⁺ and SO₄²⁻ 53 was very high, that of CO_3^{2-} was high and K⁺ was relatively lower, but HCO_3^{-} failed to have any 54 predicative power. These findings provide a new approach for the optimization of the spectral 55 model of water-soluble salt ions and improvement of its predicative precision. 56

57 Introduction

Soil salinization, one of the most important causes of land desertification and deterioration, has 58 59 posed serious threat to agricultural development and sustainable utilization of natural resources 60 (Shahid & Rahman, 2011; Abbas et al. 2013). 950 million ha of saline soil worldwide has become salinized (Schofield & Kirkby, 2003). Soil salinization is eroding and degenerating the 61 arable soil at the speed of 10 ha/min (Graciela & Alfred, 2009). Soil remediation and 62 management are very difficult in China because of such complex natural factors as climate, 63 terrain and geology, and human factors as unreasonable irrigation and disruption of ecological 64 balance. The total area of saline soil in China is 36 million ha (Li et al. 2014), accounting for 65 4.88% of the total area available nationwide (The National Soil Survey Office, 1998). Saline soil 66

67 usually has a high concentration of salt ions with a series of effects on the plants such as 68 physiological draught, ion toxicity and metabolic disorder, thus forming "salt damage" (Munns, 69 2002; Tavakkoli et al. 2011). In addition, one major cause of the inaccuracy of soil salinity 70 spectral measurement is that pure salts seldom exist in the soil because of some trace salt ion 71 elements are always fixed in soil crystals. Therefore, quick and accurate acquisition of the 72 detailed information of the various salt ions content in the soil can enhance the pertinence and 73 effectiveness of saline soil management.

74 The traditional quantitative estimation of soil salt contents usually includes such steps as field 75 soil sampling in fixed points, experiments in the laboratory and comprehensive statistical analysis (Urdanoz & Aragüés, 2011). Such method is incapable of the dynamic monitoring of 76 saline soil in a large area because of its high consumption of time and energy, small number of 77 measuring points and poor representativeness (Ding & Yu, 2014). Compared with conventional 78 79 laboratory analysis methods, remote sensing technology has been widely used due to its rich information, continuity, high precision and low cost (Ben-Dor, 2002; Viscarra Rossel et al. 2006; 80 Viscarra Rossel & Behrens, 2010; Viscarra Rossel & Webster, 2012). The various soil 81 constituents (contents of water, salt, organic matter and so forth) can be acquired conveniently 82 from remote sensing data (Gomez et al. 2008; Yu et al. 2010; Periasamy & Shanmugam, 2017). 83 84 Hence, with the abundant spectral reflection information within the VIS-NIR intervals of soil salinity, it is feasible to improve the accuracy of soil salinization inversion (Al-Khaier, 2003; 85 86 Ben-Dor et al. 2009; Abbas et al. 2013).

The application of VIS-NIR spectral analysis technique has been proved effective in 87 improving the accuracy of quantitative estimation and eliminating the external disturbance to 88 89 some extent (Dehaan & Taylor, 2002; Metternicht & Zinck, 2003; Farifteh et al. 2008). The 90 univariate linear regression on the basis of soil salinity index developed for CR (continuum removed) reflectance can be used as a method for soil salt content estimation (Weng et al. 2008). 91 Due to the strong correlation between soil electrical conductivity (EC) and soil salinity, EC is 92 93 also one of the important indicators for evaluating soil salinization degree. A variety of approaches have been used to acquire the EC in the field soil, including the partial least squares 94 regression (PLSR) and multivariate adaptive regression splines (MARS) (Volkan Bilgili et al. 95 2010; Nawar et al. 2015), logarithmic model (Xiao et al. 2016a), Bootstrap-BP neural network 96 model (Wang et al. 2018d) and satellite remote sensing technology (Nawar et al. 2014; Bannari 97 98 et al. 2018). In addition, the differential transformation (Xia et al. 2017) and fractional derivative 99 (Wang et al. 2017; Wang et al. 2018c) can fully utilize the potential spectral information and enhance model accuracy. The methods of spectral classification (Jin et al. 2015) and water 100 influence elimination (Chen et al. 2016; Peng et al. 2016; Yang & Yu, 2017) work well in 101 102 improving the quantitative inversion accuracy of soil salinity. Therefore, the remote sensing

103 technique is reliable to inverse the soil salinity quantitatively on different scales.

The quantitative analysis of VIS-NIR spectral intervals can help evaluate the content of some 104 chemical elements (Viscarra Rossel et al. 2006; Farifteh et al. 2008; Cécillon et al. 2009; Ji et al. 105 106 2016) due to the different characteristic absorption spectrum in soil chemical elements. Besides, there exists a correlation between some principal salt ions (Na⁺, Cl⁻) and spectral reflectance 107 (Jiang et al. 2017). Therefore, VIS-NIR spectroscopy technique can be used to obtain the 108 contents of the soil salt ions to a certain extent. The spectral response characteristics of mid-109 infrared (MIR) spectroscopy are better than those of VIS-NIR spectroscopy in predicting soil 110 salinity information, the latter has high predicting accuracy of the total salts content, HCO3-, 111 SO_4^{2-} and Ca^{2+} , followed by Mg²⁺, Cl⁻ and Na⁺ (Peng et al. 2016). The spectral models have 112 satisfactory prediction of the SAR (sodium absorption ratio) of soil salinization evaluation 113 parameter, which is composed of the contents of Ca^{2+} , Mg^{2+} and Na^{+} (Xiao et al. 2016b). Qu et 114 115 al. (2009) found that the contents of the total salt, SO_4^{2-} , pH and K⁺+Na⁺ have a higher inversion accuracy using spectral data to create PLSR model. The different pretreatment of the different 116 ion models varies by creating and analyzing PLSR model that demonstrates relatively good 117 predictive effects like ion contents of Ca^{2+} , Mg^{2+} , SO_4^{2-} , Cl^- , and HCO_3^- (Dai et al. 2015). 118 119 Overall, PLSR is a frequently used and robust linear model for quantitative research because it 120 has inference capabilities which are useful to model a probable linear relationship between the reflectance spectra and the salt ions content in soil. However, the non-uniform data and non-121 linear reflectance in spectral information of some soil chemical elements lead to the reduction in 122 model accuracy (Viscarra Rossel & Behrens, 2010; Nawar et al. 2015). In particular, support 123 124 vector regressions (SVR) based on kernel-based learning methods has the ability to handle nonlinear analysis case with high model accuracy (Vapnik, 1995; Peng et al. 2016; Hong et al. 125 126 2018b). Over the past several decades, the use of SVR for classification and regression has been extensively applied in soil VIS-NIR spectroscopy (Ben-Dor, 2002; Xiao et al. 2016b; Hong et al. 127 2018a). Moreover, the SVR model works well in estimating the contents of K⁺, Na⁺, Ca²⁺ and 128 SO_4^{2-} in the soil (Wang et al. 2018a). Thus, the correct way of modeling helps to guarantee the 129 model accuracy (Farifteh et al. 2007). 130

