Alleviation of Plant Stress Precedes Termination of Rich Fen Stages in Peat Profiles of Lowland Mires

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ABSTRACT

Mesotrophic rich fens, that is, groundwater-fed mires, may be long-lasting, as well as transient ecosystems, displaced in time by poor fens, bogs, forests or eutrophic reeds. We hypothesized that fen stability is controlled by plant stress caused by waterlogging with calcium-rich and nutrient-poor groundwater, which limits expansion of hummock mosses, tussock sedges and trees. We analysed 32 European Holocene macrofossil profiles of rich fens using plant functional traits (PFTs) which indicate the level of plant stress in the environment: canopy height, clonal spread, diaspore mass, specific leaf area, leaf dry matter content, Ellenberg moisture value, hummock-forming ability, mycorrhizal status and plant functional groups. Six PFTs, which formed long-term significant trends during mire development, were compiled as rich fen stress indicator (RFSI). We found that RFSI values at the start of fen development were correlated with the thickness of subsequently accumulated rich fen peat. RFSI declined in fens approaching change into another mire type, regardless whether it was shifting into bog, forest or eutrophic reeds. RFSI remained comparatively high and stable in three rich fens, which have not terminated naturally until present times. By applying PFT analysis to macrofossil data, we demonstrated that fens may undergo a gradual autogenic process, which lowers the ecosystem’s resistance and enhances shifts to other mire types. Long-lasting rich fens, documented by deep peat deposits, are rare. Because autogenic processes tend to alleviate stress in fens, high levels of stress are needed at initial stages of rich fen development to enable its long persistence and continuous peat accumulation.

Key words: ecophysiological stress; ecosystem engineers; ecosystem shift; plant macrofossils; peat deposits; rich fen vegetation succession; tree and shrub encroachment; waterlogging.

HIGHLIGHTS

- Rich fen stability is controlled by plant stress caused by waterlogging.
• Rich fens with initially higher levels of plant stress produce thicker layers of peat.
• Plant functional traits indicate that stress level declines in rich fens approaching shift to other mire types.

INTRODUCTION

Rich fens are mires (that is, peat-forming wetlands) supplied with groundwater enriched with base ions (Joosten and others 2017). Although the concept of rich fens may include mires from a range of trophic conditions (Wheeler and Proctor 2000), many authors limit it to mesotrophic ecosystems, that host low-productive, nutrient-limited vegetation dominated by low sedges and brown mosses (compare Hájek and others 2006 and references therein). Low productivity is a natural feature of mires, in which part of primary production becomes buried in peat, whereas nutrient influx from outside is absent or very limited. In the present study, we focus on this mesotrophic wing of rich fens, aiming to understand the mechanisms of their stability and shifts to other mire types. Mesotrophic rich fens developing during the Holocene were in some sites displaced in course of succession by bogs or poor fens (for example, Lamentowicz and others 2008), alder forests (for example, Marek 1965) or reed beds or aquatic systems dominated by submerged macrophytes (for example, Michaelis 2002). In some places, rich fens persisted for millennia and until the present (for example, Michaelis 2002; Jabłońska and others 2014; Hájková and others 2015). They are, however, increasingly vanishing, due to large-scale drainage for agriculture and forestry, and regional hydrological alterations (van Diggelen and others 2006). In result, many specialist fen species are on the verge of extinction (for example, Bedford and Godwin 2003; Rehell and Virtanen 2016), while fens have turned from carbon-accumulating systems into significant sources of carbon to the atmosphere (Kotowski and others 2016). To avoid further losses, external disturbing forces need to be eliminated in remaining rich fen systems, whereas the drained ones should be restored, preferably back into stable peat-forming mires (Kotowski and others 2016). However, despite the fact that rich fens are in the focus of conservation and restoration ecologists (see Lamers and others 2015) and Kotowski and others 2016 for references), the effectiveness of their restoration is limited (Klimkowska and others 2019). To learn how to restore degraded rich fens into stable and functional systems, we need to better understand preconditions of stability and instability (ecosystem shifts) in the natural ones.

