INTRODUCTION

The desert riparian ecosystem, as a groundwater-dependent ecosystem, protects the stability of the environment and provides critical ecosystem services to organisms in arid regions (Décamps et al., 2004; Ding et al., 2017; Stromberg, McCluney, Dixon, & Meixner, 2013). In arid inland river basins, particularly in the lower reaches, desert riparian forests composed mainly of Populus euphratica Oliv. and Tamarix spp. are the core components of the riparian ecosystem, providing key habitats for many different species and functioning as a critical ecological defense against sandstorms and desertification (Ding et al., 2017; Zhu et al., 2016). In these groundwater-dependent...
ecosystems, the groundwater dynamics are the key factor in the shaping and functioning of riverine ecosystems, and groundwater changes are likely the crucial determinant of the vegetation distribution of the desert riparian forest ecosystem (Tamea, Laio, Ridolfi, D’Odorico, & Rodriguez-Iiturbe, 2009). Due to global climate change and human disturbance, scarcity or variable distribution of water and nutrients make desert riparian forests highly sensitive to change, and alterations, particularly alterations in water condition, can have significant impacts on the desert riparian forest vegetation (Stromberg, Beauchamp, Dixon, Lite, & Paradzick, 2007; Tamea et al., 2009; Zhu et al., 2016).

Water availability is the major factor determining the structure and function of desert riparian ecosystems in arid inland river basins (Stromberg et al., 2013). Currently, the desert riparian vegetation-water relationship has been widely studied in eco-hydrological research (Chen et al., 2008; Loheide & Gorelick, 2007; Stromberg et al., 2013). Plant leaves, as the main organs of photosynthesis, play a key role in determining plant survival and productivity in ecosystems, and their nutrient traits, including leaf carbon, leaf nitrogen, leaf phosphorus, and leaf potassium, are closely related to the structure and function of ecosystems (Güsewell, 2004; Wright et al., 2004).

In arid regions, water conditions (e.g., precipitation) may alter the leaf nutrient traits, mainly through plant-soil feedback, and the leaf nutrient traits also change according to differences in species composition (Wang et al., 2015; Zhang et al., 2018a). Leaf decomposition and nutrient uptake and assimilation drive biogeochemical cycling to a certain degree, and these processes of resource investment and reinvestment are inherently survival strategies adapted to a chronic drought environment for plants in arid areas (Drenovský & Richards, 2004; Simões, Calado, Madeira, & Gazarini, 2011). Therefore, quantification of the patterns and the main driving factors for leaf nutrient traits is critical for revealing the patterns of nutrient variation in arid environments and forecasting the response of desert riparian forests to the changes of water condition induced by global climate change.

Ecological stoichiometry, as an emerging comprehensive approach, yields new insights for researching the complex relationships of C/N/P stoichiometric traits at different ecosystem and spatial scales (Elser, Sterner, et al., 2000; Han, Fang, Guo, & Zhang, 2005; Reich & Oleksyn, 2004; Sterner & Elser, 2002). Elser, Fagan, et al. (2000) observed that plants survive in a relatively wide range of nutrient contents in terrestrial ecosystems, and the concentrations of C, N, P, and their stoichiometric ratios within plants to an extent reflect how a plant has adapted to the local habitat (Elser, Fagan, Kerkhoff, Swenson, & Enquist, 2010). Previous studies on arid and semiarid regions suggest that plant growth is mainly limited by the N element (Wang et al., 2015; Zhang et al., 2018a), but in other studies, P-limited growth has also been detected (Zheng & Shangguan, 2007). These differences may be due to differences in nutrient limitations in different areas (Zhao et al., 2014). Currently, there has been increasing investigation on the patterns of leaf C/N/P stoichiometry and their relationships with climatic and geographical factors (Kerkhoff, Enquist, Elser, & Fagan, 2005; McGroddy, Daufresne, & Hedin, 2004; Reich & Oleksyn, 2004; Zhao et al., 2014). However, very few studies have addressed how the leaf nutrient traits of dominant riparian species respond to variations in groundwater depth, and in particular the relationship between groundwater depth and nutrient stoichiometry along a natural groundwater depth gradient in a hyperarid region.