Many researches focused on the inversion of soil salinity using spectral information. 131 Nevertheless, little research has explored the eight water-soluble salt ions (K⁺, Ca²⁺, Na⁺, Mg²⁺, 132 Cl⁻, SO_4^{2-} , HCO_3^{-} and CO_3^{2-}) using spectral information in the soil. The model fitting of ions and 133 134 spectral information still needs improving (Farifteh et al. 2008; Peng et al. 2016). Apart from the suitable multivariate statistical analysis method that can partly improve the inversion effects, 135 reduction of redundant information is another identified approach to further optimize the model 136 (Bannari et al. 2018; Stenberg et al. 2010). Plenty of studies have demonstrated that spectral 137 138 variable selection methods can not only reduce the complexity of calibration models, but also 139 improve the model predictive performance (Hong et al. 2018a). To select the optimal spectral variable subset, scholars have investigated varied methods such as gray correlation (GC) (Li et al. 140 2016; Wang et al. 2018b), stepwise regression (SR) (Zhang et al. 2018) and variable importance 141 142 in projection (VIP) (Qi et al. 2017), and have achieved satisfactory effects. In addition, all the three methods have been widely applied in many studies, such as plant physiology, food 143 engineering, mathematical statistics (Oussama et al., 2012; Maimaitiviming et al. 2017; Liu et al. 144 2015). However, few studies have concentrated on the use of variable selection algorithms in the 145 inversion of soil salt ions. 146

This study aims to: (1) build the optimal model of soil salt ions using VIS–NIR spectroscopy technique; (2) compare the models based on the sensitive spectral ranges selected using GC, SR and VIP methods for different soil ions; (3) compare the performance of PLSR and SVR models, and identify the optimal models for different ions.

151 MATERIALS AND METHODS

152 Study area

153 Hetao Irrigation District (HID), with Yin Mountains at its north, the Yellow River at its south, Ulanbuh Desert at its west and Baotou at its east, lies in Bayannur League, Inner Mongolia, 154 China. It consists of irrigation areas of Ulan Buh, Jiefangzha, Yongji, Yichang and Urat, and it is 155 China's largest irrigation district with a total size of 5740 km² (Yu et al. 2010). In addition, HID 156 is an important production base of cereal and oil plants in China with major crops of wheat, corn 157 and sunflower. Shahaogu Irrigation Area (SIA), a typical region of saline soil in HID, was 158 chosen as the study area. SIA ($107^{\circ}05' \sim 107^{\circ}10'E$, $40^{\circ}52' \sim 41^{\circ}00'N$) is located in the central 159 160 east of Jiefangzha Irrigation Area. SIA belongs to typical continental climate, having hot summers, chilly winters, rare precipitation and strong evaporation. Its mean annual temperature, 161

162 precipitation, potential evaporation is about 7.1°C, 155 mm and 2000 mm, respectively.

Physiographically, the mean elevation and slope of SIA are about 1030 m and 1/10000, respectively. According to the World Reference Base for Soil Resources (WRB), the local soil texture is mainly silty clay loam with varying degrees of saline soil. Over the years, due to its gentle terrain slope, poor groundwater runoff, intense land surface evaporation and irrational farming activities, about 60% of the land within the district has been affected by various degree of salinization, which seriously restricted the agricultural development (Wu et al. 2008; Gao et al. 2015).

170 Sample collection and chemical analysis

171 The Hetao irrigation district administration gave field permit approval to us (NO. 2017YFC0403302). To ensure the representativeness of soil samples, the samples were

randomly gathered from a total of 120 sampling units on a grid of 16 m \times 16 m (because the 173 spatial resolution of GF-1 satellite imagery is 16 m) in the study area during October $12 \sim 22$. 174 2017 (Fig. 1). In each unit, approximately 0.5 kg of topsoil (0-5 cm) was collected at four 175 randomly selected sampling sites and then mixed thoroughly to obtain a representative sample. 176 Overall, a total of 120 soil samples were acquired, and each sample was stored in a plastic bag. 177 labeled and sealed. A portable global position system (GPS) was used to determine the 178 coordinates of sampling points. Subsequently, the soil samples were transported to the lab to 179 receive a series of such treatments as sufficient natural air-drying for two weeks and rubbing 180 through a 2 mm sieve to exclude small stones and other impurities. Each sample was divided into 181 two subsamples to be used for spectra collection and physiochemical analysis. 182

183 Each 50 g of soil sample was put into a respective flask, and 250 ml of distilled water (the ratio of water to soil is 5:1) were added into each flask. The water-soluble ion contents were 184 measured in the filtrate obtained from full soaking, oscillation and filtration (Aboukila & Norton, 185 2017). Ca²⁺ and Mg²⁺ were measured using EDTA titration, Na⁺ and K⁺ flame photometry, CO₃²⁻ 186 187 and HCO₃⁻ double indicator-neutralization titration, Cl⁻ silver nitrate titration, and SO₄²⁻ EDTA indirect complexometry (Bao, 2000). The content of CO32- was too low (approximately 0) in 188 some soil samples because CO_3^{2-} is liable to integrate with Ca^{2+} and Mg^{2+} as sediment in a weak 189 alkaline solution (Table 1). Coefficient of variation (CV) reflects the degree of discreteness, and 190 191 a positive correlation exists in two variables. The high CV helps to build a robust model (Dai et 192 al. 2015). The grading of CV showed a wide range of variation among different ions, among which the ion contents of K^+ , Na^+ and SO_4^{2-} are over 100%, showing a strong variability, and 193 those of CO₃²⁻, Cl⁻, Ca²⁺, Mg²⁺ and HCO₃⁻ are between 10% and 100%, having a moderate 194 variability. 195

196 Laboratory spectral measurements and pretreatments

197 The soil samples were put into black vessels with a diameter of 10 cm and depth of 2 cm for spectral data collection and the surfaces were smoothed with a straightedge in the laboratory. 198 The spectral data of the soil samples were measured using ASD (Analytical Spectral Devices, 199 Inc., Boulder, CO, USA) FieldSpec®3 spectrometer with spectral range from 350-2500 nm. This 200 instrument is equipped with two sensors whose spectral resolutions are 1.4 nm and 2 nm, for the 201 202 region of 350-1000 nm and 1000-2500 nm, respectively. The spectral data was measured in a dark room with the light sources which have halogen lamps of 50 W, 50 cm from the sample soil 203 surfaces, and 30° incident angle to reduce the effects of external factors to the minimum. The 204 field angle of fiber-optics probe is 5°, and it is 15 cm from the sample soil surface. The light 205 206 source and spectrometer had been fully preheated, and the spectrometer had been corrected with a standardized white panel (99% reflectance) prior to each measurement to reduce measurement 207

