The CSR strategies of alpine plants and community functional diversity in the Eastern Qinghai-Tibetan mountains

Yushuang Wu*, Tatiana G. Elumeeva*, Elena S. Kazantseva**, Yanyan Yan*, Qian Wang*, Ksenia V. Dudova* & Vladimir G. Onipchenko*

ABSTRACT

Many alpine communities of the Tibetan plateau experience both harsh climates and heavy grazing, and the adaptations to such environment are realized in plant functional traits and strategies. We tested, which leaf traits allow to dominate in the two wide-spread alpine communities of the Eastern Qinghai-Tibetan mountains, Kobresia alpine meadows and alpine fens, and do meadows and fens differ by community-weighted dominance-linked traits, strategy affinities, and functional diversity. Lower leaf dry matter content (LDMC) and higher specific leaf area (SLA) provides the species dominance in Kobresia alpine meadows, while size traits do not influence the dominance. Both LDMC and SLA have low functional evenness and divergence in the two communities. The prevail of stress-tolerators reflects the abiotic environment of alpine plants, but high affinities to ruderal strategy are possibly related to grazing.

Keywords: functional traits, functional diversity, CSR strategy, alpine plants

REЗЮМЕ

У Ю., Елумеева Т.Г., Казанцева Е.С., У Я., Дудова К.В., Онипченко В.Г. Эколого-ценотические CSR стратегии альпийских растений и функциональное разнообразие сообществ на востоке Цинхай-Тибетского нагорья. Многие альпийские сообщества Тибетского нагорья существуют в условиях сурового климата и интенсивного выпаса. Адаптации к таким условиям представлены функциональными признаками, растениями и стратегиями. Мы изучили, какие признаки листьев позволяют доминировать на альпийских кобрезиевых лугах и на альпийских болотах — сообществах, которые широко распространены в альпийском поясе востока Цинхай-Тибетского нагорья. Мы проверили, различаются ли биомасса и луга по средневзвешенным признакам, связанным с доминированием, стратегиями и функциональным разнообразием. Более низкое содержание сухого вещества (LDMC) в листьях и более высокая удельная листовая поверхность (SLA) способствуют доминированию видов на альпийских кобрезиевых лугах, тогда как размерные характеристики (площадь и масса листа) не влияют на доминирование. Во взвешенных сообществах признаки LDMC и SLA имеют низкую функциональную выраженность и дивергенцию. Преобладание стресс-толерантов отражает абиотические условия обитания альпийских растений, а большой вклад в рудеральную стратегию, возможно, связан с влиянием выпаса.

Ключевые слова: функциональные признаки, функциональное разнообразие, CSR стратегии, альпийские растения

Plant functional traits and functional types or Grime's strategies affinities, measured by standardized protocols, allow to perform rather easy and informative comparative analysis of plant community structure and composition through various vegetation types and geographical zones (Grime 2001, Pérez-Harguindeguy et al. 2013, Garnier et al. 2016, Pierce et al. 2017). The following approaches are often used to study functional composition: 1) the spectrum of plant traits and functional types (strategies); 2) the community weighted means (CWMs) of plant traits or Grime's strategy affinities (Lavorel et al. 2008). The values of CWMs are usually impacted by dominant species and sometimes differ significantly from non-weighted means of the same traits (Onipchenko et al. 2022); 3) the indices of functional diversity (Mason et al. 2005, Schleuter et al. 2010), the most commonly used indices are functional richness (F(Ric), functional evenness (F(Eve)) and functional divergence (F(Div)). They can be calculated using single traits and as multtrait indices (Laliberté et al. 2014), therefore single trait indices are easier to interpret, as they often better indicate environmental gradients (Butterfield & Suding 2013, Dudova et al. 2019, Onipchenko et al. 2022).

The alpine regions are one of the most extreme environments on the earth. Therefore, alpine species have highly specialized adaptations to such habitats (Körner 1999), and the suits of functional traits, which allow them to withstand the harsh environment of high mountains, were often studied by plant ecologists. Generally, alpine plants experience stress due to high solar radiation, low temperatures, and often low nutrient availability. These factors suggest the stress-tolerators (S) are expected to be the prevailing plant functional type in terms of Grime's CSR
strategies (Grime 2001), as well as the higher affinity to stress-tolerance, calculated by the methods developed by Pierce et al. (2017). Indeed, the dominance of stress-tolerators was observed in the Tibetan plateau (Wang et al. 2018, Zhou et al. 2021, Yu et al. 2022) and increased from humid to arid alpine habitats.

