

## Dry grasslands of NW Bulgarian mountains: first insights into diversity, ecology and syntaxonomy

### Trockenrasen in den Gebirgen Nordwest-Bulgariens: erste Einblicke in Diversität, Ökologie und Syntaxonomie

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#### Abstract

We present the data of the 3<sup>rd</sup> research expedition of the European Dry Grasslands Group (EDGG), which was conducted in 2011 in two contrasting areas in NW Bulgarian mountains. The aim was to collect plot data for comparing Bulgarian dry grasslands with those of other parts of Europe in terms of syntaxonomy and biodiversity. We sampled 15 nested-plot series (0.0001–100 m<sup>2</sup>) and 68 normal plots (10 m<sup>2</sup>) covering the full variety of dry grassland types occurring in the Vratsa area (Balkan Mts.) and the Koprivshtitsa area (Sredna Gora Mt.). In the plots all vascular plants, terricolous non-vascular plants and a set of soil and other environmental parameters were determined. By applying modified TWINSPAN, we distinguished 10 floristically well characterised vegetation types at the association level. After comparison with the regional and European literature, we propose to place them within three classes and five orders: *Festuco-Brometea* with the orders *Stipo pulcherrimae-Festucetalia pallentis* (xerophilous dry grasslands of base-rich rocks; alliance *Saturejion montanae*), *Brachypodietalia pinnati* (meso-xeric, basiphilous grasslands; alliances *Cirsio-Brachypodion pinnati* and *Chyrsopogono grylli-Danthonion calyciniae*), *Calluno-Ulicetea* with the order *Nardetalia stricae* (lowland to montane *Nardus* swards; alliance *Violion caninae*), and *Koelerio-Corynephoretea* with the orders *Sedo-Scleranthetalia* (open communities of skeleton-rich, acidic soils; alliance unclear) and *Trifolio arvensis-Festucetalia ovinae* (closed, meso-xeric, acidophilous grasslands; alliance *Armerio rumelicae-Potentillion*). The *Violion caninae* with the association *Festuco rubrae-Genistelletum sagittalis* is reported from Bulgaria for the first time, while the two occurring *Koelerio-Corynephoretea* communities are described as new associations (*Cetrario aculeatae-Plantaginetum radicatae*, *Plantagini radicatae-Agrostietum capillaris*). According to DCA the main floristic gradient was largely determined by soil conditions, differentiating the *Festuco-Brometea* communities on soils with high pH and high humus content from the *Koelerio-Corynephoretea* communities on acidic, humus-poor soils, while the

*Calluno-Ulicetea* stands are the connecting link. At 10 m<sup>2</sup> *Festuco-Brometea* and *Calluno-Ulicetea* stands were richer in species across all investigated taxa and in vascular plants than *Koelerio-Corynephoretea* stands; the latter were richest in lichen species, while bryophyte richness did not differ significantly among syntaxa. Among the Bulgarian classes, the species-area relationships tended to be steepest in the *Festuco-Brometea* (i.e. highest beta diversity), but both alpha and beta diversity clearly fell behind the *Festuco-Brometea* communities in the Transylvanian Plateau, Romania, located less than 500 km north of the study region. Overall, our study contributes to a more adequate placement of the Bulgarian dry grasslands in the European syntaxonomic system and provides valuable data for large-scale analyses of biodiversity patterns.

**Keywords:** bryophyte, *Calluno-Ulicetea*, conservation, European Dry Grassland Group (EDGG), *Festuco-Brometea*, high nature value grassland, *Koelerio-Corynephoretea*, lichen, nomenclatural revision, species richness, vascular plant, vegetation classification

**Erweiterte deutsche Zusammenfassung am Ende des Textes**

## 1. Introduction

The Balkan Peninsula, as south-east corner of the European continent, is well known for its rich flora and well-preserved vegetation. GRIFFITHS et al. (2004) describe this area as glacial refuge for animal and plant species and as a crossroad for faunal and floral exchange between Central Europe and Asia Minor. Overall, biodiversity in the region is considered to be among the highest in Europe (POLUNIN 1980). According to an older count, the flora of the peninsula comprises 6,530 native vascular plant species, which would be about 62% of the European flora (POLUNIN 1980), but new records and taxonomic revisions during the last three decades certainly have increased the absolute number. The extremely high level of endemism in the Balkans compared to other parts of Europe is probably the most outstanding feature across numerous taxa (GRIFFITHS et al. 2004). For example, among the vascular plants, nearly one quarter is endemic to the peninsula (POLUNIN 1980). Moreover, in Bulgaria, which covers the north-eastern part of the Balkan Peninsula, semi-natural grasslands of high nature value (HNV grasslands) persisted until today to an extent that exceeds most other European countries (MESHINEV et al. 2005, 2009, BEAUFOY et al. 2012, STEFANOVA & KAZAKOVA 2012). However, as in many places of Europe and particularly the post-communist countries, there is a rapid decline in area and quality of these valuable habitats (e.g. VASSILEV et al. 2011).

While the first relevés following the Braun-Blanquet approach (BRAUN-BLANQUET 1965) in Bulgaria were collected in the first half of the 20th century (e.g. RUSKOV 1935, HRISTOV 1944, STEFANOV & STOYANOV 1949), classification was not a topic at that time. First attempts to classify vegetation were made in the 1950s, using the dominance approach (see review by APOSTOLOVA et al. 2012). There were also occasional studies by foreign geobotanists (e.g. HORVAT et al. 1937, SOÓ 1955, MUCINA & KOLBEK 1989), who applied the Braun-Blanquet approach, but their work received little popularity within Bulgaria at that time. In the synthesis of the Balkan vegetation by HORVAT et al. (1974), Bulgaria was represented by a single relevé. Similarly, in the overview of European high-rank syntaxa (RODWELL et al. 2002), the country was largely neglected because of the lack of comprehensive data. Until 1990 Bulgaria remained scientifically isolated as the only country in the region that exclusively applied the dominance approach. However, after the fall of the Iron Curtain and stimulated by the requirement of applying European standards in habitat classification

and site protection (e.g. NATURA 2000 network), the number of vegetation classification studies following the Braun-Blanquet approach rapidly increased (e.g. MUCINA et al. 1990, ROUSSAKOVA 2000, HÁJEK et al. 2008, TZONEV et al. 2009, ELIÁŠ et al. 2013).

Due to the abundance of rare and endemic species (VASSILEV 2012), dry grasslands have always been in the focus of the scientific interest of Bulgarian botanists. Some researchers determined limestone terrains in the continental part of the Balkan Peninsula as “Mediterranean oases”, hosting a large number of Mediterranean species (STOYANOV 1922, 1935, YORDANOV 1924, URUMOV 1935). Later VELCHEV (1962, 1971) classified communities using dominant species and characterised them according to their economic significance, whereas recently numerical classification methods have been applied and more attention has been paid to the conservation value of these grasslands (TZONEV et al. 2006, PEDASHENKO et al. 2009, 2010, TODOROVA & TZONEV 2010, VASSILEV 2012, VASSILEV et al. 2012a). However, most of the recent studies deal with few vegetation types in small geographic areas, often describing single new associations. Concerning the grassland vegetation, the first attempt to place the Bulgarian syntaxa in a European system (TZONEV et al. 2009) remained unsatisfactory in many respects mainly due to a lack of available large-scale syntheses. Already in 1991 ROYER had produced a monumental synthesis of the class *Festuco-Brometea* (basiphilous dry grasslands) in its entire range from Western Europe to Central Asia, but Bulgaria was hardly mentioned in his work, and the synthesis is also outdated in various respects. While the Bulgarian Vegetation Database (APOSTOLOVA et al. 2012) now contains a significant amount of relevés from *Festuco-Brometea* and other dry grassland classes, in the huge majority of these, bryophytes and lichens have not been treated, despite the knowledge that they generally have diagnostic value and ecological significance in dry grassland communities (BERG & DENGLER 2005, DENGLER 2005).

Dry grasslands are particularly suitable study objects for understanding biodiversity patterns as they have a very wide ecological and geographical amplitude, i.e., they are present in nearly all European regions except the Far North, distributed from the sea coast to the alpine zone and from extremely acidic to very base-rich soil conditions (DENGLER 2005). In addition to a sometimes extreme richness in vascular plants (WILSON et al. 2012), they also host significant numbers of bryophyte and lichen species (DENGLER 2005, BOCH & DENGLER 2006, LÖBEL & DENGLER 2008, JESCHKE 2012) and thus represent an ideal system for testing whether these three taxonomic groups with strongly divergent ecological properties also show different diversity-environment relationships (e.g. LÖBEL et al. 2006). Further, diversity patterns and their relationship to environmental drivers might also strongly depend on spatial scale, and species-area relationships (SARs) might reveal interesting insights into community organisation (e.g. DUPRÉ & DIEKMANN 2001, DENGLER & BOCH 2008).

In order to collect “high-quality” data on plant species composition and diversity of dry grasslands and related communities in different regions of the Palaearctic ecozone, the *European Dry Grassland Group* (EDGG; [www.edgg.org](http://www.edgg.org)) organises annual research expeditions since 2009 (VRAHNAKIS et al. 2013). With “high-quality” we refer to (1) full recording of all terricolous taxa including bryophytes and lichens, (2) uniform plot size, (3) accurate GPS coordinates (that allow repeated sampling in the future) and (4) a set of environmental variables measured in the plot. The EDGG expeditions aim at better understanding of scale-dependent plant diversity patterns and contributing to consistent large-scale classification of dry grassland syntaxa. The 1<sup>st</sup> EDGG Research Expedition was carried out in Transylvania

(Romania) in 2009, which resulted in a new classification scheme based on modern phytosociological methods (DENGLER et al. 2012a) and the documentation of the richest ever recorded plant community at the scale of 0.1 m<sup>2</sup> and 10 m<sup>2</sup> (WILSON et al. 2012).

The still poor data on the vegetation of Bulgarian dry grasslands and the complete lack of comparable biodiversity data from the region were the main motivation for the EDGG to direct its third international research expedition to this country in 2011. The expedition was carried out in two previously unstudied areas, the western part of the Balkan Mts. and the valley between the Balkan Mts. and Sredna Gora Mt., in order to compare sites with different geology, altitude and communities. The main aims were: (1) Delimitation of the major plant community types of dry grasslands and identification of their diagnostic species with modern statistical methods. (2) Placement of these community types in a supraregional classification scheme. (3) Insights into the role of environmental factors in shaping the floristic composition of the studied communities. (4) Documentation and comprehension of the plant diversity patterns of vascular plants, bryophytes and lichens in these vegetation types at various spatial scales.

## 2. Study areas

### 2.1 Location and delimitation

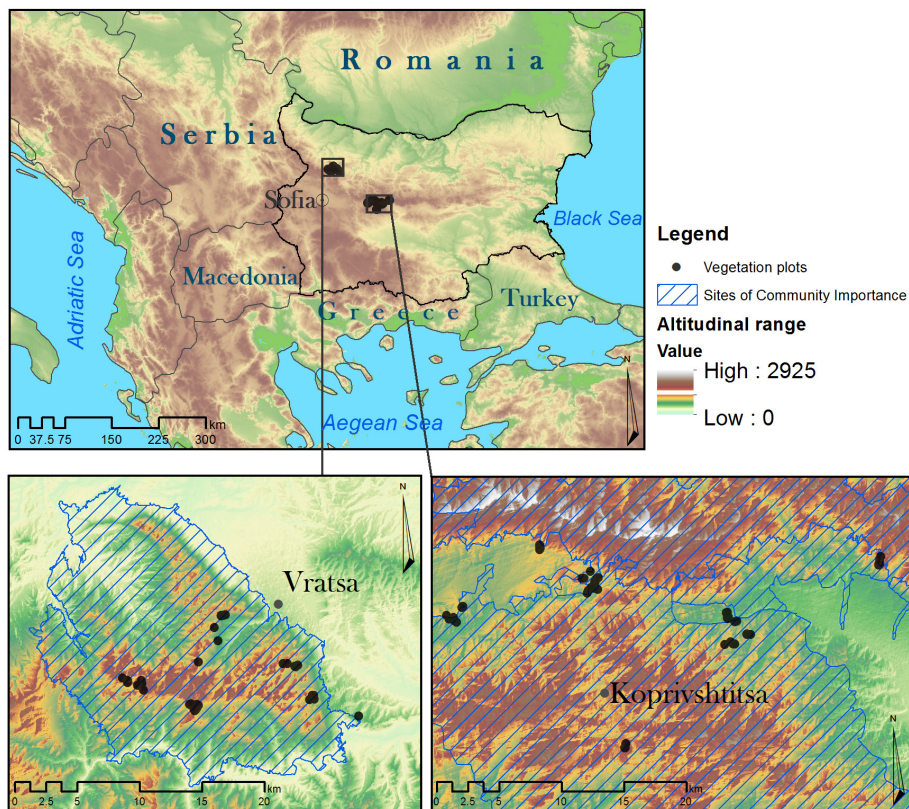
The study was conducted in two contrasting areas of northwestern Bulgaria belonging to the districts of Sofia, Plovdiv and Vratsa (Fig. 1). The first study area (Vratsa) is located in the western part of the Balkan Range (Stara planina), while the second (Koprivshitsa) comprises the valley between the Balkan Range and Sredna gora Mt., including foothills of both ridges. The studied grasslands in both areas belong to the NATURA 2000 network, specifically to four Sites of Community Interest (SCIs): Vrachanski Balkan (BG0000166), Sredna gora (BG0001389), Tsentralen Balkan (BG0000494) and Tsentralen Balkan buffer (BG0001493). In addition, the Vratsa area is located in the Vrachanski Balkan Nature Park and its reserve Vrachanski karst (IUCN category I). Parts of the Koprivshitsa area belong to the second biggest national park in Bulgaria, Tsentralen Balkan (IUCN category II), with one of the largest natural pristine European beech (*Fagus sylvatica*) forests.

### 2.2 Physical geography

The two areas were chosen to cover territories with contrasting abiotic conditions such as soil, altitude and climate. In essence, Vratsa is a high-altitude limestone area, while Koprivshitsa is located at lower altitudes on acidic bedrock (Table 1).

The climate of both areas is temperate to continental and characterised by warm summers and cold winters, resulting in a wide annual temperature amplitude. The precipitation maximum occurs in May and June and its minimum in January and February (VELEV 2002). Two main geomorphological units can be distinguished in both areas: the Balkan Mountain System, which is the most distinct mountain range in Bulgaria, crossing the whole country from west to east, and the Transitional Zone with alternating lower mountain ridges and depressions (STEFANOV 2002). Mountain karst (ANGELOVA 2003) is the most common type of bedrock in Vratsa area, which is very specific and is set in marble, marbled limestones, dolomites and limestones of different ages, predominantly Mesozoic (BENDEREV et al. 2005, MIHAYLOVA et al. 2009). Koprivshitsa area is characterised by gneiss and granite. The soils





**Fig. 1.** Location of the two study regions within Bulgaria and distribution of the analysed 98 vegetation plots within these (note that due to geographic proximity not all relevés are visible as separate symbols).