- error. Each sample soil was measured in four directions (3 turns, each is 90°), the spectrum was collected five times in each direction, and altogether there were 20 curves of the spectrum (Hong et al. 2018b). These curves were used as the raw spectral reflectance (R_{raw}) after having the arithmetic mean in ViewSpecPro software version 6.0. The gaps of the spectral curves near 1000 nm and 1800 nm were corrected using Splice Correction function (Xiao et al., 2016a).
- The fluctuation would affect the accuracy of subsequent modeling because of such disturbance 213 as the external environment, instrument noise and random error in spectral data collection. In 214 general, a series of effective pretreatment, including smoothing, resampling and transformation 215 216 etc., can eliminate the external noise to some degree, and then enhance the spectral characteristics (Ding et al. 2018). Therefore, it is necessary to pretreat R_{raw} in the following steps. 217 i) The marginal wavelength (350-399 nm and 2401-2500 nm) of higher noise in each soil sample 218 was removed, then remaining spectrum data was smoothed with filter method (window size is 5 219 220 and polynomial order is 2) using Savitzky-Golay (SG) (Savitzky & Golay, 1964) via Origin Pro software version 2017SR2. ii) The spectral data between 400 and 2400 nm was resampled with a 221 10 nm of samplee interval to keep the spectral features and remove redundant information (Xu et 222 al. 2016). A new spectral curve consisting of 200 wave bands was obtained. iii) The precise R_{raw} -223 _{SNV} was obtained by using standard normal variable (SNV) to eliminate the effects of soil particle 224 225 size, surface scattering and baseline shift on the spectrum data (Xiao et al., 2016b; Barnes et al., 1989). The spectral curves of R_{raw} and R_{raw-SNV} are shown in Fig. 2A and 2B. Notably, 226 comparison indicated that the spectral curve in Fig. 2B was much smoother than that in Fig. 2A, 227 228 which made for the subsequent modeling.

229 Gray correlation (GC)

- 230 The GC, as one grey system theory, seeks the primary and secondary relations and analyzes the
- different effects of all the factors in a system (Deng, 1982; Li et al. 2016). Its calculation process
- is as follows: the reference sequence is $X_0 = \{x_0(t), t = 1, 2, \dots, n\}$, the comparative sequence is
- 233 $X_i = \{x_i(t), t = 1, 2, \dots, n\}$, and the formula of the gray correlation degree (GCD) between X_0
- and X_i is

235
$$GCD = \frac{1}{n} \sum_{t=1}^{n} \gamma \left(x_0(t), x_i(t) \right)$$
(1)

236 where
$$\gamma(x_0(t), x_i(t)) = \frac{\min_i \min_i |x_0(t) - x_i(t)| + \rho \max_i \max_t |x_0(t) - x_i(t)|}{|x_0(t) - x_i(t)| + \rho \max_i \max_t |x_0(t) - x_i(t)|}$$

237 ρ is the distinguishing coefficient within [0,1]. ρ was set as 0.1 in this paper.

The inconsistent dimension between the spectral data and the contents of different ions has some effects on the data analysis. Therefore, normalizing the spectral data preprocessing method can reduce these disadvantageous effects (Liu et al. 2015; Wang et al. 2018b). In this paper, the larger the GCD of a certain band is, the closer relation the band and the ion content has, and vice

242 versa.

243 Variable importance in projection (VIP)

The VIP is a variable selection method based on PLSR (Oussama et al., 2012). The explanatory power of the independent variables to the dependent variables is achieved by calculating the VIP score. The independent variables are sequenced according to the explanatory power (Qi et al. 2017). The VIP score for the *j*-th variable is given as:

248
$$\operatorname{VIP}_{j} = \sqrt{\frac{p * \sum_{f=1}^{F} \operatorname{SSY}_{f} * \operatorname{W}_{jf}^{2}}{\operatorname{SSY}_{total} * F}}$$
(2)

Where *p* is the number of independent variables; *m* is the total number of components; SSY_{*f*} is the sum of squares of explained variance for the *f*-th component and *p* the number of independent variables. SSY_{total} is the total sum of squares explained of the dependent variable, and *F* is the total number of components. W_{jf}^2 gives the importance of the *j*-th variable in each *f*-th component. The higher value VIP_j has, the stronger explanatory power the independent variable has over the dependent variable. The VIP scores of independent variables have been recognized as a useful measure to identify important wavelengths when the score is more than 1 (Wold et al.

256 2001; Maimaitiyiming et al. 2017).

257 Model construction and validation

Two thirds of the samples were used for modeling (n = 80) and one third for validation (n = 40) using Kennard-Stone (K-S) to calculate the Euclidian distance among different samples to ensure

260 the statistical characteristics of modeling and the validation datasets resembled that of the whole

261 sample set (Kennard & Stone, 1969).

The PLSR and SVR models were applied to the quantitative inversion of different water-262 soluble salt ion contents in the saline soil in this paper. The PLSR model is a new stoichiometric 263 statistical model. Compared with the traditional multivariate least squares regression (MLSR), 264 PLSR can overcome the multicollinearity among the variables, reduce the dimension, synthesize 265 and filter the information, extract the aggregate variables with the strongest explanatory power in 266 the system, and exclude the noise with no explanatory power (Wold et al. 2001). The optimal 267 fitting model was built using the number of optimal principal components through full cross 268 269 validation. SVR model is a new machine learning method based on the principle of structural risk minimization provided by the statistical learning theory. This model is characterized by its 270

ability of solving such problems as limited sample size, nonlinear data processing and spatial

272 pattern recognition of high-dimension data (Vapnik, 1995). During the modeling in this study,

the type of SVR and kernel were set as epsilon-SVR and linear function, respectively; the penalty parameter *C* and nuclear parameter *g* were acquired by a grid-searching technique and a leave-one-out cross validation procedure. The optimal values of *C* and *g* were selected when the minimum $RMSE_{CV}$ (root mean squared error of cross validation) was produced (Xiao et al. 2016b). The two models were constructed and validated using the Unscrambler software version

- 278 X10.4 (CAMO AS Oslo, Norway)
- 279 Precision indices of determination coefficient of calibration (R_c^2) , determination coefficient of

280 prediction (R_p^2) , root mean squared error (RMSE) and ratio of performance to deviation (RPD)

were used to evaluate the performance of these models. RPD classification was adopted to

facilitate the interpretation of predictive results: a model is considered as excellent when $RPD \ge 2.5$

283 2.5, as very good when $2.0 \le \text{RPD} < 2.5$, as good when $1.8 \le \text{RPD} < 2.0$, and as satisfactory 284 when $1.4 \le \text{RPD} < 1.8$ and can only distinguish between high and low values when $1.0 \le \text{RPD} < 1.8$

- 285 1.4 (Viscarra Rossel et al. 2007). Generally, the most robust model would be the one with the
- largest R_c^2 , R_p^2 (approach to 1) and RPD value and the lowest RMSE value.