Ecosystem shifts in mires may be extrinsic (typically associated with allogenic influences, for example, climate change or anthropogenic alteration of catchment hydrology, for example, Hájková and others 2012; Jamrichová and others 2014) or intrinsic (caused by autogenic factors, that is, related to vegetation succession, for example, progressive accumulation of peat and raising of the plant root zone above the influence of groundwater, for example Kulczyński 1949, Zoibel 1988, Williams and others 2011). Autogenic factors are in fact ‘ecosystem engineer’ species (Jones and others 1994) with traits which may drive intrinsic shifts in mires through positive feedbacks (Waddington and others 2015). Allogenic and autogenic factors may also interact, when ecosystem engineers respond to extrinsic changes, thereby amplifying the shift (Singer and others 1996; Magyari and others 2001; Granath and others 2010; Morris and others 2015).

One example of ecosystem engineers is hummock-forming bryophytes. Following acidification of a rich fen (for example, due to partial displacement of groundwater by rainwater), they may foster a further drop of pH due to active cation exchange (Clymo 1966) for sphagna, Soudzilovskajaia and others 2010 for brown mosses), rainwater retention within hummocks (Vicherová and others 2017) and release of organic acids (van Breemen 1995; Siegel and others 2006). In effect, hummock mosses (which are intolerant to calcium, compare Vicherová and others 2015) are benefited over hollow mosses. In addition, a further expanse of sphagna may be enhanced by capturing CO₂ released from bicarbonates in water below the hummocks due to acidification (Harpenslager and others 2015) and their effective nitrogen fixation (Fritz and others 2014).

Another group of mire ecosystem engineers are trees and shrubs, which may lower water saturation in surface peat layers due to enhanced evapotranspiration (Grygoruk and others 2014), thereby facilitating establishment of new trees. Due to the lowered water saturation in surface peat, more oxygen is available for plant roots as well as ectomycorrhizal fungi. Decomposing microorganisms benefit from oxygen availability, enhancing peat mineralization and nutrient availability, which may further stimulate trees growth (Succow and Joosten 2001). Also, tussock-forming sedges have been considered ecosystem engineers (Peach and Zedler 2006) because they can be used as dry
microhabitats by non-wetland species, including
phanerophytes and their mycorrhizal associates.

Ecosystem engineers can be kept in check by
stress factors [Grime 1977; also named habitat filters
(Keddy 2017)], to which they are intolerant. In rich
fens, waterlogging of peat with base-rich ground-
water induces plant stress in several ways. Firstly,
reduced forms of calcium (Ca^{2+}), iron (Fe^{2+}),
ammonia (NH_4^+) or manganese (Mn^{2+}) are toxic for
sphagna, in contrast to calcium-tolerant rich fen
mosses, such as Hamatocaulis vernicosus and Scorpid-
ium scorioides (Vicherová and others 2015). Sec-
ondly, waterlogging limits mineralization of organic
matter, causing nutrient limitation stress. Thirdly,
anoxia of the root zone keeps out non-aerenchymic
plants and obligatorily mycorrhizal species (Peat
and Fitter 1993; Hempel and others 2013), includ-
ing trees, thereby disabling a shift into forests. Field
observations from Rospuda Fen in Poland show that
a rich fen with high level of the mentioned stresses
caused by very stable water table can persist for
millennia avoiding shifts to other mire types, in
contrast to a dynamically changing neighbouring
system with less stable hydrological conditions
(Jabłońska and others 2014). Yet, because of rarity
of undisturbed fen systems, it is difficult to assess
whether the control of rich fen stability by stress
factors is a general rule. In the present study, we
aim to assess this based on palaeoecological data.

We hypothesize that alleviation of plant stress
preceded past shifts of rich fens into other mire
types. To check this hypothesis, we analysed plant
functional traits (PFTs) spectra of subfossil plant
communities, dating back long before humans
significantly impacted the landscape. We chose
PFTs which may indicate the intensity of the fen
stressors described above (compare Table 1). Fur-
ther, based on the palaeoecological evidence, we
propose a PFT-based indicator that can help
reconstruct ecosystem’s level of plant stress (Dou-
ma and others 2012). This approach allows com-
parisons of systems with a different response to the
stress and may help to predict ecosystem shifts in
contemporary mires. An establishment of such
tools forecasting ecosystem shifts has been recog-
nized as a challenge of global change ecology
(Williams and others 2011).