However, the alpine environment is variable due to terrain, which may cause stronger differentiation of vegetation than elevation gradient, that’s why in alpine belts of humid climate the depressions are characterized by late snowmelt time and short, sometimes extremely short, growing seasons. This requires plant adaptations to rather fast growth, attributed to the ruderal (R) strategy (Onipchenko et al. 2022). The depressions, which do not suffer too late snowmelt, have enough rates of mineralization to support species with traits of competitors (Onipchenko et al. 2022).

Besides the harsh environment, the biotic interactions are also important for alpine plant selection, because non-woody alpine communities often support wild mammals and/or livestock grazing. In the areas, which develop as pastures for a considerable time, the selection also will lead to the prevailing of traits, which reflect adaptations to growth under heavy grazing. For example, the disturbance is a second important factor determining CWMs of specific leaf area (SLA) in calcareous grasslands of the Bavarian Alps (Rosbach et al. 2015), while in Tibetan rangelands, CWM SLA was not correlated with grazing intensity, but was positively correlated with precipitation and soil nitrogen content (Jäschke et al. 2020). In Tibetan alpine meadows intermediate grazing induced a shift to the ruderal strategy, while high grazing intensity promoted the dominance of species with higher C-affinities (Wang et al. 2018).

The Tibetan Plateau is an interesting object due to its large areas, covered by alpine vegetation, its high plant diversity and a long history of livestock grazing, mostly by yaks (Bos grunniens Linnaeus 1767). There are an increasing number of studies concerning functional traits of alpine plants in Tibet, including leaf traits and strategies, life form composition, floristic and functional richness, phytomass structure and CWMs of plant traits (Ma et al. 2010, Elumeeva et al. 2014, Onipchenko et al. 2014, Elumeeva et al. 2015a,b, Ren et al. 2015, Li et al. 2017, Wang et al. 2018, Jäschke et al. 2020, Vandvik et al. 2020, Zhou et al. 2021, Liu et al. 2022, Yu et al. 2022). However, the high species richness of the region and its large area suggests the importance of local studies for further incorporation of these data into large-scale regional studies.

Thus, our study aims to contribute to the investigation of alpine communities at the Eastern Qinghai-Tibetan Plateau by exploring the leaf functional traits and functional diversity of two local alpine communities, Kobresia alpine meadows and alpine fens. We tested which leaf traits allow to dominate in the communities and do meadows and fens differ by dominance-linked traits, strategy affinities, and functional diversity.

MATERIAL AND METHODS

Study area

The study was conducted in Mount Kaka (32°98’N 103°67’E). It is located in the West Sichuan Plateau, which is a part of Songpan County, Sichuan province in China (Fig. 1). The altitude of Mount Kaka is around 3950 meters. It belongs to the Min Mountains – the east of the Qinghai-Tibetan Plateau. This area is controlled by warm wet flow coming from southwest and southeast monsoon, it is a typical mountain monsoon climate. The average annual temperature in this area is +5.7º, the average temperature in January is -7.6º, and the average temperature in July is +9.7º. There is no absolute frost-free season and the average annual duration of solar radiation is 1827.5 hours, annual precipitation is 718 mm, and 72 % of precipitations falls from June to August (Hu et al. 2012, Onipchenko et al. 2014).
The timberline in the study area is at 3800–4000 m and the alpine belt is occupied by alpine meadows, fens and shrubs. The Latin names of plants are given by “Flora of China” (Editorial Board of Flora of China 1994–2013).