287 **RESULTS**

288 Correlation between water-soluble salt ions content and spectral reflectance

- 289 The correlation coefficients (Pearson correlation) between each soil salt ion content and $R_{\text{raw-SNV}}$
- in the range of 400-2400 nm were tested with the significance level of P < 0.01 ($|\mathbf{r}| = 0.234$ or
- above). The curves of correlation coefficients of soil salt ions were plotted in Fig. 3 and the

292 numbers of bands passing the significance test were counted in Table 2.

- The curve patterns of SO_4^{2-} , Cl⁻, Ca²⁺, Mg²⁺, K⁺ and Na⁺ were similar (Fig. 3). From 400 nm to
- about 550 nm, the correlation coefficients rose sharply from negative to positive, moved with a

295 gentle depression until 1400 nm, plummeted and surged up to 1560 nm (among the curves, the

- 296 change of Ca^{2+} was the sharpest), and maintained a relative stable state to 1850 nm. And then
- from 1850 to 2400 nm, dramatic oscillating variations alternated between rise and fall. In the
- intervals of 400-1400 nm and 1850-2400 nm the curve pattern of CO_3^{2-} was similar to that of
- other ions such as SO_4^{2-} . But between 1400 nm and about 1850 nm, the curve took on a unique pattern: sustained oscillating rise. The coefficient curve of HCO_3^{-} displayed a smaller variation,
- 301 smoothly fluctuating between -0.2 and 0.2. The complex variation of the coefficient curves of
- 302 different ions revealed rich spectral information.
- 303 Selection of characteristic wavelength

304 Characteristic wavelength selection based on GC method

305 The curves of gray correlation degree for soil water-soluble salt ions content and $R_{\text{raw-SNV}}$ were

shown in Fig. 4. The correlation coefficient curves of the seven ions except CO_3^{2-} resembled those of the GCD of the $R_{raw-SNV}$. Generally, the curves exhibited patterns of "oscillatory rise, fluctuation, rapid rise and fall, and oscillatory fluctuation". The gray correlation curves of CO_3^{2-} followed a pattern of "ascending, plummeting, and smooth transition". The analysis of the GC curve amplitude showed the amplitudes of Cl⁻, Mg²⁺ and Ca²⁺ were relatively large, and those of Na⁺, SO₄²⁻, K⁺ and HCO₃⁻ were relatively small, and that of CO₃²⁻ was relatively gentle.

The order of the maximal GCD was: $Cl^{-}(0.561) > Mg^{2+}(0.559) > Ca^{2+}(0.551) > Na^{+}(0.508)$ 312 $> SO_4^{2-}(0.494) > K^+(0.470) > HCO_3^{-}(0.465) > CO_3^{2-}(0.416)$. To ensure that each salt ion had 313 sensitive bands as far as possible, the GCD threshold value was set as 0.40 to select the 314 wavelength. The sensitive band was counted through gray correlation method (Table 3). The 315 numbers of sensitive bands of different ions could be sequenced from the largest to the smallest 316 as follows: $Mg^{2+}(110) > HCO_3^{-}(105) > Cl^{-}(101) > Ca^{2+}(53) > Na^{+}(36) > SO_4^{2-}(21) > K^{+}(15)$ 317 $> CO_3^{2-}$ (14). Therefore, the orders of sensitive band numbers and maximal GCD values had 318 great difference. Furthermore, the band intervals corresponding to the maximum GCD of 319 different salt ions were as follows: CO₃²⁻ was near-infrared between 1740 and 1750 nm, HCO₃⁻ 320 was green light between 560 and 570 nm, and the rest of six ions were near-infrared between 321 1650 and 1660 nm. 322

323 Characteristic wavelength selection based on SR method

Feature band intervals were selected by stepwise regression method in SPSS software version 23.0 (IBM, Chicago, USA), and the significance levels of variables acceptance and rejection were set at 0.10 and 0.15 (Zhang et al. 2018). The parameter indexes of feature band intervals selection were shown in Table 4 by stepwise regression method at maximum adjusted R^2 .

Great difference existed among the optimal SR models of different ions, and the numbers of band intervals accepted by the model range from 3 to 8 (Table 4). The SR model fitted well with the adjusted R^2 greater than 0.8 when the number of selected independent variables was considered. Meanwhile, SR model of each ion was statistically significant (p<0.001). Therefore, the band intervals selected by the SR models were used as the independent variables of PLSR and SVR models.

334 Characteristic wavelength selection based on VIP method

335 Curves of VIP scores of soil water-soluble salt ions content and $R_{raw-SNV}$ were shown in Fig. 5.

Max VIP scores and band intervals obtained from VIP method of soil water-soluble salt ions content and $R_{\text{raw-SNV}}$ were shown in Table 5.

- The curves patterns of seven ions were similar except HCO_3^- (Fig. 5). These curves exhibited
- violent oscillation in the intervals of 400-800 nm and 1900-2400 nm, gentle transition between
- 340 800 nm and around 1400 nm, and fluctuant rise from 1400 to 1900 nm. In contrast, the curve of
- 341 HCO₃⁻ showed oscillatory rise from 400 to 1400 nm, a "U" shaped motion from 1400 to 1900

- 342 nm or so, and a rapid fall and oscillation to 2400 nm. The numbers of sensitive bands based on
- 343 VIP method displayed the following sequence: $Cl^{-}(85) > Na^{+}(83) > HCO_{3}^{-}(79) > SO_{4}^{2-}(74) >$
- 344 Mg²⁺ (69) = Ca²⁺ (69) = K⁺ (69) > CO₃²⁻ (67). The sequence of the maximal VIP scores was
- 345 $HCO_3^{-}(2.37) > CO_3^{2-}(2.01) > Ca^{2+}(1.97) > SO_4^{2-}(1.74) > K^+(1.73) > Na^+(1.55) > Mg^{2+}(1.49)$
- $346 > Cl^{-}$ (1.42). The spectral interval of the maximal VIP scores of Cl⁻ was from 560 to 570 nm,
- 347 Ca^{2+} , CO_3^{2-} and HCO_3^{-} were concentrated between 1410 and 1450 nm; and K⁺, Mg^{2+} , Na^+ and
- $348 \quad SO_4^{2-}$ were from 1870 to 1890 nm.

349 Construction and analysis of PLSR model

- The sensitive bands were obtained using different band selection methods of GC, SR and VIP to build PLSR model. The results of PLSR model were shown in Table 6.
- The models of the six ions Ca^{2+} , Cl^- , CO_3^{2-} , Mg^{2+} , Na^+ and SO_4^{2-} performed well using VIP
- 353 method (R_c^2 is close to 1). The models based on the bands of Ca²⁺, Cl⁻, Mg²⁺, Na⁺ and SO₄²⁻
- 354 selected using SR method displayed good fitting effect, and those of Ca²⁺, Mg²⁺ and Na⁺ using
- 355 GC method exhibited good fitting effect.
- In terms of verification accuracy, VIP method had excellent prediction of Ca^{2+} , Na^+ , SO_4^{2-} , SR method had excellent prediction of Ca^{2+} , Mg^{2+} , Na^+ , SO_4^{2-} (the RPD of Ca^{2+} was up to 3.95), and GC method did not show strong prediction power over any ions. On the contrary, all the three models demonstrated poor forecasting power over HCO_3^- . The RPDs of SR- HCO_3^- and VIP- HCO_3^- were 0.64 and 0.93 respectively. Therefore, VIP method had the best modeling effect and SR method had the best forecasting effect, and GC method had poor modeling and forecasting effects on the salt ions inversion in the PLSR model.