Methods
Collection of Rich Fen Macrofossil
Profiles From Literature

We compiled a database of 52 peat profiles with
rich fen species present and bryophytes determined
(Table 1 in Appendix 1, map in Appendix 1, Ap-
pendix 2) and digitized all data from tables and
diagrams. A single vegetation sample in this data-
base corresponds to one peat sample analysed for
macrofossils and individually displayed on a
macrofossil diagram/table. The interval between
samples in a peat core varied between studies from
5 to 30 cm. The median interval was 10 cm. Plant
macrofossils were described with taxon names of
the lowest taxonomic rank available. When hier-
archic inclusive taxa were listed in a vegetation
sample, for example Carex sp. and Carex rostrata, we
treated them as separate taxa, assuming that a
macrofossil not recognized as a separate species
identifiable in a profile belongs potentially to an-
other species. Taxa were classified as either present
non-dominant or dominants. Dominants were
those taxa, the amount of which (number of countable
remains/volume) clearly exceeded the amount of each other taxon in a sample. Zero to
five dominants were chosen per sample.

All digitized samples of subfossil plant commu-
nities were inspected for meeting jointly three cri-
teria of rich fen, that is, (1) presence of at least one
of rich fen moss species, (2) non-dominance of
trees, bog/poor fen specialists, reed and aquatic
plants, and (3) the number of taxa mentioned in #2
not exceeding the number of fen taxa (see Ap-
pendix 1 for details). Thirty-nine profiles met these
criteria with at least one vegetation sample. In a
preliminary analysis, we found that this data set
was biased due to altitude (Appendix 3) and re-
moved upland samples (7 profiles) from further
analysis. This allowed for a better homogenization
of the data set and elimination of additional impact
of climate variation due to altitude. The resulting
data set contains 32 profiles with rich fen sections
from lowland mires (located between 0 and 300 m
a.s.l.) (Table 1 in Appendix 1). In 13 lowland pro-
files for which radiocarbon dating was provided
within rich fen sections, we checked the relation-
ship between age and thickness of these sections
and found a significant positive linear correlation.
Upland fens were outliers from this relation-
ship—compare Appendix 3.