We mainly studied two types of communities: *Kobresia* alpine meadows (KAMs). These communities occupy gentle southern slopes and ridges. The most abundant herbaceous species (from the most to the least abundant) are *Kobresia humilis* (C.A. Mey ex Trautv.), *Sibirula pauciblens* Linn., *Alchemilla nepalensis* (Spreng.) Hand.-Mazz., *Gentiana beophylla* Maxim. ex Kusnez., *Trollius farrettii* Stapf, *Festuca ovina* L., and *Pilagrostis concinna* (Hook. f.) Roshev. The cover of shrubs, for instance, *Potentilla fruticosa* L., *Caragana jubata* (Pall.) Poir., and *Spiraea alpina* Pall. range from 5 to 10 %. *Kobresia* alpine meadows have a well-developed moss layer, however, the lichens are not abundant in this community. The studied meadows are mainly used as pasture with moderate grazing by yaks (Elumeeva et al. 2014).

**Alpine fens (AFs).** Alpine fens occupy gentle slopes with poor drainage. The coverage of mosses is nearly 35 %. The most abundant vascular plants in AFs are *Carex moorcroftii* Falc. ex Boott, *Carex arctofusca* Schlkuhr, *Kobresia humilis* and *Crematedium lineare* Maxim. These communities are mainly affected by intensive grazing (Elumeeva et al. 2014).

**Leaf traits measurements and CSR strategies calculation**

The field sampling was in July 2012 and July–August 2019. We measured leaf area (LA), leaf fresh mass (LFM), and dry mass (LDM) of 94 plant species, together representing the most common vascular plant species of studied communities at the local site. In 20 species only LA and LDM were measured. Among them, the LFM of *Kobresia humilis*, *Carex cruenta* Nees, and *Carex parva* Nees were derived using regression coefficients of the relationship between LDM and LFM of the *Carex* and *Kobresia* species with flat leaves from open habitats (15 alpine and temperate grassland and fen species, $R^2=0.968, p<0.001$), the leaf dry matter content (LDMC) of *Androsace mariae* Kanitz was obtained from Ma et al. (2010), and that of *Allium sikkimensis* Baker was obtained from Ren et al. (2015). For one species, *Sansuierea stella* Maxim., all leaf measurements were extracted from the publication of Ren et al. (2015) from Maqu station, located at 3600 m. a.s.l.

To identify the plant species, we used Flora of China (Editorial Board of Flora of China. 1994–2013), the Moscov Digital Herbarium (Seregin 2022), National Plant Specimens Resource Center (2023) and the atlases of Q. Wang (2016) and Q. Zhou & Y. Gan (2016).

Standard protocols were used to measure the traits (Pérez-Harguindeguy et al. 2013). For measurements, we sampled 10–15 well-developed leaves without signs of damage, pathogenic infection, or senescence. For several species with small leaves (e.g. *Ceratium falcatum* Bunge), we collected 3–5 leaves from one plant per sample, but in all cases, the total number of samples was 10–15. Cutoff leaves were placed in plastic boxes with water and were kept in the fridge for 5–10 hours to get water-saturated status.

To measure the LFM and LDM, we used FA-N Series Electronic Analytical Balance. We used a scanner (CanoScan LiDE 120) to scan the leaf samples for LA measurements at 300 dpi for medium size leaves and 600 dpi for small leaves. Leaf samples were dried by DHG Series Heating and Drying Oven (DHG-9240B).

Specific leaf area (SLA) was defined as the ratio between LA and LDM. Leaf dry matter content (LDMC) was calculated as the ratio between LDM and LFM.

To calculate the CSR strategy affinities of collected species, based on LA, LDM and LFM, we used the global plant strategy calculator tool StrateFy (Pierce et al. 2017). Thus, strategies were defined for 99 species.

**Community-weighted means**

We calculated community biomass weighted means (CWM) of LFM, LDM, LDMC, LA, SLA, and CSR affinities in each plot by Eq.1 (Lavorel et al. 2008):

$$X_i = \sum B_i \cdot X_i$$

where $X_i$ is community-weighted means of traits or strategies; $X_i$ is the value of the trait of $i$-species; $n$ means the number of species on the plot and $B_i$ is the relative abundance (biomass proportion) of $i$-species.

The data on the biomass of each species was obtained from square plots of 0.0625 m² (25×25 cm), located in typical sites of the communities, in September 2011 and late June-July 2012 (Elumeeva et al. 2015b). There were 53 sampling plots in KAM (the total area was 3.3 m²) and 20 plots in AF (total area 1.25 m²). The distance between plots was at least 1 m. Aboveground biomass was clipped at the surface level, and vascular plants were sorted by species. Air-dried samples were dried in the oven before weighing.