**Abb. 1.** Lage der zwei Untersuchungsgebiete innerhalb Bulgariens und räumliche Verteilung der 98 Aufnahmeflächen innerhalb derselben (wegen der geografischen Nähe sind nicht alle Aufnahmen als separate Punkte erkennbar).

of Vratsa Mt. are a mosaic of Lithic Leptosols, the most shallow soil type occurring in Bulgaria, and Rendzic Leptosols, comprising more than 40% carbonates, while the area of Koprivshitsa is covered predominantly by Dystric Cambisols (NINOV 2002).

### 2.3 Landscape and vegetation

Chorologically the study areas are considered as part of the Balkan province of the sub-Mediterranean subregion of the Macaronesian-Mediterranean floristic region (MEUSEL & JÄGER 1992). The potential natural vegetation in the Vratsa area is mostly comprised of East Moesian sessile oak-hornbeam forests (*Carpinus betulus*, *Quercus dalechampii*) with *Corydalis solida* subsp. *slivenensis* (Mapping unit F67) and Moesian, partly thermophilous beech forests (*Fagus sylvatica* subsp. *moesiaca*) with *Corylus colurna* and *Hypericum umbellatum*, partly with *Acer hyrcanum* subsp. *intermedium* (Mapping unit F152) (BOHN et al. 2004). In Koprivshitsa two additional categories are present: East Moesian Balkan oak bitter

**Table 1.** Overview of the sampling intensity and the main environmental features of the study areas.

**Tabelle 1.** Übersicht der Aufnahmezahlen und wichtigsten Umweltbedingungen der beiden Untersuchungsgebiete.

Parameter	Vratsa	Koprivshtitsa
Number of 10-m <sup>2</sup> normal plots	40	58
Number of 100-m <sup>2</sup> biodiversity plots	6	9
Study area (km <sup>2</sup> )	135	310
Altitude range (m a.s.l.)	970–1400	630–1200
Mean annual temperature (°C)	6.0	7.8
Mean temperature of warmest quarter (°C)	15.0	17.0
Mean Temperature of coldest quarter (°C)	-3.0	-1.9
Annual precipitation (mm)	722	670
Dominant soil types	Lithic and Rendzic Leptosols	Dystric Cambisols
Bedrock	Carbonate	Silicate

oak forests (*Quercus cerris*, *Q. frainetto*) with *Carpinus orientalis* and *Fraxinus ornus* (G22) and Thracian oriental hornbeam-downy oak forests (*Quercus pubescens*, *Carpinus orientalis*) with *Acer tataricum*, partly with *Juniperus oxycedrus* and *Pistacia terebinthus* (G59; BOHN et al. 2004).

The two studied areas have traditionally been grazed. According to locals hundreds of flocks have grazed the pastures in the mountains in the past. The closure of the border with Greece in the middle of the 19<sup>th</sup> century led to the disappearance of specific husbandry practices of the ethnic group of Karakachans and a substantial reduction of livestock within the area. In consequence of the socio-economic changes following 1989, the use of mountain pastures decreased even more due to further reduction of cattle and sheep numbers (YANKOV 2001, MESHINEV & APOSTOLOVA 2007). Partially the mountain grasslands are also used as hay meadows (Fig. 2).



**Fig. 2.** Extensively used hay meadows at Milanovo, near Vratsa (Photo: M. Janišová).

**Abb. 2.** Extensive Wiesenwirtschaft in Milanovo oberhalb Vratsa (Foto: M. Janišová).

### 3. Methods

#### 3.1 Vegetation sampling

From 14 to 24 August 2011, the field sampling was carried out by the nine participants of the 3<sup>rd</sup> EDGG Research Expedition. We tried to cover the whole range of different dry grassland types (*Festuco-Brometea*, *Koelerio-Corynephoretea*, dry *Nardetalia strictae*) present in the two areas.

We applied two complementary sampling schemes: (i) nested-plot series ( $n = 15$ ) with plots of 0.0001, 0.001, 0.01, 1, 10 and 100 m<sup>2</sup>, where all plot sizes except the largest one were replicated in two opposite corners (DENGLER 2009b), and (ii) additional 10-m<sup>2</sup> plots (“normal plots”;  $n = 68$ ). In both cases all plots were square-shaped and placed in stands that were visually homogeneous in terms of topography, vegetation structure and floristic composition of all terricolous taxa estimated in % (for more details, see DENGLER et al. 2012a).

The taxonomy of vascular plants follows *Flora Europaea* (TUTIN et al. 1968–1993) or *Atlas Florae Europaeae* (JALAS & SUOMINEN 1994, JALAS et al. 1996, 1999, KURTTO et al. 2004–2010), depending on which of the treatments in the two series was the more recent. For mosses we adopted HILL et al. (2006), for liverworts GROLLE & LONG (2000) and for lichens SANTESSON et al. (2004). We accepted a few additions and deviations from these checklists such as definition of additional aggregates (see Tables 2 and 3). Prior to the numerical classification and ordination, we “standardised” the taxonomic information and joined layers in our relevé data following the principles outlined in DENGLER et al. (2012a), based on the recommendations of JANSEN & DENGLER (2010). The full original information is available in Tables 5 and 6 in the supplement, where it is also transparent which entries had been joined or removed.

**Table 2.** Supraspecific taxa used in the paper in addition to those defined in the given checklists.

**Tabelle 2.** Supraspezifische Taxa, die in dieser Publikation zusätzlich zu den in den angegebenen Referenzlisten definierten Aggregaten genutzt werden.

Group	Taxon	Reference	Included species from Bulgaria
Vascular plants	<i>Centaurea jacea</i> agg.	TUTIN et al. (1976: as sect. <i>Jacea</i> + sect. <i>Fimbriatae</i> )	<i>C. jacea</i> , <i>C. pannonica</i> , <i>C. rocheliana</i> , <i>C. subjacea</i>
	<i>Centaurea phrygia</i> agg.	EHRENDORFER (1973)	<i>C. phrygia</i> , <i>C. stenolepis</i>
	<i>Festuca</i> ser.	PAWLUS (1985), DENGLER (1996)	<i>F. callieri</i> , <i>F. dalmatica</i> , <i>F. illyrica</i> , <i>F. pseudovina</i> , <i>F. rupicola</i> , <i>F. stojanovii</i>
	<i>Valesiaca</i>		
	<i>Genista tinctoria</i> agg.	TUTIN et al. (1968: as <i>Genista</i> sect. <i>Genista</i> )	<i>G. depressa</i> (see Table 3), <i>G. januensis</i> , <i>G. lydia</i> , <i>G. tinctoria</i>
	<i>Quercus petraea</i> agg.	<i>ad hoc</i> definition	<i>Q. dalechampii</i> , <i>Q. petraea</i> , <i>Q. polycarpa</i>
Bryophytes	<i>Racomitrium canescens</i> agg.	NYHOLM (1998: as <i>R. sect. Canescentia</i> )	<i>R. canescens</i> , <i>R. elongatum</i> , <i>R. ericoides</i>
	<i>Schistidium apocarpum</i> agg.	NYHOLM (1998: as <i>Apocarpum</i> group)	33 species, of which many might also occur in Bulgaria (see NYHOLM 1998)
	<i>Syntrichia ruralis</i> agg.	<i>ad hoc</i> definition	<i>S. calcicola</i> , <i>S. montana</i> , <i>S. norvegica</i> , <i>S. ruralis</i>
Lichens	<i>Cladonia cervicornis</i> agg.	WIRTH (1995: as <i>C. cervicornis</i> )	<i>C. cervicornis</i> , <i>C. pulvinata</i> , <i>C. verticillata</i>
	<i>Cladonia pyxidata</i> agg.	WIRTH (1995: as <i>C. pyxidata</i> )	<i>C. chlorophaea</i> , <i>C. cryptochlorophaea</i> , <i>C. grayi</i> , <i>C. merochlorophaea</i> , <i>C. monomorpha</i> , <i>C. novochlorophaea</i> , <i>C. pocillum</i> , <i>C. pyxidata</i>

**Table 3.** Additional taxa of vascular plants used in this publication that are not accepted in the given checklists.

**Tabelle 3.** Zusätzliche Taxa, die in dieser Publikation Verwendung finden, aber in den angeführten taxonomischen Referenzlisten nicht akzeptiert sind.

Taxon	Author citation	Reference	Treatment in checklist
<i>Dactylis glomerata</i> subsp. <i>glomerata</i>	–	TUTIN et al. (1980) [ <i>vide taxon</i> ]	no infraspecific taxa formally accepted
<i>Festuca stojanovii</i>	(Acht.) Foggi & Petrova	FOGGI & MÜLLER (2009)	not mentioned, likely included in <i>F. dalmatica</i>
<i>Genista depressa</i>	Bieb.	JORDANOV (1976)	included in <i>G. tinctoria</i>
<i>Lilium jankae</i>	A. Kerner	TUTIN et al. (1980) [ <i>vide taxon</i> ]	listed under <i>L. carnolicum</i> without definitive decision about synonymy or validity as separate species

All relevés and nested plots used in this article have been stored in and are available from the *Database Species-Area Relationships in Palaearctic Grasslands* (DENGLER et al. 2012b; GIVD ID EU-00-003) and the normal plots additionally in the *Bulgarian Vegetation Database* (APOSTOLOVA et al. 2012; EU-BG-001) and the *Balkan Dry Grassland Database* (VASSILEV et al. 2012b; EU-00-013). Further details on these databases are available from the *Global Index of Vegetation-Plot Databases* (GIVD; <http://www.givd.info>; see DENGLER et al. 2011), where they are registered under the given IDs.

### 3.2 Environmental parameters

In all 10-m<sup>2</sup> plots ( $n = 98$ , normal plots and plots from nested-plot series), we estimated the cover of the shrub, herb and cryptogam layer as well as that of litter, superficially visible stones and rocks and open soil. We further recorded coordinates and altitude with a GPS as well as aspect and slope with an inclinometer. The latter two parameters were used to calculate the heat load index according to OLSSON et al. (2009), which is a relative measure that takes the highest positive values on steep southwest-facing slopes, zero in level areas and negative values on northeast-facing slopes. Microrelief was measured as maximum vertical deviation from an imaginary plane through the plot. We classified present land use roughly into the categories mown, grazed and not used, based on our knowledge of the sites and visible signs in the plots.

Furthermore, we took a mixed sample of the uppermost 10 cm of soil. Samples were air-dried at room temperature and sieved to < 2 mm. We measured pH and conductivity (determined in a solution of 10 g soil in 25 ml distilled water with a standard glass electrode) and soil organic content (by loss on ignition at 550 °C for 16 hours) (see SCHLICHTING et al. 1995). In addition, we determined soil texture with the HORIBA LA-950V2 Particle Size Distribution Analyzer (HORIBA INSTRUMENTS 2010).

### 3.3 Vegetation classification

Vegetation classification was performed on basis of the 98 relevés of the 10-m<sup>2</sup> plots. We applied the modified version of TWINSpan (Two-Way Indicator Species Analysis; HILL 1979) proposed by ROLEČEK et al. (2009) and implemented in JUICE (TICHÝ 2002). We used four pseudospecies cut levels (0%, 2%, 5% and 10%), a minimum group size of six and total inertia as heterogeneity measure, while we tried cluster numbers up to 15.

As the TWINSpan analysis showed that the three-cluster resolution corresponded closely to phytosociological classes known from the literature (“class clusters”) and the ten-cluster resolution was the lowest resolution with floristically well-defined units (“association-level clusters”), we used these two TWINSpan resolutions as starting point for further refinement. We calculated phi coefficients of association (CHYTRÝ et al. 2002) for these units using standardisation to equal cluster size at the 10-cluster level (DENGLER et al. 2012a, LUTHER-MOSEBACH et al. 2012) to avoid undue effects of widely

differing numbers of included relevés (see TICHÝ & CHYTRÝ 2006). Normally it is not sensible to translate the outcome of a numerical analysis directly into a syntaxonomic classification, in particular when the major syntaxonomic groups are represented by very unequal numbers of relevés (e.g. LUTHER-MOSEBACH et al. 2012; WILLNER et al. 2013). Therefore, in the next step, we tried to interpret the lists of diagnostic species in ecological terms and to match the units of the original TWINSpan classification with existing large-scale syntaxonomic classifications. When there was an apparent mismatch, we re-arranged the association-level units among the branches without changing the membership of relevés to these basic units.

For each re-arrangement we checked the “validity” by re-calculating phi values and assessing the number and ecological meaning of the derived groups of diagnostic taxa. Having reached what we considered the optimal solution for the dataset under present knowledge of syntaxonomy in surrounding areas, we again determined diagnostic species at all hierarchical levels. We considered species as highly diagnostic when phi was  $> 0.50$  and as diagnostic when phi was  $> 0.25$  (see CHYTRÝ 2007), provided the concentration was significant according to Fisher’s exact test at  $\alpha = 0.05$ . When a species was diagnostic for several nested syntaxa according to this definition, it was normally assigned to the level where it reached the highest phi value. However, when the alternatives were assignment as diagnostic for one superior syntaxon or for several inferior syntaxa we chose the first. We considered as “transgressive” those diagnostic species that had a higher phi value at a lower rank but were also diagnostic at a higher rank, provided they had positive phi values in more than half of the associations of the higher unit. Finally, as phi values *per se* measure only the prevalence of a species within the whole data set but not the differentiation to the floristically similar units, we accepted species as diagnostic for one syntaxon only if the phi value there was at least 0.1 higher than in any other syntaxon of the same rank (LUTHER-MOSEBACH et al. 2012). Note that the term “diagnostic species” refers to the regional context and contains both differential and character species.

Finally, we compared our units to previously described phytosociological associations, alliances and orders, which involved an extensive search for original literature and check for validity and legitimacy of names according to the *International Code of Phytosociological Nomenclature* (WEBER et al. 2000; further ICPN). However, as there is a comprehensive classification study of all Central Balkan dry grassland associations in preparation (K. Vassilev et al. in prep.), we refrained from formally describing new syntaxa above association level. The nomenclatural revision (including some typifications) of the considered syntaxa is presented in Appendices 1 and S1. Authorities of the more thoroughly treated syntaxa (the accepted ones and some representing alternative views) can be found in Section 4.1 and Appendix S1, while for all other syntaxa we give them in the text at first mentioning unless they appear together with a bibliographic reference to the protologue.

### 3.4 Other statistical analyses

We carried out a detrended correspondence analysis (DCA) of the 10-m<sup>2</sup> plots with log-transformed percent cover values in CANOCO 4.5 (TER BRAAK & ŠMILAUER 2002) using down-weighting of rare species (TER BRAAK & ŠMILAUER 2002). Environmental factors and vegetation parameters were projected *post hoc* to this ordination diagram to assist with interpretation.