Plant Functional Traits

We included 13 PFTs, which may indicate the
intensity of anoxia and/or nutrient stresses (Ta-
ble 1, extended version in Appendix 4). The rate of
yearly clonal spread (CS), that is, the increment of
clonal growth organ in the horizontal direction,
was converted into two binary traits, CS small and
CS large, as suggested by Weiher and others (1999).
| Trait | Sources | Hypothesized response to fen stress conditions |
|-------|---------|-----------------------------------------------|
| BM Brown mosses (non-Sphagnum mosses) | Own assessment | ↑<sub>AN</sub> main peat-formers, indicators of low competition for light (<i>high N</i> stress) and stable water level of groundwater origin, can tolerate calcium |
| SPH Sphagna | Own assessment | ↓<sub>A</sub> may increase with lowering groundwater feeding, indicating decrease in <i>A</i> and decrease in calcium toxicity |
| PHA Phanerophytes (trees and shrubs) | Own assessment | ↓<sub>A</sub> impeded in mires by <i>A</i>, presumably due to suppressed ectomycorrhiza; may further trigger drainage due to increased evapotranspiration |
| HF Hummock/tussock-forming species | Own classification, see Appendix 4 | ↓<sub>A</sub> hummocks offer drier safe sites for non-wetland plants, may enhance establishment of trees and other competitive species |
| EL Ellenberg moisture value F | Ellenberg and others (1991) | ↑<sub>A</sub> high EL indicates the presence of adaptations to <i>A</i> and flooding, decreases with intensifying drainage |
| CH Canopy height (m) | TRY<sup>a</sup>; Rutkowski (2012) | ↓<sub>N</sub> high CH indicates high competition for light (<i>low N</i> stress); high CH may also be attributed to plant reaction to flooding, as emergent species grow tall to stay above the water level, however, only if nutrients are sufficiently available |
| CS Yearly clonal spread present but < 1 cm | CLOPLA (Klimešová and de Bello 2009) | ↓<sub>A</sub> in wetlands species with CS small create tussocks or swards, providing stable shelters elevated above the groundwater table, which may enhance establishment of trees and other species intolerant to <i>A</i> |
| CS Yearly clonal spread > 1 cm | | ↑<sub>A</sub> CS large is frequent in wetland dominants, vegetative propagation is favoured over generative reproduction under <i>A</i>; CS large may indicate extremely productive sites where vegetative propagation is the only possible because of light deficiency for germination |
| SM Diaspore mass (mg) | TRY<sup>a</sup>; own assessment | ↓<sub>N</sub> indicates competition for light: large-seeded plants better germinate in dark and produce larger seedlings; within rich lens may decrease with growing <i>N</i> stress (excluding extremely productive sites dominated by long-distance colonizers, for example, <i>Typha</i>, <i>Phragmites</i>) |
| SLA Specific leaf area (mm<sup>2</sup> mg<sup>-1</sup>) | TRY<sup>a</sup> | ↓<sub>AN</sub> species with high SLA have high photosynthesis investments and are often ruderals or competitors but rarely stress tolerators; in wetlands SLA decreases with increasing wetness and low SLA may be an adaptation to avoid uptake of toxic forms of reduced minerals via reduced evapotranspiration |
| LDMC Leaf dry matter content (g g<sup>-1</sup>) | LEDA (Kleyer and others 2008) | ↑<sub>AN</sub> high LDMC correspond with conservative nutrient strategy indicating both <i>N</i> stress and <i>A</i> |
| Ob-M Obligatory mycorrhizal species | MycoFlor (Hempel and others 2013), Akhmetzhanova and others (2012), Veselkin, and others (2014) | ↓<sub>N</sub>↑<sub>A</sub> Ob-M may be attributed to the presence of <i>N</i> stress, impeded by <i>A</i> |
| Non-M Non-mycorrhizal species | | ↑<sub>A</sub> indicators of wet conditions (<i>A</i>) |

Arrows indicate positive (↑) or negative (↓) hypothesized relationship with anoxia (<i>A</i>) and nutrient stress (<i>N</i>). (See Appendix 4 for an extended version of the table.)

<sup>a</sup>TRY Global Plant Traits Database (Kattge and others (2011a); main data sources: Cornelissen and others (1996, 2003, 2004), Ordonez and others (2010), Kattge and others (2011b), Dainese and Bragazza (2012), Royal Botanical Gardens KEW; a list of further data sources, from which in total less than 5% of data were derived, is provided in Appendix 4.)
Two analysed types of mycorrhizal associations (Ob-M and Non-M) were defined, according to Hempel and others (2013) and Moora (2014). Each mycorrhizal type was treated as a binary trait; facultative mycorrhizal species were given 0 in both categories.

In the case of the identification of taxa at a taxonomical level higher than species (for example, Carex sp., Pteridophyta) or species with missing trait values in the mentioned sources, trait values of taxonomical relatives growing also in mires were assigned based on expert knowledge. Taxa identified to the family level were discarded from the analysis of the mycorrhizal status, with the exception of Ericaceae, which were classified as Ob-M (for example, Hempel and others (2013)).

**Data Analysis**

All the 32 profiles with rich fen sections from lowland mires were analysed for the thickness of rich fen sections and the direction of their changes. Further, we calculated community-weighted means (CWM) of traits for all vegetation samples, giving a weight 10 for the dominant taxa and 1 for the remaining taxa.