The downstream of Heihe River in northwestern China is an ideal area for research on the response of plants to environment due to there are different natural desert riparian communities along groundwater depth gradient induced by river flow (Ding et al., 2017; Zhu et al., 2016). Groundwater provided by the river is considered the key factor in affecting vegetation growth and distribution (Chen et al., 2008; Tamea et al., 2009). However, exactly how groundwater depth and soil properties affect leaf nutrient traits along a moisture gradient remains largely unclear at the local scale in this hyperarid region, and further research is needed to reveal the mechanism of nutrient utilization and uptake and the distribution patterns of leaf nutrient traits of desert riparian vegetation, which will be valuable for restoring and managing desert riparian forest ecosystems in response to climate change.

In this study, 11 sites along a groundwater depth gradient were selected to examine responses of the leaf nutrient traits of dominant desert riparian species to groundwater depth in a desert riparian ecosystem. Our study was designed to (a) examine the patterns of leaf nutrient traits of dominant species along a natural groundwater depth gradient; (b) assess the relationship of leaf nutrient traits and soil physicochemical properties; (c) disentangle the main drivers affecting the variation of leaf nutrient traits. We hypothesized that groundwater depth play a key role on leaf nutrient traits, soil properties especially soil salinity and alkalinity may have important effect on leaf nutrient traits at specific distance from river.

2 MATERIALS AND METHODS

2.1 Study sites

The Heihe River is the second largest inland river in China, with a length of 821 km. The upper reaches are covered with alpine glaciers and permafrost, becoming the main source of regional runoff; runoff is primarily consumed in the middle reaches due to extensive deserts and farmland (Ding et al., 2017; Peng, Xiao, & Xiao, 2013). Over the past decades, anthropogenic grazing and farming in the middle reaches have led to the disappearance of most of the downstream surface runoff, lakes drying up, and groundwater depth increasing sharply, which has resulted in significant degradation of the desert riparian forest of P. euphratica (Zhu et al., 2016). A water conveyance project was implemented beginning in 2000; regular water conveyance has already caused groundwater levels to rise, and the riparian vegetation has correspondingly been restored (Ding et al., 2017; Fu, Chen, & Li, 2014). Our study transect was established in a desert riparian forest ecosystem along a groundwater depth gradient in the downstream Heihe River, China (Figure 1). The region has a hyperarid desert
climate, with an annual rainfall of approximately 37.4 mm, while the mean annual pan evaporation is 3,467.56 mm; more than 75% falls during the growing season (July to September), and the annual temperature is 8.57°C (Fu et al., 2014; Li, Yu, Li, & Zhao, 2016). The elevation changes relatively little from 921 m to 925 m. Groundwater provided by the Heihe River is the major water source available for maintaining local ecosystem stability (Li et al., 2016). The regional soil is gray-brown desert soil. The vegetation types generally shift from riparian forest dominated by *P. euphratica* and *Tamarix ramosissima* Ledeb. to desert scrub dominated by *Reaumuria songarica* (Pall.) Maxim. along the river channel (Fu et al., 2014).

### 2.2 Experimental design and data collection

In early August 2015, eleven sampling sites were selected in a desert riparian ecosystem and sampled at the distances of 300, 800, 1,300, 2,200, 2,450, 2,700, 2,950, 3,200, 3,700, 4,000, and 4,500 m from Heihe River (Figure 1). Our field survey was conducted in one week. Due to the regular water conveyance project, floodwater only impacts the areas near the river bank, and the groundwater depth is relatively stable in the growing season (Ding et al., 2017; Fu et al., 2014). Groundwater depth data were obtained using a Hobo groundwater depth gauge from seven monitoring wells in 2010–2014 (http://westdc.westgis.ac.cn/), which located at 300, 2,200, 2,700, 3,200, 3,700, and 4,300 m from Heihe River. Apart from sites 1, 4, 6, 8, and 9, the groundwater depth data at other sites were obtained by cokriging interpolation method (Ahmadi & Sedghamiz, 2008; Hoeksema et al., 1989; Zhang et al., 2018b). At each site, three 5 m × 5 m plots were established randomly, where we collected leaf samples and soil samples (0–50 cm).