In total, 76 species with all measured traits and defined CSR types occurred in biomass sampling plots, accounted in an average 95 % (85–100 %) of aboveground biomass. The number of species with available LA, LDM, and SLA data was 91, and they contributed 94–100 % to the total biomass.

Besides the community-weighted means, we calculated trait means per plot, assuming that all species occurring in the plot have equal biomass. Function *functcomp* in the *FD* package in R (Laliberté et al. 2014) was used.

**Functional diversity indices**

We calculated three indices of functional diversity for single traits: functional richness (FRic), functional evenness (FEve) and functional divergence (FDiv). Function *divFD* in the *FD* package in R (Laliberté et al. 2014) was used for FRic and FEve, and FDiv was calculated by the exact formula of abundance-weighted functional variance by Eq. 2 (Mason et al. 2005):

$$\text{FDiv} = \frac{2}{\pi} \arctan \left(5 \times \sum \left(\ln X_i - \ln \bar{X} \right)^2 \times B_i\right)$$

where $n$ means the number of species, $X_i$ – trait or strategy value of the $i$-species, $\ln X_i$ is the mean logarithmic trait values, and $B_i$ is the relative abundance (biomass proportion) of $i$-species.
Data analysis

Because the numbers of sampling plots were different, to compare the leaf traits between the AF and KAM we used an approximative general independence test in the coin package in R (Hothorn et al. 2008). This test was applied for mean trait values, calculated per plot, and for community-weighted means.

To test whether community-weighted means of traits differ from means, we used paired Student’s t-test if both the variables in the community were normally distributed following the results of the Shapiro test, and the Wilcoxon signed rank exact test if the data did not comply with normality assumptions. These tests were run separately for AF and KAM.

RESULTS

Leaf functional traits

Most of the studied plants in the Eastern Qinghai-Tibetan Mountains have low leaf fresh mass (<100 mg), small leaf area (<5 cm²) and low leaf dry matter content (<30 %; Table 1). The specific leaf area of the most of leaves is less than 300 cm²/g. The smallest leaves, less than 0.1 cm² were typical for annual gentianas (*Gentiana crassuloidea*) Bureau et Franch., *G. squarrosa* Ledeb., *Androsace brachystegia* Hand.-Mazz., and the largest leaves, more than 20 cm², were in *Angelica sinesis* R.H. Shan & C.Q. Yuan and *Pleuroserum wilsonii* H. Boissieu. The lowest LDMC was in the succulent *Sedum rosei* Hamet (4.8%). Plants with the highest LDMC were shrubs (*Potentilla fruticosa* Linn., *Salix capularis* Rehd., *Spiraea alpina* Pall.), *Festuca arvenis* Linn., *Carex atrypusia* Schkuhr. and some other species of Cyperaceae. SLA varied from 54 cm²/g in *Juncus sikkimensis* Hook. f. and 99 cm²/g in *Rhoodendron thymifolium* Maxim. to 755 cm²/g in *Veronica vanellus*- *V. sandellinoides* Maxim. Other species with a very high SLA (>400 cm²/g) were *Gentiana straitae* Maxim., *Cerastium falcatum* Bunge, *Sedum rosei* Hamet, *Lomatogoniopsis alpina* T.N. Ho et S.W. Liu, *Draba eriophoda* Turcz.

The plots of both the communities had similar values of mean LDM, while LA, LFM and SLA were higher and LDMC was lower in KAM (Table 2, Fig. 2). LFM was significantly higher in the KAM, and LDMC was higher in the AF, but other CWMs did not differ significantly between communities (Table 2, Fig. 2).

Community-weighted means of all the traits, except LDM, differed significantly from the mean trait values per plot in KAM (Table 2). In the contrary, in AF community, only LDMC was significantly higher than the trait mean, and LFM was lower than the trait mean.