We tested for differences in environmental and vegetation parameters among the three distinguished classes and the ten associations with one-way analyses of variance (ANOVAs) carried out in STATISTICA 9 (STATSOFT INC. 2009). We tested whether the prerequisites of the ANOVA models (normal distribution, equal variance) were sufficiently met by visually inspecting the distribution of the residuals (QUINN & KEOUGH 2002). In addition, we used Tukey’s HSD *post hoc* test at  $\alpha = 0.05$  to identify significant differences among groups of syntaxa.

Further, we used the nested-plot data for a coarse analysis of species-area relationships (SARs). We did this with the power-law function applied with linear regression to log<sub>10</sub>-transformed area and richness data as proposed by DENGLER (2009a; for details see DENGLER et al. 2012a).

In the association descriptions (Section 4.4), we characterise the communities within the context of this work, using standardised terminology in relation to the mean value found in all our stands. For example, we classified species richness on 10 m<sup>2</sup> (where the overall mean was 38) as follows: 18–27 species: very species-poor; 28–37: moderately species-poor; 38–47: moderately species-rich and 48–57: very species-rich.

## 4. Vegetation classification

### 4.1 Proposed syntaxonomic scheme

In the following we present the syntaxonomic view of the data on which we base our further presentation, while various aspects are still tentative. Reasons for the applied scheme and alternative solutions are discussed in Sections 4.2 and 4.3. The synoptic table (Table 4 in the supplement) is based on this scheme.

Class 1: *Festuco-Brometea* Br.-Bl. & Tx. ex Klika & Hadač 1944

Order: *Stipo pulcherrimae-Festucetalia pallentis* Pop 1968

Alliance 1.1: *Saturejion montanae* Horvat et al. 1974

Association 1.1.1: *Dianthus petraeus-Sesleria latifolia*-[*Saturejion montanae*] community

Association 1.1.2: *Achillea clypeolata-Festuca stojanovii*-[*Saturejion montanae*] community

Association 1.1.3: *Satureja pilosa-Phleum phleoides*-[*Saturejion montanae*] community

Order: *Brachypodietalia pinnati* Korneck 1974

Alliance 1.2: *Cirsio-Brachypodion pinnati* Hadač & Klika in Klika & Hadač 1944

Association 1.2.1: *Ferulago campestris-Agrostis capillaris*-[*Cirsio-Brachypodion*] community

Association 1.2.2: *Tanacetum corymbosum-Sesleria latifolia*-[*Cirsio-Brachypodion*] community

Association 1.2.3: *Abietinella abietina-Sanguisorba minor*-[*Cirsio-Brachypodion*] community

Alliance 1.3: *Chrysopogono grylli-Danthonion calycinae* Kojić 1959

Association 1.3.1: *Agrostio capillaris-Chrysopogonetum grylli* Kojić 1959

Class 2: *Calluno-Ulicetea* Br.-Bl. & Tx. ex Klika & Hadač 1944

Order: *Nardetalia strictae* Preising 1950

Alliance 2.1: *Violion caninae* Schwickerath 1944

Association 2.1.1: *Festuco rubrae-Genistelletum sagittalis* Issler 1929

Class 3: *Koelerio-Corynephoretea* Klika in Klika & Novák 1941

Order: *Sedo-Scleranthetalia* Br.-Bl. 1955

Alliance 3.1: ??? (see discussion in Section 4.3.3)

Association 3.1.1: *Cetrario aculeatae-Plantaginetum subulatae* Pedashenko et al. 2013 (see Appendix 1)

Order: *Trifolio arvensis-Festucetalia ovinae* Moravec 1967

Alliance 3.2: *Armerio rumelicae-Potentillon* Micevski 1978

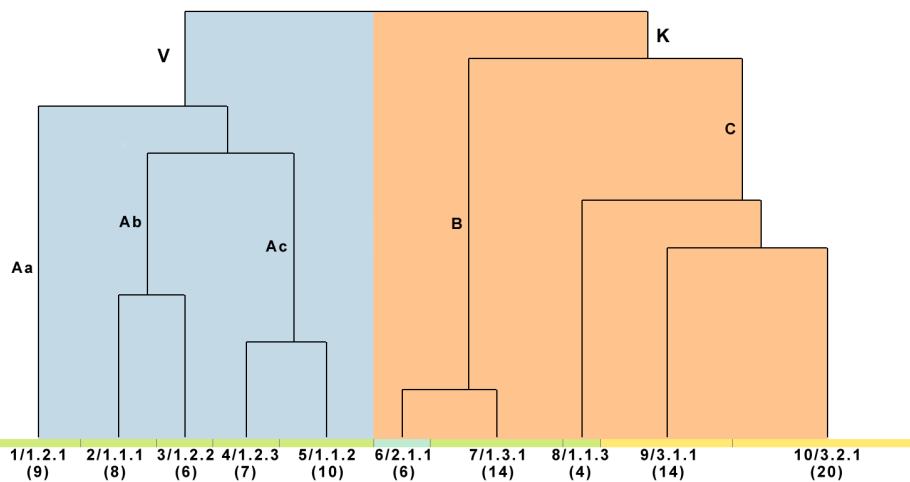
Association 3.2.1: *Plantagini subulatae-Agrostietum capillaris* Pedashenko et al. 2013 (see Appendix 1)

#### 4.2 Interpreting the numerical analyses in terms of syntaxonomy

At the 10-cluster resolution, modified TWINSpan yielded units that were floristically well characterised and therefore seem to warrant the rank of association (Fig. 3). Already the 11-cluster resolution (which split the unit 3.2.1 into two subunits; not shown) resulted in poorly characterised units, i.e. where many relevés were assigned to one of these subunits without a single diagnostic species present, and from the 12-cluster resolution onwards, additionally “chaining” occurred. Therefore we accepted the 10 clusters of the numerical analysis as association-level units.

When interpreting the cluster tree (Fig. 3), it turned out that at the higher levels, it was dominated by geographic differentiation which may or may not parallel ecological differences. For example, the highest division corresponded exactly to the two study areas, clusters 1–5 comprising only relevés from Vratsa and clusters 6–10 only relevés from Koprivshitsa. At the level of three clusters, there was a rough correspondence to phytosociological classes, with unit A (Clusters 1–5) comprising exclusively *Festuco-Brometea* communities, unit B (Clusters 6 and 7) being mainly *Calluno-Ulicetea* communities and unit C (Clusters 8–10) mainly *Koelerio-Corynephoretea* communities.

At closer look, however, Cluster 7, while having some similarities to Cluster 6, lacks all the more narrowly distributed *Calluno-Ulicetea* species and is instead characterised by wide-spread species of meso-xeric, nutrient-poor pastures from the classes *Festuco-Brometea*,



**Fig. 3.** Cluster tree of the 98 relevés achieved with modified TWINSpan at the 10-cluster resolution, which we interpret as association level in this paper. The first division corresponds exactly to the two study areas, Vratsa (V) and Koprivshitsa (K). At the bottom we provide for each cluster its original number in TWINSpan, its hierarchical position in our syntaxonomic interpretation and, in brackets, the number of relevés belonging to it.

**Abb. 3.** Clusterbaum der 98 Vegetationsaufnahmen basierend auf der modifizierten TWINSpan-Methode mit den 10 Endclustern, die wir in dieser Publikation als Assoziationen interpretieren. Die erste Teilung entspricht genau den beiden Untersuchungsgebieten, Vratsa (V) und Koprivshitsa (K). Am unteren Ende der Abbildung ist für jedes Endcluster die ursprüngliche TWINSpan-Nummer, die hierarchische Position in unserer syntaxonomischen Interpretation und in Klammern die Anzahl der zugehörigen Aufnahmen.



*Koelerio-Corynephoretea*, *Molinio-Arrhenatheretea* Tx. 1937 and *Calluno-Ulicetea*, with a slight prevalence of the first group (including the often dominating *Chrysopogon gryllus*). Further, Cluster 8, grouped by modified TWINSpan together with the two *Koelerio-Corynephoretea* communities, while containing indeed many typical species of that class (*Rumex acetosella*, *Scleranthus perennis*), still had a higher fraction of *Festuco-Brometea* species (e.g. *Phleum phleoides*, *Koeleria macrantha* agg., *Asperula purpurea*). Within the main unit A, the three TWINSpan subunits based on their diagnostic species were hardly interpretable in ecological terms and rather seemed to reflect floristic peculiarities of the individual study sites. Subunits A.b and A.c contained one xeric and one meso-xeric cluster each. In order to achieve better ecological interpretability and fit to supraregional classification schemes (ROYER 1991, RODWELL et al. 2002, DENGLER 2003), we therefore decided to combine the three meso-xeric clusters from unit A (i.e. Clusters 1, 3 and 4) into one alliance and the two xeric ones (i.e. Clusters 2 and 5) together with Cluster 8 from unit C into another alliance. Cluster 7 (from unit B), albeit being clearly meso-xeric (as shown by the occurrence of many *Molinio-Arrhenatheretea* species), lacked most of the diagnostic species of the before-mentioned meso-xeric alliance, while at the same time containing more acidophilous species. We therefore consider Cluster 7 as belonging to a third *Festuco-Brometea* alliance.

### 4.3 Identity of orders and alliances

#### 4.3.1 *Festuco-Brometea*

According to the traditional concept (e.g. BRAUN-BLANQUET & TÜXEN 1943, ROYER 1991, POTT 1995), the class *Festuco-Brometea* was organised into geographically defined orders, with the *Brometalia erecti* comprising the units in the more oceanic parts of Europe and the *Festucetalia valesiaca* the units in the more continental parts of Europe. More recent treatments instead oppose one geographically widespread meso-xeric order (*Brometalia erecti* = *Brachypodietalia pinnati*) to several xerophytic orders such as *Festucetalia valesiaca* Br.-Bl. & Tx. ex Br.-Bl. 1950 (continental lowland steppes of deep soils), *Stipo pulcherrimae-Festucetalia pallentis* (rocky dry grasslands of the Alps, Carpathians and surrounding mountains with many dealpine species) and *Artemisio albae-Brometalia erecti* Ubaldi ex Dengler & Mucina in Mucina et al. 2009 (communities of the submediterranean-subatlantic regions) (e.g. MUCINA & KOLBEK 1993, RODWELL et al. 2002, DENGLER 2004c, MUCINA et al. 2009, DENGLER et al. 2012a).

The so far only supra-national classifications of the *Festuco-Brometea* of the Central Balkans (HORVAT et al. 1974, ROYER 1991) were still based on the first concept and accordingly assigned all units to a geographically defined order *Festucetalia valesiaca*. HORVAT et al. (1974) accepted a total of three alliances within this order, which in the more modern system would nicely correspond to three orders, namely the *Festucion rupicolae* to the *Festucetalia valesiaca*, the *Chrysopogono-Danthonion* to the *Brachypodietalia pinnati* and the *Saturejion montanae* to the *Stipo pulcherrimae-Festucetalia pallentis*. ROYER (1991) presented a somewhat intermediate solution by placing all associations in the eastern half of Europe including those from the Central Balkans in the *Festucetalia valesiaca*, but grouping them into the suborders *Filipendulo-Avenulenalia*, *Stipo-Festucenalia valesiaca* and *Seslerio-Festucenalia pallentis*, which correspond to the orders *Brachypodietalia pinnati*, *Festucetalia valesiaca* and *Stipo-Festucetalia pallentis*, respectively. The situation was complicated by the proposal of a specific Central Balkan order *Astragalo-Potentilletalia* by



MICEVSKI (1971), to which later some authors subordinated any type of dry grasslands from the region, whether xeric or meso-xeric, basiphilous or acidophilous (e.g. MICEVSKI 1978, REDŽIĆ 1999, BERGMEIER et al. 2009). TZONEV et al. (2009) finally merged the various incompatible approaches of class subdivisions and thus ended up with not less than six orders for the territory of Bulgaria.

For the time being and based on the accumulated evidence, we consider the acceptance of the three orders that are well supported in recent Central European treatments (e.g. MUCINA & KOLBEK 1993, DENGLER et al. 2012a) as well as in a Europe-wide analysis of synoptic tables (J. Dengler unpubl.), i.e. *Brachypodietalia pinnati* (meso-xeric), *Festucetalia valesiaca* (xeric, deep soils, lowlands) and *Stipo-Festucetalia pallentis* (xeric, rocky soils, mountains), also for Bulgaria as the most parsimonious approach, with the first and the last being present in our study areas.

In our dataset alliance 1.1 represents the *Stipo-Festucetalia pallentis* and could be identified with the *Saturejion montanae* described from Serbia. While the *Saturejion montanae* is accepted in several recent Bulgarian treatments (SOPOTLIEVA 2008, PEDASHENKO et al. 2009, TODOROVA & TZONEV 2010, VASSILEV 2012, VASSILEV et al. 2012a), it was typically subordinated to the *Festucetalia valesiaca* as in the original description (HORVAT et al. 1974). TZONEV et al. (2009) assigned the alliance under the illegitimate name *Chrysopogono grylli-Saturejion subspicatae* Horvat & Horvatić in Horvat et al. 1974 (see TERZI 2011) to the amph-Adriatic order *Scorzonero villosae-Chrysopogonetalia grylli* Horvatić & Horvat in Horvatić 1963. To our knowledge, *Stipo-Festucetalia pallentis* were so far only once reported from Bulgaria by TZONEV et al. (2009), who assigned the alliance *Pimpinello-Thymion zygoidis* Dihoru 1970 from the Black Sea coast to this order, which is not supported floristically or ecologically according to usual concepts of the order as Circum-Pannonian and dealpine, mostly primary grasslands on steep rock cliffs. In contrast, the relevés by JOVANOVIĆ-DUNCIĆ (1955), on which HORVAT et al. (1974) mostly based their description of the *Saturejion montanae*, and our relevés fit well into the concept of the order. There is a high overlap between the diagnostic species of our study and those given for the order by MUCINA & KOLBEK (1993) for Austria and DENGLER et al. (2012a) for Transylvania, namely *Allium flavum*, *Melica ciliata*, *Sedum hispanicum*, *Sesleria rigida*, *Stipa pennata* subsp. *eriocaulis* and *Teucrium montanum*. The stands in Northern Bulgaria (see also PEDASHENKO et al. 2009, TODOROVA & TZONEV 2010, VASSILEV et al. 2012a) show some similarities to the Transylvanian *Seslerion rigidae* (compare DENGLER et al. 2012a) but are floristically sufficiently different to be considered a geographically vicariant alliance in the same order. Among our three associations, the first fits best into the *Saturejion montanae*, whereas this placement is not fully clear for the second and third. At least some relevés in the Association 1.1.2 show transitional character to the *Festucion valesiaca* (*Festucetalia valesiaca*) through the higher share of species such as *Koeleria macrantha* agg., *Poa angustifolia* or *Scabiosa ochroleuca*, which are typically more abundant in secondary steppe grasslands on deeper soils. The relevés of Association 1.1.3 are only loosely connected to the alliance and class, while several acidophilous *Koelerio-Corynephoretea* species are frequent (e.g. *Rumex acetosella*, *Plantago subulata*, *Scleranthus perennis*). In a large-scale synthesis, it might be justified to place this association in an alliance of its own, but for the time being, we prefer to leave these stands with dominant *Satureja pilosa* in the *Saturejion montanae* as did BERGMEIER et al. (2009) with stands of the same species in Northern Greece.