Along the groundwater depth gradient from S1 to S11, the dominant species were *T. ramosissima* (S1–S7, S9), *Karelinia caspica* (Pall.) Less. (S8), and *R. songarica* (S10, S11). At each site, fully expanded and healthy leaves were sampled from the dominant species in each plot, and then fresh weight of more than 100 g were gathered in envelopes and kept in an ice chest for further analysis. Leaf samples were grinded into a fine powder using a high speed mixer mill (MM400, Retsch, Haan, Germany) for nutrient analysis.

At each site, soil cores were collected every 10 cm at a soil depth of 50 cm, with three replicates. Soil moisture was determined gravimetrically via oven drying at 105°C for 72 hr. Soil bulk density was analyzed using a cutting ring (100 cm³ in volume) by the cutting ring method at 10, 20, 30, 40, 50 cm deep. Soil samples were air-dried and then passed through a 2 mm sieve for chemical analysis.

Soil electrical conductivity and pH were analyzed through 1:5 and 1:1 soil:water suspensions using a portable electronic probes (Multiline F/SET-3, WTW, Weilheim, Germany), soil available P and K content were determined according to the Olsen method and atomic absorption spectroscopy, respectively. Leaf C, Soil total C, leaf N, and soil total N were extracted by an elemental analyzer (Vario EL III, Elementar, Hanau, Germany). Leaf P and leaf K were analyzed by inductively coupled plasma optical emission spectrometer (iCAP 6300, Thermo Scientific, Waltham, USA).

### 2.3 Statistical tests

All species importance values determined the dominant species in each plant community and were obtained from the following formula (Curtis & McIntosh, 1951):

$$IV = \frac{(RC + RD + RF)}{3}$$

where IV is the importance value of the species, RC is the relative coverage, RD is the relative density, RF is the relative frequency.

The list of species composition and importance values in the studied area are shown in Supporting Information Table S1. At each site, groundwater depth and leaf nutrient traits were presented as the mean ± SD (Supporting Information Table S2). The gravimetric analysis of soil moisture was expressed as the mean value for the layers of 0–30 cm and 30–50 cm; other physicochemical properties were expressed as the mean values for the layers at 0–50 cm soil depth.

Pearson’s correlations were performed to determine the relationships between leaf nutrient traits and soil properties according to the level of significance. A generalized linear model with sampling location (distance from river) as a random effect was used to examine...
the relative importance of soil variables for the leaf nutrient traits, and one-way ANOVA was used to explore the differences in leaf nutrient traits among different sampling sites, Tukey’s test was used to determine statistically significant differences; homogeneity of variance and normality for all leaf nutrient traits data were assessed and calculated from the log10 transformation. Appropriate regression analysis was used to examine the relationships between leaf nutrient traits and groundwater depth according to high $R^2$ values and the level of significance. Significant differences were evaluated at $p < 0.05$, and all statistical analyses were performed using SPSS 22.0.

A redundancy analysis (RDA) was applied to determine the key environmental factors of the variation of leaf nutrient traits using soil properties and groundwater depth data. First, the marginal and conditional effects of environmental variables were obtained from forward selection in the redundancy analysis. Statistical testing for added environmental explanatory variables was performed with Monte Carlo tests (9,999 permutations) (Ter Braak & Smilauer, 2012). Compared to Marginal effects, conditional effects represented the effects of the explanatory variables on the leaf nutrient traits after the previous variable was eliminated through forward selection (Ding et al., 2017; Ter Braak & Smilauer, 2012). After the RDA, the key variables ($p < 0.05$) were included in the group of the key environment variables. Related to our hypotheses, variation partitioning model (VPM) was used to decompose the two groups of significant variables: groundwater depth and soil properties, and quantify the interpretation rate of the variation of leaf nutrient traits by separating the significant variables into the unique effects of each variable and the joint effects between variables (Heikkinen, Luoto, Kuussaari, & Pöyry, 2005). The redundancy analysis, variation partitioning, and forward selection were performed using CANOCO 5.0 for Windows (Ter Braak & Smilauer, 2012).