363 Construction and analysis of SVR model

- 364 The sensitive bands were obtained by using different band selection methods of GC, SR and VIP
- to build SVR model. The results of SVR model were shown in Table 7.
- The modeling accuracy of SVR model was similar to that of PLSR model. But the verification
- 367 accuracy of ions was different between the two models. VIP method had the excellent prediction
- of Ca²⁺, Cl⁻, Mg²⁺, Na⁺, SR method had the excellent prediction of Ca²⁺, Mg²⁺, Na⁺, SO₄²⁻, and
- 369 GC method did not show strong prediction power over any ions. The prediction results of Ca^{2+}
- were the best: the RPD of VIP and SR models were 3.93 and 3.97, respectively. Overall, in the
- 371 SVR model, VIP method exhibited the best performance for modeling and predicting the salt
- ions content, SR method was the second, and GC method was relatively poorer.

373 **DISCUSSION**

374 Comparison among the results of different salt ions content in estimating

- 375 The optimal band selection method varied in some degree from the optimal modeling method
- 376 (Table 6 and 7). The comparison was made between the measured value and the estimated value

of all the ions concerned under the optimal model (Fig. 6). The sequence of the forecasting power of the ions was $Ca^{2+} > Na^+ > Cl^- > Mg^{2+} > SO_4^{2-} > CO_3^{2-} > K^+ > HCO_3^-$, and it was the same as that of the modeling power.

Obviously, the verification result showed that most data points of the five ions, Ca²⁺, Na⁺, Cl⁻, 380 Mg^{2+} and SO_4^{2-} , were concentrated near line 1:1. The optimal models of these five ions had very 381 strong predicative power with the RPD above 2.5 (Tables 6 and 7). Compared with the previous 382 researches, model prediction effects of K⁺ and Na⁺ (Qu et al. 2009); Ca²⁺, Na⁺ and Mg²⁺ 383 (Viscarra Rossel & Webster, 2012); SO₄²⁻, HCO₃⁻, Ca²⁺, Cl⁻, Mg²⁺ and SO₄²⁻ (Dai et al. 2015); 384 HCO_{3}^{-} , Ca^{2+} and SO_{4}^{2-} (Peng et al. 2016); K⁺, Na⁺, Ca²⁺ and SO_{4}^{2-} (Wang et al., 2018a) were 385 satisfactory. Although the results of this study are not exactly the same as these previous 386 387 researches, it still shows the rationality own to some extent. In addition, this result shows that band selection has realized the goal of removing the irrelevant information, and plays a major 388 389 role in improving the inversion accuracy of salt ions.

In Figure 6, the data points of CO_3^{2-} and K⁺ were relatively dispersed in the verification result. 390 The CO_3^{2-} had a relatively good predictive power (RPD = 1.80) and the K⁺ had a normal 391 predictive power (RPD = 1.43). Notably, HCO_3^- had no predicative power (RPD = 0.96) because 392 393 the slope were under the 1:1 line and the data points were most discrete (Figs. 6D). The 394 predicting effect of HCO_3 was different from that of Peng et al. (2016) and Dai et al. (2015), but similar to that of Wang et al. (2018a). The cause of this result needs to be further studied. Overall, 395 it is vital to make some efforts to improve the robustness and accuracy of these ion models. Xiao 396 et al. (2016b) failed to predict Na⁺, Mg²⁺ and Ca²⁺, but applied the SVR model to forecasting 397 SAR after the SNV transformation and the performance was satisfactory (RPD = 2.13). 398 Analogously, first derivative reflectance (FDR) index was calculated to effectively predict SAR 399 400 by Xiao et al. (2016a). In addition, Viscarra Rossel & Webster (2012) forecasted the content of Na⁺ after logarithmic pretreatment with VIS-NIR spectral technique (RPD = 2.10). Thus, salt ion 401 indexes construction and variable transformation processing are helpful approaches to improve 402 403 the correlation with the spectra so as to establish satisfactory models.

A little difference existed in the applicability between PLSR and SVR models on inversing the 404 content of ions. Both methods could produce satisfactory results in conformity with that of Peng 405 et al. (2016). In addition, the optimal inversion models and prediction models for each ion were 406 different: SR-PLSR model and SR-SVR model for Ca²⁺, VIP-SVR model and SR-PLSR model 407 for CO32-, SR-PLSR model and VIP-PLSR model for K+, VIP-PLSR model and GC-PLSR 408 model for HCO_3^- , respectively. Among them, the performance of the optimal inversion model of 409 Ca²⁺ resembled that of the prediction model. The results suggested that the ion models with 410 poorer performance frequently demonstrated uncertainty in the inversion process (Peng et al. 411 412 2016). Generally, as the major water-soluble ion components in the two highly soluble salts of

413 sodium and kali, Na⁺ and K⁺ exhibit great difference in the spectral characterization degree (Dai

414 et al. 2015). Therefore, the spectral characters of water-soluble salt ions are not necessarily

415 determined by the number of dissociative ions, so more pertinent experiments and analysis

416 should be conducted to explore the response mechanism.

417 Correlation analysis and inversion performance

The raw spectral reflectance curve of each soil sample presented distinct shapes (Fig. 2A). One of the prime reasons for this phenomenon is that the absorption features in these soil samples were related to soil salt crystal contents and types, as well as various chemical bonds (e.g., C-H, O-H, N-H). The results were in accordance with those in previous studies (Viscarra Rossel et al. 2006; Viscarra Rossel & Webster, 2012; Dai et al. 2015; Peng et al. 2016; Wang et al. 2018a), which demonstrated that soil VIS-NIR spectra could be used to determine part of soil salt ions contents in some degree.