Alliance 1.2 could clearly be identified by its species composition with the *Cirsio-Brachypodium pinnati* as it has been extensively described in the eastern parts of central Europe (MUCINA & KOLBEK 1993, CHYTRÝ et al. 2007, ŠKODOVÁ 2007, DENGLER et al. 2012a). This alliance was not listed in the syntaxonomic overview of Bulgaria by TZONEV et al. (2009) but had been reported in three more recent studies (PEDASHENKO et al. 2010, VASSILEV 2012, VASSILEV et al. 2012a).

Alliance 1.3 could be identified with the *Chrysopogono grylli-Danthonion calycinae* described from Serbia (KOJIĆ 1959) and already reported several times from Bulgaria (APOSTOLOVA & MESHINEV 2006, TZONEV et al. 2009, VASSILEV et al. 2012a). The syntaxonomic assignment of this alliance to an order is debated controversially. APOSTOLOVA & MESHINEV (2006) still follow the traditional concept of the original description and place it in the *Festucetalia valesiaca*, TZONEV et al. (2009) are in favour of a placement in the acidophilous order *Koelerio-Phleetalia phleoidis* Korneck 1974, while BERGMEIER et al. (2009) prefer the Balkan order *Astragalo-Potentilletalia*. As the alliance has a clearly meso-xeric character and there are still doubts whether *Koelerio-Phleetalia* (see e.g. DENGLER 2004b) and *Astragalo-Potentilletalia* (see above) are justified as orders at all, we consider a preliminary placement in the *Brachypodietalia pinnati* presently as most parsimonious.

#### 4.3.2 *Calluno-Ulicetea*

Alliance 2.1 shows a perfect match with the Central European alliance *Violion caninae*, the matt-grass swards of the lowland to montane belts (compare PEPLER-LISBACH & PETERSEN 2001, KRAHULEC et al. 2007, UJHÁZY 2007). Astonishingly, neither the alliance nor the order *Nardetalia strictae* or the class *Calluno-Ulicetea* are given for Bulgaria in the overview by TZONEV et al. (2009). Instead only the *Potentillo ternatae-Nardion* is indicated, which is a group of subalpine to alpine matt-grass communities typically placed in the class *Juncetea trifidi* and clearly not the same as our stands, which essentially lack alpine species.

#### 4.3.3 *Koelerio-Corynephoretea*

The class *Koelerio-Corynephoretea* so far found very little attention on the Balkan Peninsula. In HORVAT et al. (1974), only sandy steppes along the Danube and its tributaries as well as at the Black Sea coast are briefly mentioned (mainly *Festucion vaginatae* Soó 1929), while TZONEV et al. (2009) – in addition to two alliances of sandy steppes (placed in the *Festuco-Brometea* by them) – report only two other *Koelerio-Corynephoretea* alliances for the territory of Bulgaria: *Alysso-Sedion* Oberd. & T. Müller in T. Müller 1961 and *Thero-Airion* Tx. ex Oberd. 1957. At the European scale, synthetic works comparable to that of ROYER (1991) for the *Festuco-Brometea* are missing. We only can refer to the supranational syntheses developed by MORAVEC (1967) and those of the senior author of this paper (largely unpublished, but see DENGLER 2003, 2004a). The latter author accepts for those large parts of Europe, for which he compiled synoptic tables, eight orders, two in the subclass *Sedo-Scleranthenea* (Br.-Bl. 1955) Dengler in Dengler et al. 2003 (communities of shallow skeletal soils) and six in the subclass *Koelerio-Corynephorenea* (Klika in Klika & V. Novák 1941) Dengler in Dengler et al. 2003 (communities of deep sandy soils).

Following the concept of DENGLER (2003, 2004a) and similarly that of MORAVEC (1967), alliance 3.1 as an open cryptogam-rich community of skeletal soils derived from acidic bedrock belongs to the order *Sedo-Scleranthetalia*. Within this order, DENGLER (2003) accepts three geographically vicariant alliances: *Sedo albi-Veronicion dillenii* (often errone-

ously termed *Arabidopsis thalianae*) in low mountain ranges of temperate Europe, *Sedo-Scleranthion* Br.-Bl. 1955 in the Alps and *Sedion pyrenaici* Tx. ex Rivas-Martínez et al. in Díaz González & Fernández Prieto 1994 on the Iberian Peninsula. Recently BERGMEIER et al. (2009) described the new alliance *Diantho pinifolii-Jasionion heldreichii* from northern Greece, but their data are hardly comparable because they did not record bryophytes and lichens, though they make up approximately half of the species composition of such communities. Whether our association belongs to the *Sedo albi-Veronicion dillenii*, to the *Diantho pinifolii-Jasionion heldreichii* or even to an undescribed third alliance, can only be answered by analysing a more comprehensive dataset from a larger area. With the strong representation of *Plantago subulata* in terms of both frequency and cover, a species that is absent from the previously known associations of both alliances, our association reminds of the Iberian alliance *Hieracio castellani-Plantaginion radicatae* Rivas-Martínez & Cantó 1987 (*Plantago radicata* = *P. subulata*) in terms of ecology, physiognomy and floristics (see DENGLER 2004c), an alliance that is placed into the separate Iberian order *Jasiono sesseliflorae-Koelerietalia crassipedis* Rivas-Martínez & Cantó 1987 (DENGLER 2003).

Alliance 3.2 shows some resemblance with the Atlantic order *Thero-Airetalia* Rivas Goday 1964 – which is indeed present in Bulgaria with the *Vulpietum myuri* Philippi 1973 (SOPOTLIEVA 2008) – via the presence of *Vulpia myurus* and *Aira elegantissima*. Overall, however, the closed sward, the prevalence of hemicryptophytes over annuals, the presence of typical species such as *Potentilla argentea* agg., *Trifolium arvense* and *T. campestre* and the frequent dominance of *Agrostis capillaris* support the subordination to the *Trifolio arvensis-Festucetalia ovinae* (MORAVEC 1967, WEEDA et al. 1996, DENGLER 2003, 2004a), an order which, to our knowledge, has never before been reported from the Balkan Peninsula. DENGLER (2003, 2004a) accepts three largely vicariant alliances in temperate Europe: *Sedo-Cerastion arvensis* Sissingh & Tideman 1960 in the Atlantic parts, *Hyperico perforati-Scleranthion perennis* Moravec 1967 in the mountains of Central Europe, on the British Isles and in Fennoscandia and *Armerion elongatae* Pötsch 1962 in the subcontinental lowlands. Evidently, none of these alliances fits the communities in Bulgaria geographically or floristically. The only alliance with some resemblance is the *Armerio rumelicae-Potentillion* described by MICEVSKI (1978) from siliceous mountains in Macedonia and placed in the *Astragalo-Potentilletalia (Festuco-Brometea)*. While the two associations included in the protologue show some transitional character between the *Festuco-Brometea* and the *Koelerio-Corynephoretea*, the selected type association (*Genisto carinalis-Agrostietum byzanthinae*, see Appendix S1) is clearly dominated by species of the latter class (e.g. *Aira elegantissima*, *Trifolium arvense*, *Rumex acetosella*, *Potentilla argentea* agg., *Scleranthus perennis*). Typified that way, we consider the *Armerio rumelicae-Potentillion* as belonging to the *Trifolio arvensis-Festucetalia ovinae* and identical with our alliance 3.2 as indicated by many floristic coincidences (e.g. the high frequency of *Plantago subulata*). Whether the other association placed by MICEVSKI (1978) in this alliance (*Koelerio macranthae-Festucetum stajanovii*) also completely belongs to the alliance emended in such a way, should to be tested in a large-scale analysis.

#### 4.4 Description of the associations (Tables 5 and 6 in the supplement, Tables 7 and 8)

While we are convinced that each of the 10 distinguished clusters warrants association status and some on the basis of more data could be split even further, there are only few valid names available that have been published for dry grassland syntaxa in Bulgaria or

**Table 7.** Characteristics of the three dry grassland classes (FB = *Festuco-Brometea*, CU = *Calluno-Ulicetea*, KC = *Koelerio-Corynephoretea*). Means and standard deviations are given. Different letters indicate significant differences between communities at  $\alpha = 0.05$  from Turkey's HSD test; *P*-values derived from ANOVAs.

**Tabelle 7.** Eigenschaften der Bestände der drei Trockenrasen-Klassen (FB = *Festuco-Brometea*, CU = *Calluno-Ulicetea*, KC = *Koelerio-Corynephoretea*). Es sind Mittelwerte und Standardabweichungen angegeben. Unterschiedliche Kleinbuchstaben bezeichnen signifikante Unterschiede ( $\alpha = 0,05$ ) zwischen den Klassen gemäß Tukeys HSD-Test; in der letzten Spalte sind die *P*-Werte aus den Varianzanalysen angegeben.

Parameter	FB	CU	KC	<i>P</i>
No. of relevés ( <i>n</i> )	58	6	34	
Altitude (m a.s.l.)	1108 ± 219 <sup>b</sup>	1059 ± 12 <sup>ab</sup>	910 ± 154 <sup>a</sup>	<0.001
Inclination (°)	18 ± 14	17 ± 8	16 ± 8	0.639
Heat index	0.08 ± 0.43	-0.19 ± 0.21	0.03 ± 0.21	0.201
Microrelief (cm)	15.3 ± 15.1	8.2 ± 2.1	11.9 ± 8.9	0.267
pH (H <sub>2</sub> O) of the soil	6.2 ± 1.1 <sup>b</sup>	4.9 ± 0.3 <sup>a</sup>	5.0 ± 0.4 <sup>a</sup>	<0.001
Conductivity (µS/cm) of the soil	177 ± 76 <sup>a</sup>	239 ± 288 <sup>a</sup>	72 ± 33 <sup>b</sup>	<0.001
Loss at ignition of the soil (%)	19.4 ± 8.3 <sup>b</sup>	11.6 ± 4.2 <sup>a</sup>	7.6 ± 3.3 <sup>a</sup>	<0.001
Proportion of sand in the soil (%)	75.5 ± 9.8	78.7 ± 3.8	78.0 ± 6.0	0.325
Proportion of silt in the soil (%)	22.9 ± 9.6 <sup>b</sup>	19.8 ± 2.7 <sup>ab</sup>	18.6 ± 6.4 <sup>a</sup>	0.056
Proportion of clay in the soil (%)	0.5 ± 0.8 <sup>b</sup>	0.1 ± 0.1 <sup>ab</sup>	0.1 ± 0.1 <sup>a</sup>	<0.001
Total vegetation cover (%)	82 ± 19 <sup>a</sup>	93 ± 4 <sup>a</sup>	73 ± 18 <sup>b</sup>	0.012
Cover of herb layer (%)	80 ± 22 <sup>a</sup>	92 ± 6 <sup>a</sup>	57 ± 23 <sup>b</sup>	<0.001
Cover of cryptogam layer (%)	5 ± 7 <sup>a</sup>	3 ± 2 <sup>a</sup>	22 ± 19 <sup>b</sup>	<0.001
Cover of litter (%)	26 ± 25 <sup>b</sup>	8 ± 2 <sup>ab</sup>	11 ± 13 <sup>a</sup>	<0.001
Cover of stones and rocks (%)	11 ± 19	0 ± 1	13 ± 14	0.243
Cover of bare soil (%)	5 ± 11	3 ± 5	11 ± 12	0.053
Species richness of all taxa (10 m <sup>2</sup> )	41.3 ± 10.7 <sup>b</sup>	41.0 ± 7.4 <sup>ab</sup>	33.2 ± 10.7 <sup>a</sup>	0.002
Species richness of vascular plants (10 m <sup>2</sup> )	38.0 ± 10.2 <sup>b</sup>	37.0 ± 7.3 <sup>ab</sup>	26.9 ± 9.4 <sup>a</sup>	<0.001
Species richness of bryophytes (10 m <sup>2</sup> )	2.3 ± 2.2	3.2 ± 1.5	2.9 ± 1.7	0.268
Species richness of lichens (10 m <sup>2</sup> )	0.9 ± 1.7 <sup>a</sup>	0.8 ± 1.7 <sup>a</sup>	3.5 ± 2.7 <sup>b</sup>	<0.001

neighbouring countries. When we found a convincing match, we apply the association names here; otherwise we use an informal name, discuss potential matches under “Syntaxonomy” and leave the resolution to the planned large-scale analysis (K. Vassilev et al. in prep.). Only in two cases where the literature review yielded no close resemblance while we could document the same type from multiple places with more than 10 relevés in total, we provide formal descriptions of new associations.

**Association 1.1.1:** *Dianthus petraeus*-*Sesleria latifolia*-[*Saturejion montanae*] community (Fig. 4)

**Characterisation:** Moderately species-rich community with open horizontal structure, dominated by *Sesleria latifolia*, *Potentilla cinerea* agg., *Dianthus petraeus*, *Trifolium alpestre*, *Helianthemum nummularium* and *Carex humilis*. The cryptogam layer is weakly developed without any species dominating over others.

**Ecology and distribution:** This community mostly grows on steep east or south-east inclined terrains (mean heat index: 0.65) at the highest altitudes in the study area (mean elevation: 1409 m a.s.l.). Soils are derived from calcareous bedrock, with many outcrops at the

**Table 8.** Characteristics of the ten dry grassland associations. Means and standard deviations are given. Different letters indicate significant differences between communities at  $\alpha = 0.05$  from Turkey's HSD test; P-values derived from ANOVAs.

**Table 8.** Eigenschaften der Bestände der drei Trockenrasen-Assoziationen. Es sind Mittelwerte und Standardabweichungen angegeben. Unterschiedliche Kleinbuchstaben bezeichnen signifikante Unterschiede ( $\alpha = 0,05$ ) gemäß Tukeys HSD-Test; in der letzten Spalte sind die P-Werte aus den Varianzanalysen angegeben.