3 | RESULTS

3.1 | Leaf nutrient trait changes in response to groundwater depth

In all plant species, the average leaf C, leaf N, leaf P, and leaf K were 327.29, 13.88, 0.58, and 6.71 mg/g, respectively, and the leaf C/N, leaf N/P, and leaf C/P ratios were 24.41, 26.12, and 614.94, respectively (Supporting Information Table S2).

The ANOVA results showed significant differences for leaf C ($F = 12.437, p < 0.001$) among the different sampling sites. Leaf C ranged from 173.71 to 381.34 mg/g for different sampling sites (Supporting Information Table S2). After the RDA, the key variables ($p < 0.05$) were included in the group of the key environment variables. Related to our hypotheses, variation partitioning

![FIGURE 2](Image)

**FIGURE 2** The variation of leaf nutrient traits with the groundwater depth gradient
The variation partitioning model indicated that groundwater depth, soil pH, and their interaction explained 35% of the variation. The independent effect of groundwater depth had a large contribution, accounting for 20% of the variation in leaf nutrient traits, and the independent effect of soil pH had a small contribution, accounting for 16.4% of the variation in leaf nutrient traits.

4 | DISCUSSION

4.1 | Leaf nutrient traits in desert riparian ecosystem

In the lower reaches of Heihe River, our result showed that the average leaf C of the dominant species was 327.29 mg/g (Supporting Information Table S2), and lower than the mean values of global flora and of Loess Plateau (Elser, Fagan, et al., 2000; Zheng & Shangguan, 2007). Leaf N and leaf P were lower than that in the Gobi desert region of the Heihe River, and lower than the mean values of global flora and of other arid and semiarid regions (Reich & Oleksyn, 2004; Wang et al., 2015; Zhang et al., 2018a; Zheng & Shangguan, 2007). Leaf N and leaf P were lower than that in other regions (Han et al., 2005; Reich & Oleksyn, 2004; Wang et al., 2015; Zheng & Shangguan, 2007). It is suggested that the dominant species was characterized by lower leaf C, leaf N, and leaf P, but higher leaf N/P and leaf C/P. In this study, the lower values of leaf C might be due to drought and salt stress. In this hyperarid region, plants need to reduce their water potential and stomatal conductance to minimize drought and salt stress during the growing season. In this case, photosynthesis may have been restricted, and the salt stress (high soil electrical conductivity) increasing the metabolic costs may have led to decreased C fixation (Chaves, Flexas, & Pinheiro, 2009; Dodd & Donovan, 1999; McCree, 1986). Leaf C dropped rapidly when groundwater depth was higher than 3 m (Figure 2), because R. songarica community suffered serious water stress in this hyperarid region (Fu et al., 2014), thus C fixation deceased, and species difference possibly was another reason, further research is needed in controlled conditions in future. The lower mean values of leaf N and leaf P along the groundwater
depth gradient appeared to be attributable mainly to the dominant species (*T. ramosissima*). Because *T. ramosissima* is in flowering and fruiting stages from July to September, it is undergoing storage and the transfer of genetic material to reproductive organs (Rong, Liu, Xia, Lu, & Guo, 2012); therefore, leaf N and leaf P along the groundwater depth gradient were relatively low. Our results were based on samples taken during the growing season, and leaf N and leaf P may vary seasonally; thus, long-term (e.g., in different seasons or years) field investigation is necessary in the future.

In this study, the high values of leaf C/P might also be attributable mainly to the dominant species (*T. ramosissima* and *R. songarica*), which had a slow growth rate and low photosynthetic rate due to drought (annual precipitation 30–40 mm) and salt stress (high soil electrical conductivity), thereby leading to greater carbon allocation compared with nitrogen and phosphorus allocation to leaves. Our results showed that leaf C/N and leaf C/P in desert riparian region were significantly higher than that in Gobi desert region indicating that the Gobi desert plants grew more rapidly in the growing season due to the limited rainfall; however, the desert riparian vegetation rather defense against habitat stress and differed from the Gobi desert vegetation's growth adaptation strategy in the growing season.