In a subset of 12 profiles, rich fen sections were at least 1 m long, allowing for regression analyses of PFT changes. To explore whether CWMS of stress-related PFTs within the rich fen phases formed directional trends leading to—or indicating—a shift in mire vegetation, we constructed mixed effect models including nine profiles in which rich fen phase finished in the past. (Depth of rich fen end was at least 90 cm—Table 1 in Appendix 1.) We used the distance (measured in cm) of a vegetation sample to the position of a vegetation shift in the peat profile (that is, the first non-rich fen sample above a rich fen section within a peat core) as an independent variable. The variable, further referred to as ‘distance to rich fen finish’, has negative values for samples lying in the peat core below the shift and positive values for samples above the shift; the first non-rich fen sample above the rich fen section within a peat core has value zero. Vegetation samples were grouped by their profile, and a model with a random intercept and a slope was used. We used the Gaussian distribution for quantitative traits (CH, SLA, SM and EL) and binomial distribution for the traits with CWMS within a 0–1 range (SPH, PHA, HF, CS small, CS large, LDMC, Ob-M and Non-M). All models were calculated using glmmPQL function in MASS package in R (Venables and Ripley 2002). The traits, which were significantly related to the distance to rich fen finish according to the above-described models, were then aggregated as a rich fen stress indicator (RFSI), and the relation of RFSI itself to the distance to rich fen finish was checked. Although RFSI values were within 0–1 range, the relationship between RFSI and the distance to the rich fen end was analysed using glmm with the Gaussian distribution and a simple linear regression, because RFSI values were between 0.3 and 0.8 and thus located in the middle (linear) section of the sigmoidal distribution curve. Finally, we calculated also the relation of RFSI to the distance to rich fen finish for three rich fen mires that have persisted to the present or terminated due to a recent human impact; in the latter case, we excluded the uppermost samples of degraded peat from further analysis.

In a subset of 20 rich fen sections, including at least three samples classified as rich fen, needed to calculate a median value, forming a 30–50-cm-long continuous section, we checked the relationship between median RFSI of this lowermost section and the length of the whole rich fen section, expressed as a difference between the first non-rich fen sample above the rich fen finish in a profile and the uppermost vegetation sample in the lowermost 30–50 cm section (‘distance to rich fen finish’).

**RESULTS**

The analysis of 32 profiles showed that most peat profiles included only one rich fen section and the length of the longest rich fen section in a profile, as well as the summary length of rich fen sections, rarely exceeded 200 cm (Figure 1 in Appendix 5). The majority of rich fen sections did not exceed 50 cm. When shifts occurred, they led equally often towards bogs, forests and reed beds. The majority of rich fen sections start not deeper than 300 cm; only a few rich fen deposits begin at deeper layers (Figure 1).

Out of nine profiles analysed for PFT trends, three changed into bogs or poor fens, another three into fen woodlands and the remaining three into reeds or tall sedge beds. The analysis of PFTs in nine profiles revealed that CWMS values for HF, CS small and CH increased and CWM values for Non-M decreased during rich fen development (Table 2). By analysing rich fen time series together with the peat samples directly following a shift, we found increasing CWM values for PHA and decreasing CWM values for BM.

We assumed that the shares of Non-M and BM (both CWMS decreasing during mire development) are positively related to the level of ecophysiological stress and thus positively related to rich fen
stability, whereas the shares of HF, CS small, PHA and CH (all CWMs increasing during mire development) are negatively related to the level of stress and, similarly, negatively related to rich fen stability (compare Table 1). On the basis of this assumption, we calculated an additive rich fen stress indicator (RFSI) for each vegetation sample (the higher the RFSI value, the more stressed and thus more stable the fen) (equation 1):

\[
RFSI = \left( \frac{CWM_{Non-M} + CWM_{BM} + (1 - CWM_{HF})}{C0_{CWM}} \right) + \left( \frac{1 - CWM_{CS small}}{C0_{CWM}} \right) + \left( \frac{1 - CWM_{PHA}}{6} \right)
\]

To obtain RFSI values ranging from 0 to 1, CWM of canopy height was standardized for each vegetation sample to 0–1 range, by dividing all values by the highest CH among the species in our data set (2.75 m).