Parameter	Assoc. 1.1.1	Assoc. 1.1.2	Assoc. 1.1.3	Assoc. 1.2.1	Assoc. 1.2.2	Assoc. 1.2.3	Assoc. 1.3.1	Assoc. 2.1.1	Assoc. 3.1.1	Assoc. 3.2.1	P
No. of relevés (n)	8	10	4	9	6	7	14	6	14	20	
Altitude (m a.s.l.)	1409 ± 39 <sup>c</sup>	1003 ± 34 <sup>a</sup>	819 ± 202 <sup>a</sup>	1271 ± 125 <sup>b</sup>	1275 ± 163 <sup>bc</sup>	1083 ± 66 <sup>ac</sup>	929 ± 149 <sup>a</sup>	1059 ± 12 <sup>ac</sup>	901 ± 128 <sup>a</sup>	916 ± 172 <sup>a</sup>	<0.001
Inclination (°)	30 ± 25	19 ± 14	31 ± 5	13 ± 7	19 ± 17	12 ± 5	14 ± 7	17 ± 8	17 ± 9	15 ± 8	0.019
Heat index	0.65 ± 0.71 <sup>b</sup>	-0.14 ± 0.28 <sup>a</sup>	0.33 ± 0.25 <sup>ab</sup>	-0.06 ± 0.17	-0.10 ± 0.54 <sup>a</sup>	0.05 ± 0.17	0.03 ± 0.19 <sup>a</sup>	-0.19 ± 0.21 <sup>a</sup>	0.09 ± 0.27 <sup>a</sup>	-0.01 ± 0.16 <sup>a</sup>	<0.001
Microrelief (cm)	22.9 ± 16.1 <sup>bc</sup>	28.5 ± 18.1 <sup>c</sup>	32.5 ± 15.0 <sup>c</sup>	11.8 ± 9.9 <sup>ab</sup>	13.0 ± 13.1 <sup>abc</sup>	7.0 ± 4.2 <sup>ab</sup>	4.1 ± 2.6 <sup>a</sup>	8.2 ± 2.1 <sup>ab</sup>	16.0 ± 11.3 <sup>abc</sup>	9.1 ± 5.5 <sup>ab</sup>	<0.001
pH (H <sub>2</sub> O) of the soil	7.0 ± 0.6 <sup>b</sup>	7.3 ± 0.4 <sup>b</sup>	5.3 ± 0.6 <sup>a</sup>	5.3 ± 0.3 <sup>a</sup>	6.8 ± 0.9 <sup>b</sup>	7.3 ± 0.7 <sup>b</sup>	5.0 ± 0.2 <sup>a</sup>	4.9 ± 0.3 <sup>a</sup>	4.8 ± 0.3 <sup>a</sup>	5.1 ± 0.4 <sup>a</sup>	<0.001
Conductivity (µS/cm) of the soil	246 ± 36 <sup>b</sup>	189 ± 38 <sup>ab</sup>	74 ± 25 <sup>ac</sup>	161 ± 78 <sup>abc</sup>	216 ± 90 <sup>ab</sup>	213 ± 79 <sup>ab</sup>	136 ± 62 <sup>abc</sup>	239 ± 288 <sup>ab</sup>	61 ± 22 <sup>c</sup>	81 ± 37 <sup>c</sup>	<0.001
Loss at ignition of the soil (%)	29.7 ± 8.5 <sup>e</sup>	23.6 ± 5.0 <sup>bc</sup>	3.9 ± 1.7 <sup>a</sup>	21.0 ± 4.7 <sup>b</sup>	23.4 ± 3.4 <sup>bc</sup>	18.3 ± 3.1 <sup>b</sup>	12.9 ± 4.0 <sup>cd</sup>	11.6 ± 4.2 <sup>cd</sup>	7.0 ± 3.6 <sup>a</sup>	8.1 ± 3.0 <sup>ac</sup>	<0.001
Proportion of sand in the soil (%)	81.5 ± 6.6 <sup>a</sup>	79.5 ± 4.8 <sup>ab</sup>	75.7 ± 2.8 <sup>ab</sup>	74.0 ± 10.4 <sup>a</sup>	71.2 ± 17.7 <sup>ab</sup>	67.0 ± 12.2 <sup>ab</sup>	76.3 ± 6.1 <sup>ab</sup>	78.7 ± 3.8 <sup>ab</sup>	79.6 ± 4.4 <sup>ab</sup>	76.9 ± 6.8 <sup>a</sup>	0.021
Proportion of silt in the soil (%)	18.1 ± 6.9 <sup>ab</sup>	19.4 ± 4.7 <sup>ab</sup>	23.1 ± 2.2 <sup>ab</sup>	24.8 ± 9.9 <sup>ab</sup>	25.1 ± 18.1 <sup>ab</sup>	31.5 ± 11.3 <sup>ab</sup>	21.8 ± 6.8 <sup>ab</sup>	19.8 ± 2.7 <sup>ab</sup>	16.7 ± 5.2 <sup>a</sup>	20.0 ± 6.9 <sup>a</sup>	0.016
Proportion of clay in the soil (%)	0.1 ± 0.2 <sup>ab</sup>	0.6 ± 0.4 <sup>abc</sup>	0.2 ± 0.2 <sup>abc</sup>	0.8 ± 0.8 <sup>bcd</sup>	1.0 ± 1.2 <sup>cd</sup>	1.4 ± 1.1 <sup>d</sup>	0.0 ± 0.1 <sup>a</sup>	0.1 ± 0.1 <sup>abc</sup>	0.0 ± 0.0 <sup>a</sup>	0.1 ± 0.1 <sup>a</sup>	<0.001
Total vegetation cover (%)	77 ± 19 <sup>abc</sup>	72 ± 16 <sup>ac</sup>	40 ± 20 <sup>d</sup>	97 ± 2 <sup>b</sup>	94 ± 7 <sup>ab</sup>	87 ± 10 <sup>ab</sup>	89 ± 10 <sup>ab</sup>	93 ± 4 <sup>ab</sup>	61 ± 17 <sup>cd</sup>	81 ± 15 <sup>ab</sup>	<0.001
Cover of herb layer (%)	70 ± 30 <sup>ab</sup>	70 ± 16 <sup>ab</sup>	33 ± 15 <sup>d</sup>	97 ± 2 <sup>c</sup>	93 ± 8 <sup>abc</sup>	84 ± 10 <sup>abc</sup>	87 ± 11 <sup>bc</sup>	92 ± 6 <sup>abc</sup>	39 ± 17 <sup>d</sup>	70 ± 17 <sup>a</sup>	<0.001
Cover of cryptogam layer (%)	8 ± 11 <sup>ab</sup>	4 ± 5 <sup>ab</sup>	8 ± 9 <sup>abc</sup>	2 ± 5 <sup>ab</sup>	2 ± 3 <sup>ab</sup>	7 ± 11 <sup>ab</sup>	3 ± 7 <sup>a</sup>	3 ± 2 <sup>ab</sup>	28 ± 14 <sup>c</sup>	18 ± 21 <sup>bc</sup>	<0.001
Cover of litter (%)	18 ± 14 <sup>ab</sup>	15 ± 21 <sup>ab</sup>	1 ± 1 <sup>a</sup>	37 ± 24 <sup>bc</sup>	28 ± 26 <sup>abc</sup>	11 ± 10 <sup>ab</sup>	46 ± 25 <sup>c</sup>	8 ± 2 <sup>ab</sup>	4 ± 4 <sup>a</sup>	16 ± 15 <sup>ab</sup>	<0.001
Cover of stones and rocks (%)	17 ± 20 <sup>abcd</sup>	28 ± 22 <sup>cd</sup>	43 ± 29 <sup>d</sup>	0 ± 1 <sup>a</sup>	3 ± 6 <sup>ab</sup>	6 ± 7 <sup>abc</sup>	0 ± 0 <sup>a</sup>	0 ± 1 <sup>ab</sup>	19 ± 13 <sup>bcd</sup>	8 ± 13 <sup>ab</sup>	<0.001
Cover of open soil (%)	6 ± 4 <sup>ab</sup>	4 ± 3 <sup>ab</sup>	31 ± 34 <sup>c</sup>	2 ± 2 <sup>a</sup>	3 ± 3 <sup>ab</sup>	6 ± 6 <sup>ab</sup>	2 ± 2 <sup>a</sup>	3 ± 5 <sup>ab</sup>	16 ± 16 <sup>bc</sup>	7 ± 7 <sup>ab</sup>	<0.001
Species richness all taxa (10 m <sup>2</sup> )	40.8 ± 5.6 <sup>abcd</sup>	48.4 ± 10.0 <sup>d</sup>	27.3 ± 10.1 <sup>ac</sup>	41.9 ± 11.1 <sup>a</sup>	46.7 ± 7.4 <sup>abd</sup>	47.9 ± 7.9 <sup>b</sup>	34.5 ± 7.8 <sup>abc</sup>	41.0 ± 7.4 <sup>abcd</sup>	28.6 ± 6.6 <sup>c</sup>	36.4 ± 12.0 <sup>abc</sup>	<0.001
Species richness vascular plants (10 m <sup>2</sup> )	36.5 ± 5.0 <sup>abcd</sup>	44.8 ± 8.7 <sup>c</sup>	22.0 ± 7.5 <sup>de</sup>	39.8 ± 11.4 <sup>a</sup>	43.2 ± 8.9 <sup>abc</sup>	43.6 ± 8.2 <sup>b</sup>	32.5 ± 7.7 <sup>abd</sup>	37.0 ± 7.3 <sup>abcd</sup>	20.8 ± 6.3 <sup>c</sup>	31.1 ± 9.1 <sup>ad</sup>	<0.001
Species richness bryophytes (10 m <sup>2</sup> )	2.0 ± 2.3	3.0 ± 2.7	2.8 ± 1.9	2.1 ± 1.8	3.2 ± 3.4	2.7 ± 1.7	1.4 ± 1.5	3.2 ± 1.5	3.1 ± 1.1	2.8 ± 2.0	0.441
Species richness lichens (10 m <sup>2</sup> )	2.1 ± 3.3 <sup>ab</sup>	0.6 ± 0.7 <sup>a</sup>	2.8 ± 2.5 <sup>ab</sup>	0.0 ± 0.0 <sup>a</sup>	0.3 ± 0.5 <sup>a</sup>	1.4 ± 1.3 <sup>a</sup>	0.6 ± 1.2 <sup>a</sup>	0.8 ± 1.6 <sup>a</sup>	4.9 ± 1.6 <sup>b</sup>	2.5 ± 2.9 <sup>a</sup>	<0.001

surface and with almost neutral reaction (mean pH: 7.00), high conductivity and organic matter content. Stands were recorded in the area of Beglichka mogila peak and above the village of Milanovo, where they were used as low-intensity pastures or not at all (because of steepness of the terrain or abandonment).

**Classification:** While this community type has not been described in phytosociological terms before, some decades ago VELCHEV (1971) documented similar types in the same region with the dominance approach. Accordingly, three of his “associations” more or less correspond to our community: “*Sesleria latifolia-Fetuca dalmatica*”, “*Sesleria latifolia-Stipa pennata*” and “*Sesleria latifolia-Satureja montana*”.

**Association 1.1.2:** *Achillea clypeolata-Festuca stojanovii*-[*Saturejion montanae*] community

**Characterisation:** Very species-rich community with high occurrence of the dwarf shrub *Satureja montana* subsp. *kitaibelii*. Stands are rich of forbs (e.g. *Teucrium chamaedrys*, *Asperula purpurea*, *A. cynanchica*, *Veronica austriaca* and *Euphorbia cyparissias*), while grasses (*Koeleria macrantha* agg., *Poa angustifolia*, *P. compressa* and *Festuca stojanovii*) play a lesser role. Bryophytes and lichens are poorly presented in terms of richness and cover.



**Fig. 4.** *Dianthus petraeus*-*Sesleria latifolia*-[*Saturejion montanae*] community (order *Stipo pulcherrimae-Festucetalia pallentis*) in Parshevitsa, near Vratsa. Among the recognisable plants are *Achillea clypeolata*, *Jovibarba heufelii* and *Satureja montana* subsp. *kitaibelii* (Photo: J. Dengler, JD115374).

**Abb. 4.** *Dianthus petraeus*-*Sesleria latifolia*-[*Saturejion montanae*]-Gesellschaft (Ordnung *Stipo pulcherrimae-Festucetalia pallentis*) in Parshevitsa, nahe Vratsa. Man erkennt unter anderem *Achillea clypeolata*, *Jovibarba heufelii* und *Satureja montana* subsp. *kitaibelii* (Foto: J. Dengler, JD115374).

**Ecology and distribution:** Occurs on both steep and slightly inclined calcareous terrains with rocky outcrops at south, west or north-west exposition and with rough microrelief. Soils are shallow, rich in skeleton material and sand fraction. Organic matter content is high, and soil reaction is neutral to slightly alkaline. Relevés were taken in the area of Kravya and on the slopes of the Okolchitsa peak on extensively used pastures.

**Classification:** Floristically and ecologically similar stands, which likely belong to the same association, have been described by TZONEV (2009) as *Potentillo pilosae-Achilleetum clypeolatae* from the Chernelka Canyon, approx. 80 km East of Vratsa. Unfortunately, this publication is not valid according to Art. 1 ICPN because it appeared only online but not in print.

**Association 1.1.3:** *Satureja pilosa-Phleum phleoides*-[*Saturejion montanae*] community

**Characterisation:** Very species-poor community developed on silicate terrains, characterised by open horizontal structure (mean total vegetation cover: 40%). Like in Assoc. 1.1.2, a dwarf shrub is dominating – in this case *Satureja pilosa*. Numerous grasses like *Koeleria macrantha* agg., *Phleum phleoides*, *Dichanthium ischaemum*, *Festuca valesiaca* and *Chrysopogon gryllus* add to the vegetation cover as well as a high number of lichens. Transgressive species to the class *Koelerio-Corynephoretea* like *Plantago subulata*, *Scleranthus perennis*, *Rumex acetosella* and *Ceratodon purpureus* are also abundant.

**Ecology and distribution:** Stands cover steep south- or southeast-facing slopes with rough microrelief and many outcrops on siliceous bedrock. The soils have a high proportion of skeleton, acid reaction (mean pH = 5.3), a low conductivity value and scarce organic matter. Stands of this community were found on slopes above the towns Anton and Klisura, which are unsuitable for agricultural use because of their high inclination.

**Classification:** The *Artemisia alba*-[*Saturejion montanae*] community described by BERGMEIER et al. (2009) from northern Greece has also frequent *Satureja pilosa* but otherwise only limited floristic overlap.

**Association 1.2.1:** *Ferulago campestris-Agrostis capillaris*-[*Cirsio-Brachypodion*] community

**Characterisation:** Moderately species-rich community with closed horizontal structure and strongly dominated by *Agrostis capillaris*. Other mesophilous grasses like *Anthoxanthum odoratum*, *Festuca nigrescens*, *Brachypodium pinnatum* and *Briza media* show much lower cover and constancy values unlike some forbs: *Trifolium alpestre*, *Stachys officinalis*, *Achillea millefolium* agg., *Primula veris* and *Galium verum*. The cryptogam layer is poorly presented, whereas accumulation of litter is substantial.

**Ecology and distribution:** Occurs on both steep and slightly inclined terrains with varying exposition and microrelief at high altitude (mean 1271 m a.s.l.). Soils are derived from calcareous bedrock, have a prevailing sandy texture and are rich in skeleton material and organic matter. Communities were found in the area of Okolchitsa, Beglichka mogila peak and above the village of Milanovo, where they were used as low-intensity pastures; some were already abandoned.