425 Traditionally, correlation analysis helps reveal the relationships between soil salt ions content and VIS-NIR spectra, and it indicates modeling effects to some degree (Weng et al. 2008). In the 426 current research, the number of the significant bands of different ions could be sequenced from 427 the largest to the smallest as follows: Cl⁻ (96%) > Ca²⁺ (95%) > Mg²⁺ (93%) > Na⁺ (90.5%) > K⁺ 428 429 $(89\%) = SO_4^{2-} (89\%) > CO_3^{2-} (73\%) > HCO_3^{-} (0.5\%)$, the correlation coefficients of different ions ranged from the largest to the smallest as: $Cl^{-}(-0.882) > Ca^{2+}(-0.877) > Mg^{2+}(-0.848) >$ 430 Na⁺ (-0.752) > SO₄²⁻ (0.749) > K⁺ (0.630) > CO₃²⁻ (0.552) > HCO₃⁻ (0.235) (Table 2). Thereby, 431 five ions (Cl⁻, Ca²⁺, Mg²⁺, Na⁺ and SO₄²⁻) had more significant relationship with reflectance 432 spectra. Although there were some differences between forecasting power ranking and 433 434 correlation ranking, the optimal models of these five ions had the excellent predictive results (Fig. 6). Nevertheless, the other three ions (K^+, CO_3^{2-}) and HCO_3^{-} had weak correlations and 435 436 unsatisfactory predictive power. In particular, HCO₃⁻ had only one significant band and the worst prediction effects. But in most cases, the sensitive band numbers of HCO₃- were not the least in 437 comparing the results of the three wavelength selection methods (Tables 3-5). Thus, we 438 439 conjecture that the different calculation mechanisms cause a certain inconsistency between modeling performance and sensitivity. In addition, the optimal method of finding out their 440 responding spectrum varies from one ion to another in the soil. In future study, it is practically 441 significant to adopt various methods to select the optimal bands in the inversion of soil ions. 442

443 Effects of wavelength selection on estimation models

444 The massive complex spectra often contain a large amount of redundant information irrelevant to

the ions contents. The selection of feature spectra is hence a critical step to create a robust model.

- 446 From Tables 3-5, we could see the great difference exist in the number of wavelength selected
- 447 with the three methods: VIP method had the largest number of wavelengths $(34.5\% \sim 42.5\%)$,
- 448 SR method had the smallest number of wavelengths $(1.5\% \sim 4\%)$ and number of wavelengths

449 $(7\% \sim 55\%)$ varied greatly by GC method.

Our experiment with three wavelength selection methods also indicated that different methods 450 yielded different results. Among the three methods, the VIP method produced the best results, 451 followed by SR method, while the GC method performed least ideally. We argue that the GC 452 method is not necessarily an inappropriate method as some results are still acceptable. However, 453 GC method could distinguish the primary relationships among the factors in the system by 454 calculating and comparing GCD (Deng, 1982; Liu et al. 2015). In the field of spectral analysis, 455 the application of GC method could better identify sensitive spectral indices, select sensitive 456 457 bands and optimize inversion model (Li et al. 2016). On the other hand, Wang et al (2018b) used 458 GC method to extract the feature bands of soil organic matter content to construct the model with stronger generalization capability. Therefore, the soil compositions have a strong impact on the 459 performance of spectral model. This conclusion is consistent with previous research results 460 461 (Viscarra Rossel et al. 2006; Viscarra Rossel & Webster, 2012; Xiao et al. 2016b). The VIP values were calculated with VIP method, in the process of PLSR analysis to further evaluate the 462 significance of each wavelength for model prediction (Wold et al. 2001; Maimaitiyiming et al. 463 2017; Qi et al. 2017). VIP method often produces the best results in the modeling set because it 464 can distinguish between useful information and inevitable noises in the set. Oussama et al. (2012) 465 adopted this method to reduce almost 75% of the total data set for a simplified model of high 466 accuracy. Additionally, as a simplified regression linear model, SR method not only preserves 467 significant bands but also solves multicollinearity problems effectively (Xiao et al. 2016a; Xiao 468 et al. 2016b). It has great optimization effect on model complexity by adjusting the significance 469 470 level of selected and excluded variables (Zhang et al. 2018). Compared with the selection results with VIP method, SR method could be used to extract fewer bands to establish ions (except for 471 K^+ , CO_3^{2-} and HCO_3^{-}) forecasting models with RPD above 1.80. Therefore, it is meaningful to 472 make further simplification of the model while ensuring its accuracy. 473

474 Research limitations

This study clearly demonstrated that VIS-NIR spectral analysis technique is an effective method 475 476 to detect salt ions content of salinity soil in the irrigated district. In terms of extracting feature wavelengths to estimate ions content, our work provides a comprehensive comparison and 477 evaluation approaches. Such endeavor is critically and practically important to further enhance 478 the model performance of the soil salt ions. The application of machine learning algorithms with 479 480 strong applicability to solve nonlinear relationship between variables, such as Ant Colony 481 Optimization-interval Partial Least Square (ACO-iPLS), Recursive Feature Elimination based on Support Vector Machine (RF-SVM), and Random Forest (RF) has been proved to be a useful 482 approach to obtain the effective information of soil organic matter (Ding et al. 2018). To further 483 improve the prediction accuracy, the more machine learning algorithms should be applied to the 484

analysis of sensitive spectral regions and the construction of stable models in future study. In
addition, the application of multi-source remote sensing platforms such as Landsat, GaoFen-5,
Hyperion and unmanned aerial vehicle (UAV) in soil salt ions estimation has not been
investigated. Therefore, further research should focus on the possible combination of multiple
approaches and remote sensing data at different scales to estimate soil salt ions content.

490 CONCLUSIONS

491 This study investigated the feasibility of estimating soil water-soluble salt ions content via VIS-NIR spectral model. Different methods were applied to the selection of response bands interval 492 to construct robust inversion models. Among them, VIP method could select larger number of 493 wavebands with the highest accuracy, SR method could select the smallest number of wavebands 494 495 with good accuracy. However, the number of wavebands obtained using GC method varied greatly with poor accuracy. The PLSR and SVR models achieved good effects on the modeling 496 and forecasting of most ions content. Moreover, the PLSR model was slightly more than the 497 SVR model in terms of the number of ion models with good predictive effects (RPD over 2.0). 498 499 The models of Ca^{2+} , Na^+ , Cl^- , Mg^{2+} and SO_4^{2-} displayed the highest prediction accuracy, and the RPDs were 3.97, 3.15, 2.98 and 2.75, respectively, while those of other ions were poor. Overall, 500 the best wavelength selection methods, models and inversion results of soil salt ions were 501 different. In the future, the combination of band selection methods and spectral model will have 502 503 a great potential for predicting some soil salt ions content in the salinization area. Such an approach can be utilized to assist decision makers toward the determination of soil salinization 504 levels. 505

506 ACKNOWLEDGEMENTS

507 The authors want to thank A.P. Junying Chen for her help in language standardization of this 508 manuscript and providing helpful suggestions. We are especially grateful to the reviewers and 509 editors for appraising our manuscript and for offering instructive comments.

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Table 1(on next page)

Descriptive statistics of soil water-soluble salt ions content.

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Statistical index	Minimum/ (g•kg ⁻¹)	Maximum/ (g•kg-1)	Mean/ (g•kg ⁻¹)	Standard deviation	Coefficient of variation/%
CO32-	0.000	0.066	0.020	0.020	98.86
HCO ₃ -	0.171	0.666	0.316	0.099	31.27
SO4 ²⁻	0.047	40.892	9.073	10.828	119.34
Cl-	0.145	23.234	4.825	4.711	97.65
Ca ²⁺	0.08	4.111	0.697	0.669	95.95
Mg^{2+}	0.039	1.952	0.706	0.606	85.91
\mathbf{K}^{+}	0.001	5.727	0.936	1.358	145.14
Na ⁺	0.016	23.035	5.014	5.563	110.94

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Table 2(on next page)

Max correlation coefficient and band intervals of soil water-soluble salt ions content with standard normal variable reflectance.