**CSR strategies of alpine plants**

The prevailing strategies of plants were ruderal R-strategy (ca. 38 %; Table 3) and combined ruderal and stress-tolerant SR-strategy (28 %). The most of plants (97 out 99 species) had expressed R-strategy scores, while S and competitive C-strategies were assigned to 66 and 55 species, respectively. There were no species with C-strategy. The typical R-selected (ruderal) species are *Galamus elegans* Wall. ex Roxb. (C:S:R= 0:0:100 %) and *Koenigia islandica* Linn. (C:S:R= 0:0:100 %), which have a very small area of leaves.

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Table 1. The descriptive statistics of leaf functional traits in alpine plants. n – number of species accessed, LFM – leaf fresh mass; LDM – leaf dry mass; LA – leaf area; LDMC – leaf dry matter content; SLA – specific leaf area.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Value</th>
<th>Kobresia alpine meadow (Mean ± SE, n=33)</th>
<th>Alpine fen (Mean ± SE, n=20)</th>
<th>Between-community differences, p</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>(Mean ± SE, n=33)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Trait mean</td>
<td>2.77±0.06</td>
<td>2.27±0.12</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>2.55±0.11</td>
<td>2.27±0.12</td>
<td>0.169 n.s.</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>0.956 n.s.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Trait mean</td>
<td>63.0±1.8</td>
<td>51.5±2.5</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>54.5±3.4</td>
<td>41.9±2.9</td>
<td>0.034*</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>&lt;0.001***</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Trait mean</td>
<td>13.6±0.3</td>
<td>12.5±0.7</td>
<td>0.131 n.s.</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>13.2±0.6</td>
<td>12.9±0.9</td>
<td>0.764 n.s.</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>0.622 n.s.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Trait mean</td>
<td>25.0±0.1</td>
<td>26.2±0.4</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>29.2±0.4</td>
<td>31.5±0.5</td>
<td>0.003**</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>&lt;0.001***</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Trait mean</td>
<td>218±1</td>
<td>194±2</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>196±2</td>
<td>191±5</td>
<td>0.275 n.s.</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>&lt;0.001***</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Mean S-strategies scores of species, occurring in biomass sampling plots were higher in AF and R-scores were higher in KAM, while C-scores did not differ between communities (Table 4). The community-weighted means of all C and R affinities were higher in KAM, while S-strategy CWMs were higher in AF. Therefore, in both the communities CWM values of S-strategy exceeded mean values, and CWMs of C- and R-strategies were lower than trait values, so from the available local species pool in both the communities’ dominant species with the better-expressed stress-tolerance (Fig. 3).

**DISCUSSION**

**Plant functional traits**

The plants in the study area have general characteristics, typical for alpine plants at the global scale. Almost all studied plant species were shrubs or herbs with generally light and small leaves. Among other mountain systems, the mean values of leaf traits were close to those in the Northwestern Caucasus (Onipchenko et al. 2022) and the Alps (Körner 2003).

**The CSR-types of alpine plants**

The total vascular plant species pool of alpine communities at Mt. Kaka is not still estimated, though earlier we obtained 108 species from the total 375 m² recorded plots in KAM and 60 species from the total 100 m² in AF (Elumeeva et al. 2015a), with the maximum species richness of 82 per 100 m² in KAM (Onipchenko et al. 2014). Indeed, there are many species not covered by sampling plots, therefore our data by leaf traits include the most known diversity and the most common species at the altitude range of 3930–3960 m a.s.l.

The KAM are richer than AF both floristically (Onipchenko et al. 2014) and functionally (Table 5), and this reflects specific ecological conditions of alpine fens with a high water saturation of soils and lower aeration. For example, alpine fens of the Northwestern Caucasus also have a rather low species richness (Onipchenko & Semenova 1995). Lower functional richness indicates that there are fewer species can take advantage from this environmental condition (Tilman et al. 1997). The comparison of the local functional richness with other studies seems not to be relevant because the studies from different regions are based on different species pools.
Both communities have rather low functional evenness, which suggests the existence of one or more groups of species with similar trait values, which dominate the community. The lowest functional divergence is observed for SLA, thus the dominance is shifted to a single group of species. In contrary, the size traits have high FDiv indices and there can be two or more functionally distinct groups.