**Classification:** Relevé BGR024 is dominated by *Festuca paniculata* subsp. *paniculata* and might therefore justify placement in a separate association or even a different higher syntaxon. GRABHERR (1993), for example, considers this taxon in Austria as diagnostic for the order *Festucetalia spadiceae* Barbero 1970 within the class *Caricetea curvulae* Br.-Bl. 1948 (alpine-subalpine acidic grasslands), but as we had only one such relevé, we considered any decision premature.



**Association 1.2.2:** *Tanacetum corymbosum*-*Sesleria latifolia*-[*Cirsio-Brachypodion*] community (Fig. 5)

**Characterisation:** Moderately species-rich community with closed horizontal structure, dominated by *Sesleria latifolia* and many forbs like *Trifolium alpestre*, *Primula veris*, *Helianthemum nummularium*, *Teucrium chamaedrys*, *Stachys officinalis* and *Prunella grandiflora*. The *Sesleria* dominance is similar to Assoc. 1.1.1, with which it often grows in close contact, but the sward is much denser and the species composition more mesophytic. Bryophytes and lichens have very low cover values.

**Ecology and distribution:** This community grows on calcareous terrains at high altitude (mean 1275 m a.s.l.), mostly on steep north- and northwest-facing slopes. The soils have sandy texture, neutral reaction (mean pH = 6.8), high conductivity values and high organic matter content. Stands were found in the area of Beglichka mogila peak, Okolchitsa peak and above the village of Milanovo, where they were used as low-intensity pastures. This community might be a transitional stage between grassland and forest vegetation. *Tanacetum corymbosum* is ecologically much related to forest edges, and *Sesleria latifolia* occurs in different plant communities, including as herb layer under tree canopy, for example in the black pine communities, and has thus been considered diagnostic for the *Seslerio-Pinetum nigrae* Em 1962. Unlike many other plants growing beneath the canopy, *S. latifolia* can survive or even spread in open places. Due to its strong competitive ability and low forage quality, *S. latifolia* becomes dominant in grasslands under different environmental conditions.

**Classification:** To our knowledge, no similar community type has been described so far.



**Fig. 5.** *Tanacetum corymbosum*-*Sesleria latifolia*-[*Cirsio-Brachypodion*] community (order *Brachypodietalia pinnati*) in Parshevitsa, near Vratsa. Among the recognisable plants are *Vincetoxicum hirsundinaria*, *Tanacetum corymbosum* and *Brachypodium pinnatum* (Photo: J. Dengler, JD115377).

**Abb. 5.** *Tanacetum corymbosum*-*Sesleria latifolia*-[*Cirsio-Brachypodion*]-Gesellschaft (Ordnung *Brachypodietalia pinnati*) in Parshevitsa, nahe Vratsa. Man erkennt unter anderem *Vincetoxicum hirsundinaria*, *Tanacetum corymbosum* und *Brachypodium pinnatum* (Foto: J. Dengler, JD115377).



**Association 1.2.3:** *Abietinella abietina-Sanguisorba minor*-[*Cirsio-Brachypodium*] community

**Characterisation:** Very species-rich community with semi-closed horizontal structure and many legumes present: *Medicago lupulina*, *M. sativa* subsp. *falcata*, *Lotus corniculatus*, *Trifolium alpestre* and *T. pratense*. Stands do not show clear dominance, but *Teucrium chamaedrys* and *Achillea millefolium* have high cover values among the most constant species. Cryptogams are also well presented and form a distinct layer.

**Ecology and distribution:** Occurs on slightly inclined terrains with varying aspects on slightly alkaline grounds and high conductivity values. Soils are rich in skeleton material, and the sand/silt fraction is the lowest among all studied communities. Records were taken at the area of Kravya, above the village of Milanovo and on slopes of Okolchitsa peak on extensively used pastures

**Classification:** While the community composition mainly consists of common species so that the association could be expected to be widespread, we are not aware of any previous publication of such a vegetation type from Bulgaria.

**Association 1.3.1:** *Agrostio capillaris-Chrysopogonetum grylli* (Fig. 6)

**Characterisation:** Moderately species-poor community, dominated by grasses like *Chrysopogon gryllus*, *Agrostis capillaris*, *Anthoxanthum odoratum*, *Festuca valesiaca* and *F. rubra*. Frequent forbs are *Achillea millefolium* agg., *Carex caryophyllea*, *Galium verum* and *Plantago lanceolata*. Closed horizontal structure and dense litter cover prevent the formation of a cryptogam layer.



**Fig. 6.** *Agrostio capillaris-Chrysopogonetum grylli* (alliance *Chrysopogono grylli-Danthonion calycinae*, order *Brachypodietalia pinnati*) north of Koprivshtitsa. The dominating grass is *Chrysopogon gryllus* (Photo: J. Dengler, JD115601).

**Abb. 6.** *Agrostio capillaris-Chrysopogonetum grylli* (Verband *Chrysopogono grylli-Danthonion calycinae*, Ordnung *Brachypodietalia pinnati*) nördlich von Koprivshtitsa. Das dominante Gras ist *Chrysopogon gryllus* (Foto: J. Dengler, JD115601).

**Ecology and distribution:** Stands cover mostly slightly inclined northeast- or northwest-facing slopes (mean heat index: 0.03) on acid, sandy soils poor in organic matter. This community was recorded in several areas at lower altitudes (mean: 929 m a.s.l.): north of Koprivshitsa, near the towns Pirdop and Klisura as well as areas around Beklemeto mountain pass. Most stands were parts of large pastures used at very low intensity.

**Classification:** The *Agrostio-Chrysopogonietum* is the type association of the *Chrysopogono grylli-Danthonion calycinae* described in Serbia (KOJIĆ 1959) and so similar to our stands that, for the time being, it seems justified subordinating them here. According to VASSILEV (2012), at least 30 different associations have later been published within the alliance *Chrysopogono-Danthonion* throughout the Balkans, whose floristic differences are often only subtle. Therefore, after the intended critical plot-based revision, probably only few of them will remain.

**Association 2.1.1:** *Festuco rubrae-Genistelletum sagittalis* (Fig. 7)

**Characterisation:** Moderately species-rich community with closed horizontal structure, dominated by *Nardus stricta*, *Agrostis capillaris*, *Danthonia decumbens*, *Trifolium alpestre*, *Chamaespartium sagittale*, *Briza media*, *Anthoxanthum odoratum*, *Potentilla erecta* and *Viola canina*. Stands are rich in cryptogams, but they do not form a separate layer under the dense cover of grasses. Litter is weakly accumulated.

**Ecology and distribution:** This community mostly grows on west or northwest-inclined terrains on skeleton-rich, sandy soils derived from siliceous bedrock. Grounds are very acid (mean pH: 4.9), poor in organic matter and have high conductivity values. In the study areas, they were found only on low-intensity pastures south of the town of Koprivshitsa.



**Fig. 7.** *Festuco rubrae-Genistelletum sagittalis* (alliance *Violion caninae*, order *Nardetalia strictae*) south of Koprivshitsa. Among the recognisable plants are *Leontodon hispidus*, *Danthonia decumbens* and *Nardus stricta* (Photo: J. Dengler, JD115761).

**Abb. 7.** *Festuco rubrae-Genistelletum sagittalis* (Verband *Violion caninae*, Ordnung *Nardetalia strictae*) südlich von Koprivshitsa. Man erkennt unter anderem *Leontodon hispidus*, *Danthonia decumbens* und *Nardus stricta* (Foto: J. Dengler, JD115761).



**Classification:** Unexpectedly, the *Festuco-Genistelletum* described from the Alsace (France) and widespread in southern Germany (PEPLER-LISBACH & PETERSEN 2001, BECKER et al. 2012) matches the stand in Bulgaria to such a large degree that we suggest that they are one and the same association.

**Association 3.1.1:** *Cetrario aculeatae-Plantaginetum subulatae* (Fig. 8)

**Characterisation:** Moderately species-poor community, characterised by a high proportion of cryptogams: *Cladonia foliacea*, *Cetraria aculeata*, *Ceratodon purpureus*, *Polytrichum piliferum* and *Racomitrium canescens* agg., where lichens are dominating (mean cover of cryptogam layer: 29%). Stands have open horizontal structure (mean vegetation cover = 61%) and a low amount of accumulated litter. The most frequent vascular plants are *Plantago subulata*, *Rumex acetosella*, *Agrostis capillaris*, *Chrysopogon gryllus*, *Scabiosa triniifolia* and *Hypericum perforatum*.

**Ecology and distribution:** Occurs on both steep and slightly inclined terrains with varying exposition and microrelief and a high proportion of stones and rocks as well as bare soil. Grounds are derived from siliceous bedrock and very acid (mean pH = 4.8), poor in organic matter and of low conductivity. This community was found in several areas at lower altitudes (mean: 901 m a.s.l.): north of Koprivshitsa, near the towns Pirdop and Klisura as well as a stand around Beklemeto mountain pass in pastures of varying intensity. There the *Cetrario-Plantaginetum* is found at small scale within extensive stands of the *Agrostio-Chrysopogonetum* (Assoc. 1.3.1) and the *Plantagini-Agrostietum* (Assoc. 3.2.1) surrounding outcrops or on hillocks of coarse gravel.



**Fig. 8.** *Cetrario aculeatae-Plantaginetum subulatae* (alliance unknown, order *Sedo-Scleranthetalia*) in Klisura, near Koprivshitsa. Among the recognisable plants are *Plantago subulata* and *Scabiosa triniifolia*, *Polytrichum piliferum* and *Cladonia foliacea* (Photo: J. Dengler, JD115649).

**Abb. 8.** *Cetrario aculeatae-Plantaginetum subulatae* (Verband ungeklärt, Ordnung *Sedo-Scleranthetalia*) in Klisura, nahe Koprivshitsa. Man erkennt unter anderem *Plantago subulata* und *Scabiosa triniifolia*, *Polytrichum piliferum* und *Cladonia foliacea* (Foto: J. Dengler, JD115649).

**Classification:** To our knowledge a similar vegetation type has never before been documented from Bulgaria or surrounding countries. We therefore publish a new association here based on our 14 relevés (see Appendix 1).

**Association 3.2.1:** *Plantagini subulatae-Agrostietum capillaris* (Fig. 9)

**Characterisation:** Similar to the *Cetrario aculeatae-Plantaginetum subulatae* (Assoc. 3.1.1), but with a much more closed sward dominated by grasses. On the other hand, lichens play a much smaller role in the species composition, with *Cetraria aculeata* being largely missing.

**Ecology and distribution:** Occurs on slightly inclined terrains with varying aspects. Soils are acid (mean pH = 5.3), poor in organic matter and of low conductivity. This community is the most frequent type in the siliceous study area and was found in all localities there.

**Classification:** SOPOTLIEVA (2008) described in her PhD thesis a *Trifolio arvensis-Festucetum valesiaca* nom. inval. (according to Art. 1 ICPN, but validation in preparation) from East Bulgaria (about 180 km away). Her unit, which she placed in the *Festucion valesiaca* (*Festuco-Brometea*), shows some similarities with our stands, but while in her case *Festuco-Brometea* species prevail, in our case *Koelerio-Coryneporetea* species do. There have also been reports from the Sofia region based on the dominance approach and without species lists of a “*Dichantium ischaemum-Festuca valesiaca+Plantago carinata*” and “*Dichantium ischaemum-Trifolium alpestre+Plantago carinata*” (*Plantago carinata* = *P. subulata*) (see review by APOSTOLOVA & SLAVOVA 1997), which might mean a similar community. As our community is very frequent but was not formally described before, we publish it as a new association here based on our 20 relevés (see Appendix 1).



**Fig. 9.** *Plantagini subulatae-Agrostietum capillaris* (alliance *Armerio rumelicae-Potentillion*, order *Trifolio arvensis-Festucetalia ovinae*) north of Koprivshtitsa. Among the recognisable plants are *Jasione heldreichii*, *Armeria rumelica* and *Racomitrium canescens* agg. (Photo: J. Dengler, JD115603).

**Abb. 9.** *Plantagini subulatae-Agrostietum capillaris* (Verband *Armerio rumelicae-Potentillion*, Ordnung *Trifolio arvensis-Festucetalia ovinae*) nördlich von Koprivshtitsa. Man erkennt unter anderem *Jasione heldreichii*, *Armeria rumelica* und *Racomitrium canescens* agg. (Foto: J. Dengler, JD115603).

## 5. Ecology and biodiversity

### 5.1 Vegetation-environment relationships

The most common type of land use in the studied grasslands was grazing (77% of plots), with 87% of the active pastures being used at low intensity. Unused grasslands were either abandoned pastures and meadows or natural grasslands that showed no evidence of management. Mechanical properties of soils were similar in all sampled areas with prevalence of the sand fraction (mean 76.6%, range 37.4–91.0%). The soil chemistry of the two study areas differed considerably. In the area of Vratsa, soils were developed from carbonate bedrock and showed a high humus content (mean 23.3%, range 13.4–43.9%), were base-rich with a mean pH (H<sub>2</sub>O) of 6.7 (range 4.8–8.0) and had high conductivity values (mean 202  $\mu$ S/cm, range 78–380  $\mu$ S/cm). In the area of Koprivshitsa, soils were developed from silicate bedrock, had a low humus content (mean 9.1%, range 2.3–22.0%), were acid with a mean pH (H<sub>2</sub>O) of 5.0 (range 4.4–6.4) and had low conductivity values (mean 104  $\mu$ S/cm, range 19–311  $\mu$ S/cm).

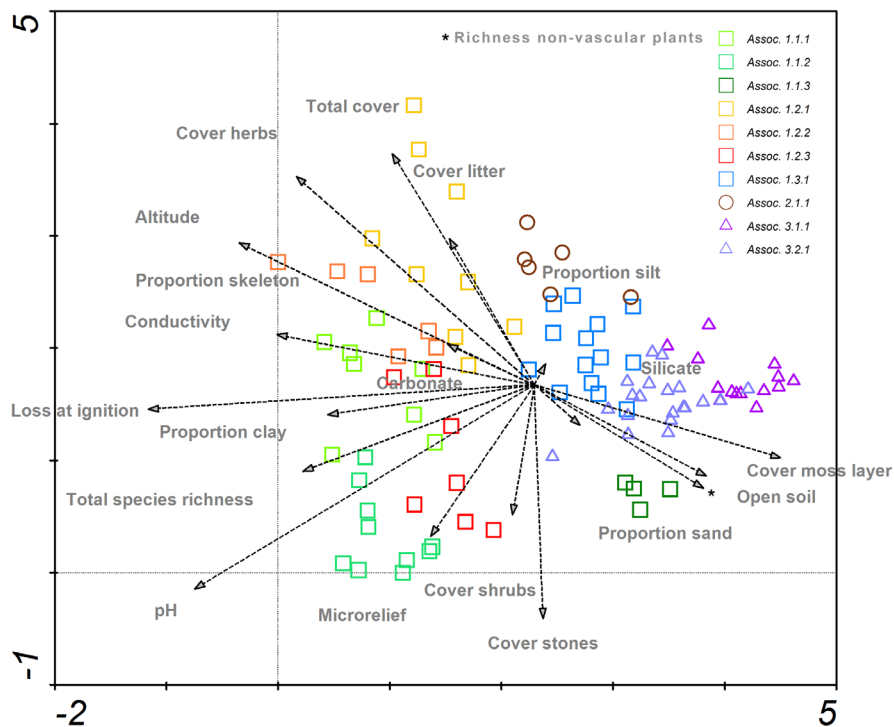
The first DCA axis was strongly negatively correlated with loss at ignition, proportion of clay and conductivity as well as geology, with carbonate bedrock having a positive loading (Fig. 10). This implies that soil properties were the main factors determining vegetation differences in the study region. Accordingly, the DCA like the TWINSpan tree (Fig. 3) separated two main vegetation types: (i) species-rich communities distributed on carbonate terrains, with base-rich soils having a high content of organic matter and a high clay fraction; (ii) less species-rich communities distributed on siliceous bedrock, with acid, sandy soils poor in organic matter. The second axis was strongly negatively correlated with cover of stones and positively correlated with total cover and cover of litter, indicating a gradient from open grasslands with many cryptogam species at rocky sites to dense grasslands with fewer cryptogam species on consolidated substrates.