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Water-soluble salt ions	Number of significant bands	Maximum correlation coefficient	Maximum correlation band intervals/nm
Ca ²⁺	190	-0.877	1940~1950
Cl-	192	-0.882	1990~2000
CO ₃ ²⁻	146	0.552	1870~1880
HCO ₃ -	1	0.235	2200~2210
K^+	178	0.630	1850~1860
Mg^{2+}	186	-0.848	1990~2000
Na ⁺	181	-0.752	2010~2020
SO4 ²⁻	178	0.749	1860~1870

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Table 3(on next page)

Max gray correlation degree and band intervals of soil water-soluble salt ions content with standard normal variable reflectance.

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Water-soluble salt ions	Sensitive band numbers	Maximum gray correlation degree	Maximum gray correlation degree intervals/nm
Ca ²⁺	53	0.551	1650~1660
Cl	101	0.561	1650~1660
CO ₃ ²⁻	14	0.416	1740~1750
HCO3 ⁻	105	0.465	560~570
K^+	15	0.470	1650~1660
Mg^{2+}	110	0.559	1650~1660
Na ⁺	36	0.508	1650~1660
SO4 ²⁻	21	0.494	1650~1660

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Table 4(on next page)

Parameter indexes of feature band intervals selection by stepwise regression method.

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Water-soluble salt ions	Sensitive band numbers	Band intervals/nm	Adjusted R ²	Standard error	Sig.
Ca ²⁺	7	1040~1050, 1090~1100, 1900~1910, 1920~ 1930, 2200~2210, 2310~2320, 2370~2380	0.942	0.529	<0.001
Cl	8	730~740, 910~920, 1890~1900, 1970~ 1980, 1990~2000, 2180~2190, 2200~2210, 2290~2300	0.975	1.063	<0.001
CO ₃ ²⁻	4	1280~1290, 1360~1370, 1380~1390, 1420~ 1430	0.836	0.012	< 0.001
HCO3 ⁻	3	2200~2210, 2260~2270, 2290~2300	0.934	0.085	< 0.001
K ⁺	6	740 \sim 750, 810 \sim 820, 1160 \sim 1170, 1890 \sim 1900, 2210 \sim 2220, 2390 \sim 2400	0.817	0.706	< 0.001
Mg^{2+}	6	1130~1140, 1930~1950, 1990~2000, 2100~ 2110, 2170~2180	0.973	0.152	< 0.001
Na ⁺	6	740 \sim 750, 820 \sim 830, 1860 \sim 1870, 2210 \sim 2220, 2260 \sim 2270, 2390 \sim 2400	0.942	1.812	< 0.001
SO_4^{2-}	6	610~620, 1140~1150, 1960~1970, 2210~ 2220, 2290~2300, 2390~2400	0.947	3.255	< 0.00

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Table 5(on next page)

Max VIP scores and band intervals of soil water-soluble salt ions content and standard normal variable reflectance.

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Water-soluble salt ions	Sensitive band numbers	Maximum VIP scores	Maximum VIP scores intervals/nm
Ca ²⁺	69	1.97	1440~1450
Cl-	85	1.42	560~570
CO ₃ ²⁻	67	2.01	1440~1450
HCO ₃ -	79	2.37	1410~1420
K^+	69	1.73	1880~1890
Mg^{2+}	69	1.49	1870~1880
Na ⁺	83	1.55	1880~1890
SO4 ²⁻	74	1.74	1880~1890

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Table 6(on next page)

Calibration and validation results of soil water-soluble salt ions content from the PLSR inversion models using GC, SR and VIP wavelength selection methods.

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Wavelength selection methods	Water-soluble salt ions Latent variab		Calibration sets	Validation sets		
			R_c^2	R_p^2	RMSE/(g·kg ⁻¹)	RPD
	Ca ²⁺	7	0.897	0.724	0.362	1.71
	Cl-	7	0.796	0.565	3.150	1.35
	CO ₃ ²⁻	5	0.660	0.649	0.012	1.21
Conservation	HCO ₃ -	7	0.646	0.285	0.088	0.96
Gray correlation	K^+	1	0.388	0.258	1.209	0.85
	Mg^{2+}	6	0.891	0.767	0.295	1.99
	Na ⁺	7	0.840	0.805	2.589	1.88
	SO4 ²⁻	4	0.561	0.360	8.711	0.87
	Ca ²⁺	7	0.965	0.937	0.168	3.95
	Cl-	2	0.861	0.729	2.434	1.80
	CO ₃ ²⁻	4	0.685	0.742	0.010	1.80
	HCO ₃ -	3	0.340	0.154	0.094	0.64
Stepwise regression	K^+	5	0.722	0.563	0.931	1.37
	Mg^{2+}	4	0.933	0.849	0.236	2.52
	Na ⁺	3	0.901	0.868	2.145	2.67
	SO4 ²⁻	5	0.918	0.889	3.807	2.75
	Ca ²⁺	3	0.909	0.865	0.249	2.57
	Cl-	4	0.930	0.862	1.725	2.48
	CO ₃ ²⁻	9	0.865	0.617	0.012	1.44
Variable importance in	HCO ₃ -	9	0.704	0.263	0.090	0.93
projection	K^+	5	0.664	0.566	0.945	1.43
	Mg^{2+}	3	0.910	0.840	0.243	2.34
	Na ⁺	8	0.939	0.902	1.801	3.15
	SO4 ²⁻	8	0.919	0.872	4.038	2.75

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Table 7(on next page)

Calibration and validation results of soil water-soluble salt ions content from the SVR inversion models using GC, SR and VIP wavelength selection methods.