The differences between mean trait values reflect the differences in ecological conditions between sites, which influence the selection of species that are able to survive in a given environment, while differences between CWMs reflect the dominance of species with certain trait values.

In the case of the studied KAM and AF, our data suggests, that there can be the same factor, which acts on selection by the SLA and leads to the similar CWMs of this trait (196 and 191 cm$^2$/g, respectively). This selection to dominance can be caused by intensive grazing, which both affects meadows and fens. These CWMs of SLA are close to the value, reported for alpine snowbeds in the Caucasus (Elumeeva et al. 2015a, Onipchenko et al. 2022), which structure is supposed to be due to overgrazing and kept despite long-term reservation. In contrary, Li et al. (2017) found that SLA$_{CWM}$ of alpine swamps in the eastern part of the Tibetan plateau north of our site, where standing water is present during the growing season, is ca. 120–130 cm$^2$/g and lower than that of swamp meadows and typical meadows, where its values are also close to ours (ca. 180–190 cm$^2$/g). They also reported lower LDMC$_{CWM}$ of the alpine swamp (25-30%), which kept the range of our values for both communities.

In contrast to “economics” traits, the size traits show high functional divergence, which suggests the dominance of at least two groups with small and relatively large leaves.

The plants of alpine communities of the studied site suffer from two factors, which in opposite ways influence SLA. Intensive grazing promotes species with high SLA, which is usually associated with a high relative growth rate (Wright & Westoby 2001) and is important for fast regrowth. On the other hand, grazing reduces plant height and the growth of shrubs, which improves light availability and increases ultraviolet stress in high altitudes, promoting the species with lower SLA (Li et al. 2010). The higher mean values of SLA show that many species of KAM are comfortable to withstand light shading of alpine shrubs such as Potentilla fruticosa or Spiraea alpina, but grazing disturbance shifts down CWMs of SLA with the corresponding increase of affinities to stress-tolerance, while, for example, SLA$_{CWM}$ decreases under Potentilla fruticosa shrubs (Wang et al. 2021).

Table 4. Community weighted means and strategies means of species in *Kobresia* alpine meadows and alpine fens. SE – standard error. C – C-selected proportion of species; S – S-selected proportion of species; R – R-selected proportion of species. Different letters indicate significant difference between CWMs and strategies means in the same community at p < 0.05, * – p < 0.05; ** – p < 0.01, *** – p<0.001, n.s. – not significant. Approximative general independence test was used to check between-community differences, and paired Student’s t-test or Wilcoxon signed rank exact test were used to check within-community differences.

<table>
<thead>
<tr>
<th>Strategy, affinity, %</th>
<th>Value</th>
<th><em>Kobresia</em> alpine meadow (Mean ± SE, n=53)</th>
<th><em>Alpine fen</em> (Mean ± SE, n=20)</th>
<th>Between-community differences, p</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>13.0±0.2</td>
<td>12.4±0.4</td>
<td>0.161 n.s.</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>11.7±0.6</td>
<td>9.6±0.3</td>
<td>0.032*</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>0.003**</td>
<td>&lt;0.001***</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>41.0±0.4</td>
<td>46.9±1.4</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>52.6±1.3</td>
<td>60.7±1.9</td>
<td>0.001**</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>&lt;0.001***</td>
<td>&lt;0.001***</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>45.9±0.4</td>
<td>40.6±1.2</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td></td>
<td>CWM</td>
<td>35.6±0.8</td>
<td>29.7±1.8</td>
<td>&lt;0.001***</td>
</tr>
<tr>
<td></td>
<td>Within-community differences, p</td>
<td>&lt;0.001***</td>
<td>&lt;0.001***</td>
<td></td>
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</tbody>
</table>
CONCLUSION

The leaf traits of plants in the communities of the eastern Qinghai-Tibet plateau are typical for the alpine belt and are affected by local environmental factors, such as grazing by yaks and excess water availability. KAMs are more functionally rich than alpine fens, but both have low functional evenness. The abiotic environment causes the prevalence of stress-tolerant strategy, while grazing enhances the role of R-selection and dominance of species with a narrow value range of some traits. The most indicative trait was SLA, where means in AF were significantly lower than in KAM, but SLAvm were similar.

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