Vegetation differences related to variations in ecological conditions are revealed by comparison of plant communities at class level (Table 7). *Festuco-Brometea* communities were found at the highest elevations in the study area (1108 m a.s.l.) on base-rich soils (mean pH = 6.2) rich in organic matter (19.4%), on steep slopes and rough microrelief. Horizontal structure of stands was open (82% vegetation cover), but litter cover was the highest among all classes (26%). These conditions appear to be most suitable to support high species richness (41.3 taxa on 10 m<sup>2</sup>). In contrast, *Koelerio-Corynephoretea* communities occurred at lower altitudes (910 m a.s.l.), on slightly acidic soils (pH = 5.0) poor in organic matter (7.6%). Total vegetation cover and species richness were the lowest among all studied classes (73% and 33%, respectively), while cover (mean 22%) and richness of lichens (3.5 species) as well as cover of open soil covers were the highest (11%). Most ecological parameters characteristic for the class *Calluno-Ulicetea* had intermediate values between those of the other two classes except for total vegetation cover, which was the highest, and cover of litter, which was the lowest.

### 5.2 Biodiversity patterns

With a mean total richness of 38.5 species on 10 m<sup>2</sup> (Table 4), the studied dry grasslands were intermediately species-rich at the European scale, where DENGLER (2005) listed association means ranging from 8.4 to 75.2 at that scale. Compared to *Festuco-Brometea* and *Koelerio-Corynephoretea* associations of NE Germany (DENGLER 2005), all associations of





**Fig. 10.** DCA ordination diagram of the 98 10-m<sup>2</sup> vegetation plots. The plots are coded according to the accepted classification (squares: *Festuco-Brometea*; circles: *Calluno-Ulicetea*; triangles: *Koelerio-Corynephoretea*). Rare species were down-weighted. Eigenvalues: 1st axis: 0.59; 2nd axis: 0.37.

**Abb. 10.** DCA-Ordinationsdiagramm der 98 Aufnahmeflächen von 10 m<sup>2</sup> Größe. Die Symbole der Plots entsprechend der angenommenen Klassifikation (Quadrate: *Festuco-Brometea*; Kreise: *Calluno-Ulicetea*; Dreiecke: *Koelerio-Corynephoretea*). Seltene Arten wurden für die Berechnung heruntergewichtet. Eigenwerte: 1. Achse: 0,59; 2. Achse: 0,37.

the respective classes contained on average 10 species more on that spatial scale, while their richness was similar to corresponding vegetation types of the Baltic islands of Saaremaa and Öland, respectively (BOCH & DENGLER 2006, LÖBEL & DENGLER 2008; Tables 7–9). In contrast, the corresponding communities (orders *Stipo pulcherrimae-Festucetalia pallentis* and *Brachypodietalia pinnati*) of Transylvania, only about 450 km away, had generally higher species richness than those of this study: 46.4 vs. 38.8 and 66.0 vs. 41.0, respectively (DENGLER et al. 2012a).

In Bulgaria *Festuco-Brometea* communities tended to be richer than those of the *Koelerio-Corynephoretea*, a typical pattern throughout Europe. However, with only about eight species (24%), the difference was smaller than in other regions, where the surplus is rather around 50% and more (e.g. HOBÖHM 1998, DENGLER 2005), and was significant only for the full dataset (Table 7), not for just the biodiversity plots (Table 9). Our single *Calluno-Ulicetea* association was as rich as the *Festuco-Brometea* stands (Tables 7–9), which is in line with HOBÖHM (1998: Anhang 12.5), who found that communities of the order *Nardetalia strictae* are among the richest vegetation types in Central Europe. Within the classes the

analysed associations generally showed only little differentiation in total richness, except for the *Satureja pilosa-Phleum phleoides* community (Assoc. 1.1.3), which was much poorer than the other *Festuco-Brometea* associations (Table 8).

Considering the three investigated taxonomic groups, all associations had much higher vascular plant than bryophyte and lichen species richness. Thus vascular plant richness closely corresponded to overall richness patterns. Bryophyte species richness on 10 m<sup>2</sup> was similar in all 10 associations with means of 2.0–3.2 species (Table 8). In contrast, lichen richness was distributed quite unevenly: *Koelerio-Corynepherea* stands showed higher values than the two other classes. In particular, the *Cetrario aculeatae-Plantaginietum subulatae* (Assoc. 3.1.1) had by far the highest values with 4.9 species per 10 m<sup>2</sup> (Table 8).

When analysing the seven spatial scales (Table 9), only few comparisons between the syntaxa were significant due to the low number of replicates. However, there was a non-significant tendency of *Festuco-Brometea* communities being richer in vascular plant species and all taxa than those of the two other classes, but only at the larger grain sizes. In contrast, at scales below 1 m<sup>2</sup>, the *Calluno-Ulicetea* and partly also the *Koelerio-Corynepherea* were richer in all taxa and vascular plants than *Festuco-Brometea*, albeit only insignificantly. This pattern was also reflected in the slope of the species-area-relationships, where the *z*-values of the *Festuco-Brometea* tended to be higher (mean: 0.253) than those of the other classes (0.210 and 0.218), although these differences were not significant. Higher *z*-values would indicate a faster increase of richness with area or, in other words, higher beta diversity. While the *z*-values in our study were within the typical range of dry grasslands across Europe (HOBOM 1998, DENGLER 2005), they were clearly lower than in Transylvanian dry grasslands (DENGLER et al. 2012a).

## 6. Conclusions and outlook

Considering that so far there is no comprehensive classification on the Bulgarian dry grasslands, our study is a valuable contribution to the knowledge of the variety of the dry grassland vegetation in the country and on the Balkan Peninsula in general. It provides an important input to the emerging efforts towards a consistent classification of grassland vegetation in Europe (DENGLER et al. 2013). While the overall number of plots was limited, it seems they are the first published from this region that thoroughly recorded not only vascular plants but also terricolous bryophytes and lichens. This aspect of sampling together with the different perspectives of phytosociologists from various countries outside Bulgaria in the expedition team allowed contributing to a clarification of the syntaxonomic position of Bulgarian dry grassland syntaxa in the European context.

We presented the first explicit documentation of the alliance *Violion caninae* and the order *Trifolio arvensis-Festucetalia ovinae* from Bulgaria. Further, we provided arguments to split up the heterogeneous Balkan dry grassland order *Astragalo-Potentilletalia* to place its content into the orders *Stipo pulcherrimae-Festucetalia pallentis* (*Saturejion montanae*), potentially *Festucetalia valesiaca*, *Brachypodietalia pinnati* (*Chrysopogono grylli-Danthonion calycinae*) and *Trifolio-Festucetalia* (most parts of the *Armerio rumelicae-Potentillion*). As these ideas are tentative at this stage, a much more comprehensive analysis of dry grassland syntaxa in Bulgaria as well as its neighbouring countries is needed for further clarification. Improving the knowledge about the mentioned vegetation types also con

**Table 9.** Scale-dependent richness values as well as slope parameters of the power-law species-area relationships ( $z$ ) (means  $\pm$  standard deviations) of the three phytosociological classes, based on the 15 “biodiversity plots”. Different letters indicate significant differences between communities at  $\alpha = 0.05$  from Turkey’s HSD test;  $P$ -values derived from ANOVAs.

**Tabelle 9.** Skalenabhängiger Artenreichtum sowie Steigungsparameter der mit dem Potenzgesetz modellierten Artenzahl-Areal-Beziehungen ( $z$ ) im Vergleich der drei pflanzensoziologischen Klassen, basierend auf 15 „Biodiversitätsplots“. Unterschiedliche Kleinbuchstaben bezeichnen signifikante Unterschiede ( $\alpha = 0,05$ ) gemäß Tukeys HSD-Test; in der letzten Spalte sind die  $P$ -Werte aus den Varianzanalysen angegeben.

Plot size	Min	Max	<i>Festuco-Brometea</i>	<i>Calluno-Ulicetea</i>	<i>Koelerio-Coryneporetea</i>	$P$
$n$			10	2	3	
Species richness of all taxa						
0.0001 m <sup>2</sup>	1	4	2.5 $\pm$ 0.9	3.3 $\pm$ 1.1	3.0 $\pm$ 0.5	0.395
0.001 m <sup>2</sup>	1	7	3.9 $\pm$ 1.8	6.0 $\pm$ 0.7	5.0 $\pm$ 1.3	0.233
0.01 m <sup>2</sup>	2	15	8.1 $\pm$ 3.9	11.8 $\pm$ 3.9	7.8 $\pm$ 2.8	0.441
0.1 m <sup>2</sup>	4	27	14.6 $\pm$ 6.6	17.8 $\pm$ 3.9	13.3 $\pm$ 2.5	0.711
1 m <sup>2</sup>	8	38	25.6 $\pm$ 9.0	27.0 $\pm$ 7.8	24.5 $\pm$ 5.4	0.949
10 m <sup>2</sup>	19	56	40.3 $\pm$ 11.3	41.3 $\pm$ 6.7	35.8 $\pm$ 10.3	0.802
100 m <sup>2</sup>	47	89	68.3 $\pm$ 14.5	60.0 $\pm$ 5.7	58.7 $\pm$ 7.6	0.466
Species richness of vascular plants						
0.0001 m <sup>2</sup>	1	4	2.1 $\pm$ 0.8	3.3 $\pm$ 1.1	2.3 $\pm$ 1.2	0.241
0.001 m <sup>2</sup>	1	7	3.3 $\pm$ 1.9	6.0 $\pm$ 0.7	4.3 $\pm$ 2.0	0.187
0.01 m <sup>2</sup>	2	15	7.2 $\pm$ 3.6	11.5 $\pm$ 3.5	6.7 $\pm$ 3.4	0.293
0.1 m <sup>2</sup>	4	22	13.3 $\pm$ 6.0	17.0 $\pm$ 3.5	11.2 $\pm$ 4.5	0.536
1 m <sup>2</sup>	8	33	23.3 $\pm$ 8.3	26.0 $\pm$ 7.8	19.0 $\pm$ 5.7	0.601
10 m <sup>2</sup>	16	55	36.5 $\pm$ 11.3	37.3 $\pm$ 6.7	27.2 $\pm$ 8.3	0.409
100 m <sup>2</sup>	38	87	61.2 $\pm$ 15.9	53.0 $\pm$ 4.2	44.3 $\pm$ 3.5	0.210
Species richness of bryophytes						
0.0001 m <sup>2</sup>	0	2	0.3 $\pm$ 0.5	0.0 $\pm$ 0.0	0.3 $\pm$ 0.3	0.704
0.001 m <sup>2</sup>	0	2	0.4 $\pm$ 0.5	0.0 $\pm$ 0.0	0.3 $\pm$ 0.3	0.639
0.01 m <sup>2</sup>	0	3	0.6 $\pm$ 0.8	0.3 $\pm$ 0.4	0.5 $\pm$ 0.5	0.878
0.1 m <sup>2</sup>	0	3	1.0 $\pm$ 1.0	0.8 $\pm$ 0.4	1.0 $\pm$ 0.9	0.951
1 m <sup>2</sup>	0	4	1.6 $\pm$ 1.3	1.0 $\pm$ 0.0	2.3 $\pm$ 0.8	0.453
10 m <sup>2</sup>	1	7	2.7 $\pm$ 2.0	3.8 $\pm$ 0.4	3.3 $\pm$ 0.8	0.684
100 m <sup>2</sup>	2	11	4.8 $\pm$ 2.7	6.5 $\pm$ 2.1	5.0 $\pm$ 1.0	0.678
Species richness of lichens						
0.0001 m <sup>2</sup>	0	1	0.1 $\pm$ 0.3	0.0 $\pm$ 0.0	0.2 $\pm$ 0.3	0.831
0.001 m <sup>2</sup>	0	2	0.2 $\pm$ 0.5	0.0 $\pm$ 0.0	0.2 $\pm$ 0.3	0.893
0.01 m <sup>2</sup>	0	3	0.3 $\pm$ 0.7	0.0 $\pm$ 0.0	0.5 $\pm$ 0.9	0.731
0.1 m <sup>2</sup>	0	2	0.4 $\pm$ 0.7	0.0 $\pm$ 0.0	0.8 $\pm$ 0.8	0.440
1 m <sup>2</sup>	0	4	0.6 $\pm$ 0.8 <sup>a</sup>	0.0 $\pm$ 0.0 <sup>a</sup>	2.8 $\pm$ 0.8 <sup>b</sup>	0.002
10 m <sup>2</sup>	0	6	0.9 $\pm$ 1.2 <sup>a</sup>	0.3 $\pm$ 0.4 <sup>a</sup>	4.5 $\pm$ 1.3 <sup>b</sup>	0.001
100 m <sup>2</sup>	0	11	1.8 $\pm$ 2.1 <sup>a</sup>	0.5 $\pm$ 0.7 <sup>a</sup>	7.7 $\pm$ 2.9 <sup>b</sup>	0.004
Species-area relationship for all taxa						
$z$ -value	0.196	0.341	0.253 $\pm$ 0.042	0.210 $\pm$ 0.019	0.218 $\pm$ 0.008	0.201



tributes to a better understanding of habitat diversity. This particularly applies to species-rich *Nardus* grasslands (listed as habitat 6230\* in Annex I of the Habitats Directive) and to eastern sub-Mediterranean dry grasslands (62A0).