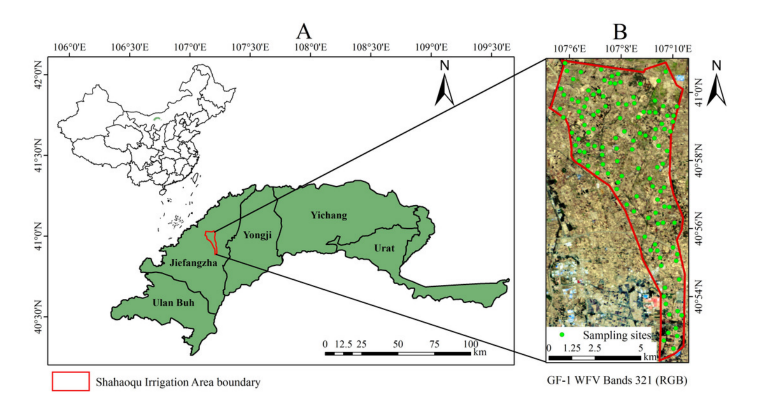
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Wavelength selection methods	Water-soluble salt ions	Calibration sets		Validation sets		
		R_c^2	R_p^2	RMSE/(g·kg-1)	RPD	
	Ca ²⁺	0.910	0.752	0.337	1.73	
	Cl-	0.652	0.500	3.275	1.05	
	CO3 ²⁻	0.688	0.664	0.012	1.14	
Gray correlation	HCO ₃ -	0.563	0.328	0.083	0.70	
Gray correlation	K^+	0.421	0.269	1.155	0.61	
	Mg ²⁺	0.934	0.781	0.289	2.07	
	Na ⁺	0.809	0.764	2.851	1.85	
	SO4 ²⁻	0.565	0.397	9.046	0.52	
	Ca ²⁺	0.964	0.940	0.164	3.97	
	Cl-	0.893	0.790	2.186	2.15	
	CO3 ²⁻	0.605	0.583	0.013	1.16	
Stepwise regression	HCO ₃ -	0.327	0.164	0.095	0.56	
Stepwise regression	K^+	0.717	0.578	0.874	1.26	
	Mg^{2+}	0.936	0.875	0.214	2.75	
	Na ⁺	0.903	0.864	2.171	2.61	
	SO4 ²⁻	0.915	0.893	3.862	2.71	
	Ca ²⁺	0.960	0.935	0.173	3.93	
	Cl-	0.949	0.897	1.483	2.98	
	CO3 ²⁻	0.883	0.664	0.012	1.56	
variable importance in	HCO ₃ -	0.669	0.280	0.088	0.91	
projection	K^+	0.645	0.565	0.888	1.23	
	Mg^{2+}	0.965	0.877	0.214	2.51	
	Na ⁺	0.958	0.872	2.211	2.76	
	SO4 ²⁻	0.914	0.865	4.106	2.48	

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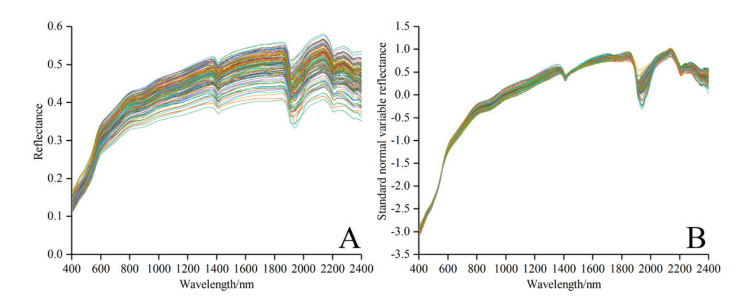
Distribution of sampling sites in the study area.

(A) Location map of Shahaoqu Irrigation Area. (B) Sampling location in Shahaoqu Irrigation Area.

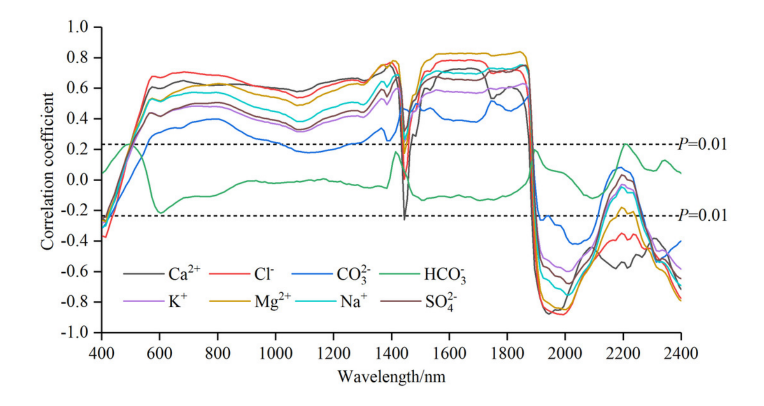


Spectral curves of all soil samples.

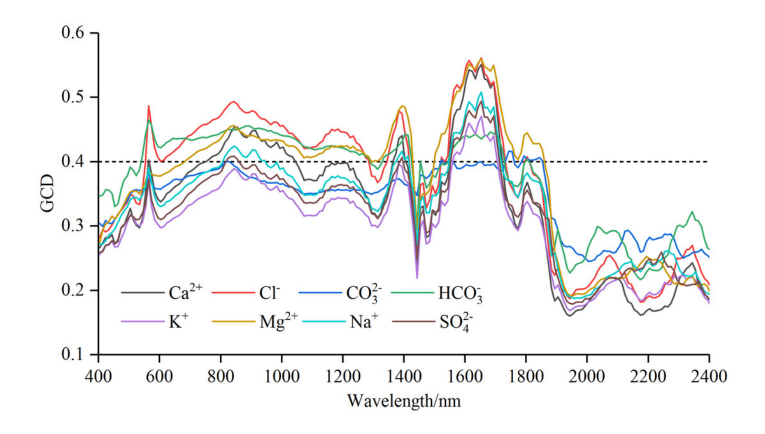
(A) Reflectance spectral curves. (B) Standard normal variable reflectance curves.



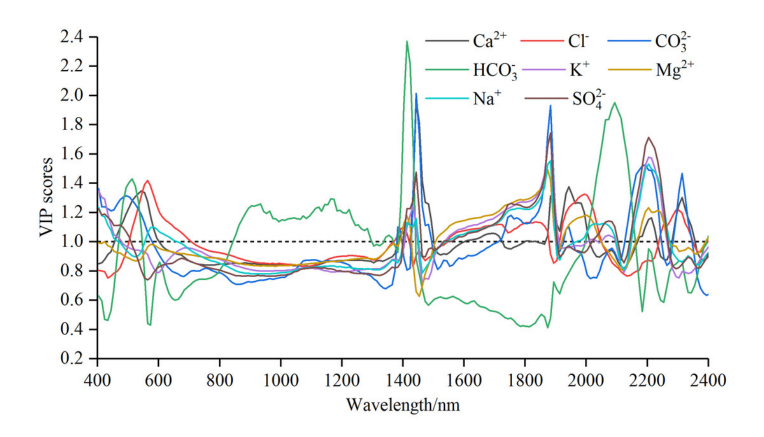
Correlation coefficients of soil water-soluble salt ions content with standard normal variable reflectance.



Gray correlation degree (GCD) for soil water-soluble salt ions content with standard normal variable reflectance.



The Variable importance in projection (VIP) scores for soil water-soluble salt ions content with standard normal variable reflectance.



Validation of soil water-soluble salt ions content based on the best model.

(A) Ca^{2+} with SR-SVR model. (B) Cl^{-} with VIP-SVR model. (C) CO_{3}^{2-} with SR-PLSR model. (D) HCO_{3}^{-} with GC-PLSR model. (E) K⁺ with VIP-PLSR model. (F) Mg²⁺ with SR-SVR model. (G) Na⁺ with VIP-PLSR model. (H) SO_{4}^{2-} with VIP-PLSR model.