In arid terrestrial ecosystems, scarce precipitation causes poor soil leaching, low loss of P from soil weathering, and a low biodiversity level and vegetation cover, leading to soil P content being relatively abundant compared to soil N content, which could explain why nitrogen is more likely to be the key limiting factor in desert ecosystems (Vitousek & Howarth, 1991; Vitousek, Porder, Houlton, & Chadwick, 2010). These results were also confirmed by our results (Zhang et al., 2018a) and those of other studies (Vitousek et al., 2010; Wang et al., 2015; Wang, Yang, & Ma, 2008). Previous studies on the different ecosystems suggest that leaf N/P < 14 demonstrates N limitation, leaf N/P > 16 demonstrates P limitation, and 14 < leaf N/P < 16 indicates either N or P limitation, or both (Aerts & Chapin, 2000; Koerselman & Meuleman, 1996). Our result showed that the average leaf N/P was 26.12, indicating that dominant desert riparian vegetation was largely constrained by P. This result differs from previous studies in other desert ecosystems (Vitousek & Howarth, 1991; Wang et al., 2015; Zhang et al., 2018a) while is consistent with the results from the Loess Plateau study in China (Zheng & Shangguan, 2007). Some research indicated that P is the main growth-limiting nutrient of plant communities because of low soil phosphorus levels in China (Han et al., 2005). In this hyperarid region, this difference apparently was in response to a combination of factors: first, the dominant species of the community (*T. ramosissima*) is a typical shrub associated with endophytic diazotrophic bacteria, resulting in an increase in N fixation by plants (Xu, Luo, Wang, & Wang, 2014). Second, soil salt stress affects the absorption of P because many Cl\(^{-}\), SO\(_4^{2-}\) and other anions exist in the soil and compete with P, resulting in a decrease in P uptake by plants (Balba, 1995), thereby resulting in a high N/P ratio. Moreover, to survive under the stress of extremely arid environments with high temperature, plants tend to have lower growth rates, which might result in a higher N/P ratio. In fact, some research indicated that the leaf N/P stoichiometric ratio was impacted by plant genetic characteristics.

| TABLE 2 Effects of soil properties on the variations in leaf C, leaf N, leaf P, leaf K, and leaf C/N/P ratios |
|----------------|---------|---------|---------|---------|
| Leaf C         | Leaf N  | Leaf P  | Leaf K  | Leaf C/N |
| SS% p          | SS% p   | SS% p   | SS% p   | SS% p   |
| Soil moisture (0–30 cm) | 1.0 0.623 | 8.6 0.310 | 9.0 0.261 | 0.6 0.944 | 0.0 0.996 |
| Soil moisture (30–50 cm) | 1.3 0.569 | 3.6 0.507 | 8.5 0.274 | 0.0 0.944 | 0.0 0.996 |
| Soil bulk density | 19.7 0.042 | 0.7 0.755 | 4.7 0.430 | 0.3 0.961 | 0.0 0.996 |
| Soil total carbon | 22.5 0.032 | 3.6 0.475 | 1.7 0.621 | 0.6 0.796 | 0.5 0.857 |
| Soil total nitrogen | 36.1 0.009 | 3.9 0.490 | 10.6 0.233 | 1.5 0.670 | 0.2 0.912 |
| Soil available K | 5.1 0.866 | 0.1 0.269 | 1.7 0.780 | 17.3 0.321 | 1.4 0.690 |
| Soil pH | 6.6 0.271 | 8.4 0.317 | 1.5 0.638 | 0.9 0.721 | 0.9 0.796 |
| Soil electrical conductivity | 0.1 0.806 | 3.3 0.756 | 0.5 0.353 | 5.2 0.353 | 0.2 0.853 |

* p < 0.05, ** p < 0.01.
and survival strategies and was closely associated with habitat complexity (Aerts & Chapin, 2000; Kerkhoff et al., 2005; Koerselman & Meuleman, 1996). Due to the low plant diversity in arid regions, so the leaf N/P of dominant species can be indicative of nutrient limitation of the desert riparian plant community, to some extent. Güsewell and Koerselman (2002) reported that the leaf N/P accurately determined the nutrient limitation of plant growth at the community level. Therefore, further detailed field investigations in different seasons and artificial controlled experiment should be performed to reveal the nutrient limitation patterns of different plant functional groups or at the community level.