Finally, this study was probably the first on the Balkan Peninsula thoroughly recording diversity patterns of dry grassland vegetation across spatial scales and taxa. In many respects the results matched those in other dry grasslands of Europe. However, while most types of dry grasslands are among the richest plant communities of any region in Europe (DENGLER 2005, WILSON et al. 2012), one big question remains unsolved at present: Why are the dry grasslands of a few narrow regions in Europe, such as Transylvania, the White Carpathians and the hemiboreal alvar sites, so much richer than all others including those in the study area, though they seem to be similar in site conditions and species pool size? Despite first explanatory approaches (e.g. HÁJKOVÁ et al. 2011, DENGLER 2012, DENGLER et al. 2012a, MERUNKOVÁ et al. 2012), we are still far from truly understanding this astonishing pattern. The best chance to solve this riddle is the compilation of standardised high-quality richness data together with information on as many predictors across many different regions in Europe as possible and their use in a large-scale meta-analysis.

## Erweiterte deutsche Zusammenfassung

**Einleitung** – Die Balkanhalbinsel ist innerhalb Europas überdurchschnittlich reich an Pflanzenarten und Endemiten, während zugleich halbnatürliche Grasländer von hohem Naturschutzwert dort noch großflächig erhalten sind. Bulgarien hinkt, was die pflanzensoziologische Inventarisierung angeht, den meisten anderen europäischen Ländern hinterher, da dort bis zum Fall des Eisernen Vorhangs die wenigen Vegetationskundler nahezu ausschließlich der russischen Schule folgten. Seither wurden zahlreiche pflanzensoziologische Studien durchgeführt, was zu einer besonders in den Graslandklassen gut mit Aufnahmen bestückten nationalen Vegetationsdatenbank führte (APOSTOLOVA et al. 2012). Allerdings fehlten bislang überregionale Analysen, die auf die Eingliederung der bulgarischen Einheiten in ein europaweites syntaxonomisches System abzielten. Europäische Trockenrasen sind ein besonders interessantes Studienobjekt für Biodiversitätsanalysen, unter anderem, da sie auf Flächengrößen unter 100 m<sup>2</sup> die globalen Rekorde im Pflanzenartenreichtum halten (DENGLER 2005, 2012, WILSON et al. 2012). Zum Verständnis der Diversitätsmuster in paläarktischen Trockenrasen und der sie steuernden Faktoren sowie als Beitrag zu einer konsistenten supranationalen Klassifikation derselben führt die *European Dry Grassland Group* (<http://www.edgg.org>) seit 2009 jährliche Forschungsexpeditionen in noch unzureichend erforschte Regionen durch, um dort Grundlagendaten mit einer standardisierten Erfassungsmethode zu erheben (vgl. DENGLER et al. 2012). Die vierte derartige Expedition führte im Sommer 2012 nach Westbulgarien mit den folgenden Hauptzielen: (1) Abgrenzung der vorkommenden Typen von Trockenrasen und Ermittlung ihrer Kennarten mit statistischen Methoden; (2) Einordnung dieser Einheiten in das europäische syntaxonomische System; (3) Grundverständnis der ökologischen Faktoren, die zur Differenzierung der Einheiten führen; (4) Dokumentation der Diversitätsmuster von Gefäßpflanzen, Moosen und Flechten auf verschiedenen räumlichen Skalenebenen.

**Untersuchungsgebiete** – Für die Expedition wählten wir zwei zuvor kaum untersuchte Regionen in Nordwestbulgarien aus, Vratsa im westlichen Teil des Balkangebirges (Stara planina) und Koprivshitsa im Tal zwischen Stara planina und den südlich angrenzenden Mittelgebirgszug Sredna gora, einschließlich der niederen Lagen beider Höhenzüge (Abb. 1). Die beiden Regionen liegen größtenteils in *Sites of Community Interest* des Natura 2000-Netzes und beinhalten großflächig extensiv genutzte, meist trockene Grasländer (Abb. 2). Sie unterscheiden sich deutlich in ihren physisch-geografischen Voraussetzungen (Tab. 1), Vratsa mit Kalkgestein liegt auf 970–1400 m Meereshöhe und Koprivshitsa mit Silikatgestein auf 630–1200 m Meereshöhe.

**Methoden** – Wir analysierten eine möglichst große Bandbreite der in den beiden Untersuchungsgebieten vorkommenden Trockenrasentypen mit zwei komplementären Erhebungsmethoden: (a) zum einen erfassten wir sogenannte Biodiversitätsplots ( $n = 15$ ) von 100 m<sup>2</sup> Größe, in die je zwei Subplots der Größen 0,0001, 0,001, 0,01, 0,1, 1 und 10 m<sup>2</sup> in zwei gegenüberliegenden Ecken geschachtelt waren (vgl. DENGLER 2009b), (b) zum anderen fertigten wir zusätzliche normale Vegetationsaufnahmen auf 10 m<sup>2</sup> großen Flächen an ( $n = 68$ ). Auf sämtlichen Aufnahmeflächen beider Erfassungsmethoden wurden sämtliche epigäische Gefäßpflanzen, Moose und Flechten erfasst, auf den 10 m<sup>2</sup> großen Flächen (insgesamt  $n = 98$ ) zusätzlich die prozentuale Deckung, topografische, Boden-, und Landnutzungsvariablen. Für die Vegetationsklassifikation wurden die 10 m<sup>2</sup> großen Flächen mit der modifizierten TWINSPAN-Methode (ROLEČEK et al. 2009) gegliedert. Für die erhaltenen Endcluster wurden diagnostische Arten mittels phi-Werten in Kombination mit Fishers exaktem Test (CHYTRÝ et al. 2002) nach Standardisierung der Aufnahmezahl auf 1/10 ermittelt, für die übergeordneten Einheiten nach den darauf aufbauenden Prinzipien von LUTHER-MOSEBACH et al. (2012). Eine DCA mit passiv hinein projizierten Umweltvariablen diente der Visualisierung der Anordnung der Vegetationseinheiten entlang von floristischen Hauptgradienten. Unterschiede in den Umweltbedingungen und Biodiversitätsparametern zwischen den Vegetationseinheiten analysierten wir mittels Varianzanalyse.

**Vegetationsklassifikation** – Die 10-Cluster-Lösung von TWINSPAN war die feinste Auflösung, die noch floristisch klar unterscheidbare Einheiten erbrachte und wurde daher als Assoziationsebene angenommen. Während wir die Zuordnung der Aufnahmen zu den Endclustern unverändert ließen, modifizierten wir die Anordnung dieser Einheiten in der Hierarchie geringfügig (Abb. 3), um unsere regionalen Ergebnisse in das existente und gut belegte überregionale Klassifikationssystem besser einpassen zu können. Entsprechend konnten wir drei Vegetationsklassen mit Trockenrasen im Gebiet nachweisen. Die Klasse *Festuco-Brometea* ist vertreten mit den Felstrockenrasen (*Stipo pulcherrimae-Festucetalia pallentis*) und den Kalk-Halbtrockenrasen (*Brachypodietalia pinnati*). Zur ersten Ordnung stellen wir den balkanischen Verband *Saturejion montanae*, der bislang verschiedenen anderen Ordnungen untergeordnet war, und der in den Untersuchungsgebieten mit drei assoziationsgleichen Einheiten vertreten ist, der *Dianthus petraeus-Sesleria latifolia*-Gesellschaft (Abb. 4), der *Achillea clypeolata-Festuca stojanovii*-Gesellschaft und der azidoklinen *Satureja pilosa-Phleum phleoides*-Gesellschaft. Die *Brachypodietalia pinnati* umfassen einerseits den Verband *Cirsio-Brachypodium pinnati* mit der *Ferulago campestris-Agrostis capillaris*-Gesellschaft, der *Tanacetum corymbosum-Sesleria latifolia*-Gesellschaft (Abb. 5) und der *Abietinella abietina-Sanguisorba minor*-Gesellschaft, andererseits den azidoklinen Verband *Chrysopogono grylli-Danthonion calycinae* mit dem *Agrostio capillaris-Chrysopogonetum grylli* (Abb. 6). Aus der Klasse *Calluno-Ulicetea* konnte erstmalig der Verband *Violion caninae* (Ordnung *Nardetalia strictae*) für Bulgarien nachgewiesen werden, vertreten durch die zuvor nur aus dem südlichen Mitteleuropa bekannte Assoziation *Festuco rubrae-Genistelletum sagittalis* (Abb. 7). Die Klasse *Koelerio-Corynephoretea* war zuvor so gut wie nicht aus Bulgarien dokumentiert worden, so dass wir die beiden von uns gefundenen, in der Region von Kovprivishtitsa weit verbreiteten Assoziationen neu beschreiben mussten. Das *Cetrario aculeatae-Plantaginetum subulatae* ass. nov. (Abb. 8) ist eine offene, kryptogamenreiche Felsgrusflur saurer Gesteine und gehört wohl zu einem noch unbeschriebenen Verband der Ordnung *Sedo-Scleranthetalia*. Das *Plantagini subulatae-Agrostietum capillaris* ass. nov. (Abb. 9) ist dagegen ein Sandhalbtrockenrasen mit meist recht geschlossener Grasnarbe, der sich zwanglos in den zuvor beschriebenen Verband *Armerio rumelicarum-Potentillion* einordnen lässt, welcher aber entgegen der üblichen Praxis in der Region nicht in die Ordnung *Astragalo-Potentilletalia* (*Festuco-Brometea*), sondern die Ordnung *Trifolio arvensis-Festucetalia ovinae* (*Koelerio-Corynephoretea*) gestellt werden sollte. Alle Assoziationen bzw. assoziationsgleichen Einheiten werden vegetationsökologisch charakterisiert, ihre Ökologie und Verbreitung beschrieben und ihre Klassifikation im regionalen wie überregionalen Kontext diskutiert. Im Anhang S1 werden nomenklatorische Probleme erörtert und ggf. Typisierungen vorgenommen.

**Ökologie und Biodiversität** – Die DCA (Abb. 10) trennte entlang der ersten Achse (a) artenreiche Gesellschaften auf humus- und tonreichen Böden der basenreichen Gesteine des Gebiets um Vratsa von (b) relativ artenarmen Gesellschaften auf humus- und tonarmen Böden saurer Ausgangsgesteine um

Kovprivshitsa. Entsprechend unterscheiden sich die Aufnahmen der beiden Klassen *Festuco-Brometea* und *Koelerio-Corynephoretea* signifikant in ihren Umweltbedingungen- und in ihrer Biodiversität, während die *Calluno-Ulicetea* meist eine intermediäre Stellung einnehmen (Tab. 7). Mit 47–89 Arten auf 100 m<sup>2</sup> sind die untersuchten Bestände artenreich, aber erreichen nicht Extremwerte wie etwa in Siebenbürgen oder den Weißen Karpaten. Mit einem durchschnittlichen  $z$ -Wert von 0.25 ist zumindest in den *Festuco-Brometea*-Gesellschaften auch die  $\beta$ -Diversität ungewöhnlich hoch (Tab. 9).

**Resümee und Ausblick** – Der gemeinsame Blick von Trockenrasenspezialisten aus Bulgarien und verschiedenen anderen europäischen Ländern half im Rahmen der EDGG-Expedition, die vielfältigen Trockenrasentypen Nordwestbulgariens besser in ein überregionales System einzuordnen und leistet damit einen wichtigen Beitrag zu einer konsistenten europaweiten Klassifikation. Die standardisierte Erfassung der Biodiversitätsmuster von Gefäßpflanzen, Moosen und Flechten auf verschiedenen räumlichen Skalenebenen lieferte wertvolle Grundlagendaten (vor allem bei den Nicht-Gefäßpflanzen) und stellt einen wichtigen Beitrag zum Verständnis der treibenden Kräfte der Biodiversitätsmuster paläarktischer Trockenrasen dar.

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### Supplements and Appendices

**Supplement 1.** Table 4. Abridged synoptic table of the dry grassland communities in the study region in NW Bulgaria.

**Beilage 1.** Tabelle 4. Gekürzte Stetigkeitstabelle der analysierten Trockenrasengesellschaften Nordwest-Bulgariens.

**Supplement 2.** Table 5. Relevé table of the *Festuco-Brometea* communities in the study region in NW Bulgaria.

**Beilage 2.** Tabelle 5. Aufnahmetabelle der untersuchten *Festuco-Brometea*-Gesellschaften Nordwest-Bulgariens.

**Supplement 3.** Table 6. Relevé table of the *Calluno-Ulicetea* and *Koelerio-Corynephoretea* communities in the study region in NW Bulgaria.

**Beilage 3.** Tabelle 6. Relevé table of the *Calluno-Ulicetea* and *Koelerio-Corynephoretea* communities in the study region in NW Bulgaria.

**Appendix 1.** New descriptions and typifications of syntaxa.

The references to all given authorities are included in the reference list.

**Anhang 1.** Neubeschreibungen und Typisierungen von Syntaxa.

The Quellen zu allen Autoren von Syntaxa sind im Quellenverzeichnis aufgeführt.

***Cetrario aculeatae-Plantaginetum subulatae* Pedashenko et al. ass. nov. hoc loco**

Type: Relevé BGR039 in Table 6 of this publication [Holotypus hoc loco]

***Plantagini subulatae-Agrostietum capillaris* Pedashenko et al. ass. nov. hoc loco**

Type: Relevé BGR056 in Table 6 of this publication [Holotypus hoc loco]

***Tuberario guttatae-Corynephoretea* Hohenester 1967**

Type: *Corynephoretalia* Tx. ex Hohenester 1967 [Lectotypus hoc loco]

***Armerio rumelicarum-Potentillion* Micevski 1978**

Protologue: “*Armerio-Potentillion*” MICEVSKI (1978: pp. 21 et seq.)

Type: *Genisto carinalis-Agrostietum byzanthinae* Micevski 1978\* [Lectotypus hoc loco]

***Saturejion montanae* Horvat et al. 1974**

Protologue: “*Saturejion montanae* Horvat 62” (HORVAT et al. 1974: p. 264)

Type: *Carici humilis-Stipetum grafianae* Jovanović-Dunjić 1955 [Lectotypus hoc loco]

***Agrostio capillaris-Chrysopogonetum grylli* Kojić 1959**

Protologue: “Asocijacija *Agrostideto-Chrysopogonetum grylli* (As. *Agrostis capillaris-Chrysopogon gryllus*)” (KOJIĆ 1959: pp. 12–30)

Type: KOJIĆ (1959: Table 9, relevé 6) [Lectotypus hoc loco]

***Carici humilis-Stipetum grafianae* Jovanović-Dunjić 1955**

Protologue: “*Humileto-Stipetum grafianae*” (JOVANOVIĆ-DUNJIĆ 1955)

Type: JOVANOVIĆ-DUNJIĆ (1955: Table 4, relevé 5) [Lectotypus hoc loco]

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**Zusätzliche unterstützende Information ist in der Online-Version dieses Artikels zu finden.**

**Appendix S1.** Overview and nomenclatural revision of the syntaxa discussed.

**Anhang S1.** Übersicht und nomenklatorische Revision der diskutierten Syntaxa.

**Appendix S2.** Origin of the relevés

**Anhang S2.** Herkunftsnachweis der Aufnahmen

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