4.2 Variations of leaf nutrient traits and its influencing factors

Leaf trait–environment relationships are important to explain and predict the underlying mechanisms of the pattern of leaf nutrient traits along environmental gradients and to identify ecosystem nutrient limitations (Güsewell, 2004; Kerkhoff et al., 2005; Reich & Oleksyn, 2004; Zhao et al., 2014). In the Gobi desert region of Heihe River, the patterns of leaf nutrient traits controlled by precipitation were not significant, but soil moisture was better able to explain the variations in leaf nutrient traits (Zhang et al., 2018a). These results indicated that water conditions might be the main driving factor affecting the variations of leaf nutrient traits in arid regions. Our results showed that groundwater depth and soil pH jointly influenced the pattern of leaf nutrient traits in the desert riparian vegetation (Table 3). Apart from groundwater depth, soil properties especially saline-alkali properties may also have had important effect on leaf nutrient traits at local scale in this hyperarid region. In addition, the interactions of groundwater depth and soil pH negatively contributed to the explained variation in leaf nutrient traits; however, the independent effects of groundwater depth and soil pH were positively contributed to explaining this variation. The negative contribution of

| Marginal effects | Conditional effects |
|------------------|---------------------|
| Variables        | Percentage          | Variables        | Percentage |
| Groundwater depth| 21.2                | Groundwater depth| 21.2        |
| Soil pH          | 17.7                | Soil pH          | 17.9        |
| Soil bulk density| 17.3                | Soil available K | 3.9         |
| Soil available K | 12.2                | Soil C/N         | 1.7         |
| Soil electrical conductivity | 12.0 | Soil bulk density | 1.6 |
| Soil total nitrogen| 10.5               | Soil total carbon| 1.5         |
| Soil total carbon | 10.4                | Soil total nitrogen| 1.3 |
| Soil available P | 8.7                 | Soil moisture in 0–30 cm | 0.9 |
| Soil moisture in 0–30 cm | 7.6           | Soil electrical conductivity | 0.9 |
| Soil moisture in 30–50 cm | 7.5           | Soil available P | 0.6         |

The interaction indicated that the majority of the relationships between the two factors (groundwater depth and soil pH) are likely suppressive rather than additive (Helikkinen et al., 2005).

Leaf nutrient traits showed a significant groundwater depth gradient in the desert riparian ecosystem of the Heihe River (Figure 2). Groundwater was the critical water resource for the survival of desert plants, and it also regulated plant growth and nutrient fluxes through interactions with environmental factors (Tamea et al., 2009; Zhang et al., 2018b). In our previous study, soil nutrients decreased significantly.
along the groundwater depth gradient (Zhang et al., 2018b), and leaf nutrient traits were closely associated with soil properties (Table 1); increased groundwater depth and soil nutrient properties affected the leaf C, leaf N, leaf P, and leaf K, as these factors all decreased significantly with groundwater depth. In the desert riparian, vegetation with shallow groundwater, water conditions, and soil nutrients at these sites might be generally sufficient, and more nutrients could be used by plants. With increasing groundwater depth, the depletion of soil nutrients from low nutrient availability in the soil resulted in a decrease of nutrient uptake by plants. In this hyperarid region, soil nutrients might be the main factor affecting the leaf nutrient traits at the local scale; however, the soil nutrient content is regulated by groundwater, and this could explain why groundwater depth was the largest contributor to the leaf nutrient traits in the variation partitioning model. Meanwhile, soil pH was better able to explain the variations in leaf nutrient traits, in our results and together with those from previous studies, suggests that soil saline-alkali imposes a common limitation on plant growth and distribution patterns, largely due to the soil parent material in arid regions (Liu, Wang, & Xi, 2008; Zhou, Chen, & Li, 2010).

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CONFLICTS OF INTEREST

None declared.

AUTHOR CONTRIBUTIONS

YRZ conceived and designed the experiment; TYG and XLZ performed materials and data analysis; XLZ wrote the paper; LHJ, LML, NNG, JHZ, XLZ, WTC, and TYG carried out field investigations.

DATA ACCESSIBILITY

Leaf nutrient traits and soil properties contents data have been deposited in the Dryad Digital Repository. https://doi.org/10.5061/dryad.rg25909.

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