RFSI was then confirmed to decrease in rich fen time series, which gave the basis for its calculation (Figure 2). RFSI was more or less constant or even increasing for those three rich fens, which had not terminated until the present times (Figure 3). They are characterized by quite high stress level, with RFSI fluctuating between 0.6 and 0.8 (Figure 3), whereas it dropped below 0.6 in those fens that abruptly changed to other ecosystems (Figure 2).

Analysing RFSI in 20 lowermost rich fen sections, we found that it was significantly negatively related to the distance to the rich fen finish (Figure 4).

**DISCUSSION**

In our research hypothesis, we proposed that ecosystem shifts in rich fens are preceded by decreases in stress intensity, which facilitates ecosystem engineer species and enhances ecosystem shifts. Indeed, by constructing a composite indicator based on stress-related functional traits, we were able to support this hypothesis, indicating that stress intensity decreased in most of the long-lasting mesotrophic rich fens that underwent a shift to another mire vegetation type. We interpret the decreasing stress as an autogenic cumulative process related to biotic feedbacks by ecosystem engineer species. We also found that such autogenic cumulative changes were absent in rich fens which produced very deep deposits, probably indicating a continuous peat accumulation over thousands of years until now. This finding suggests that under certain circumstances rich fens can be long-lasting and very stable ecosystems. However, as most of the profiles have not been dated, we have to consider that the thicker peat produced by fens with higher stress levels may also result from higher peat accumulation rates under such conditions and not from longer periods of accumulation, or that both explanations are true.

**Table 2.** The 95% Confidence Intervals for Average Change in Functional Traits CWM and Rich Fen Stress Index (RFSI) over 5-cm-Thick Peat Deposit (That is, During Rich Fen Development Lasting on Average about 100 Years), Calculated Over the Nine Rich Fens Which Finished in the Past

| trait | CWM | HF | CS small | CH (mm) | Non-M | RFSI |
|-------|-----|----|----------|---------|-------|------|
| 95% CI for average change/5 cm | 0.03–0.09 | 0.02–0.06 | 1–9 | –0.09 to –0.01 | –0.0038 to –0.0002 |

*As for the model with random intercept and slope iteration limit was reached without convergence, the results are shown for the model with random intercept only.*

*Only the results for glmm models for rich fen time series significant at 0.05 probability level and with 95% CI for average change not covering 0 are shown.*
Transitions of rich fens towards bogs or forests could happen as mere effects of autogenic (intrins-ic) changes, driven by a gradual expansion of sphagna or trees. On the contrary, the shift to tall helophytes (reed beds, eutrophic tall sedges) seems rather driven by some external factors causing flooding or increased discharge (extrinsic change). Therefore, it may look surprising why a decrease in stress-based RFSI index was found in most of the profiles documenting shifts of rich fens, regardless whether they changed into bogs, forests or semi-aquatic/helophyte/reed bed vegetation. Below, we propose an eco-hydrological explanation how the mentioned autogenic mechanism may, through an increased nutrient availability and a lowered hydrological buffering capacity of the peat, reduce fen resilience to both increased and decreased water inflow.

While analysing changes in stress-related PFTs during the development of mires, we noticed an increase in the share of hummock-forming species, species with a small clonal spread, a larger canopy height and a decrease in the share of non-mycorrhizal species (Table 2). We see several non-ex-clusive mechanisms that may explain coincidence of these trends towards an abrupt change in rich fen systems. First, as a result of the changes mentioned above, an increasingly hummocky habitat develops. Hummock-forming species are consid-ered ecosystem engineers in mires (Jones and others 1994; van Breemen 1995; Granath and others 2010; Hodges and Rapson 2010). With an increasing height and cover of hummocks, a mire becomes increasingly prone to peat mineralization, due to prolonged oxic conditions within hummocks. An increasing CH may indicate vegetation response to the increase in nutrient availability due to higher peat decomposition rate in drier hummocky habitat, as well as the community response to flooding, as tall species can tolerate it better than small ones (Moor and others 2017). The above explanations are in concert with the decreasing contribution of Non-M species during fen develop-ment. On the one hand, drier conditions/microhabitats created by an increase in HF and CS small species may promote root colonization by mycorrhizal fungi that usually respond negatively to anoxia (for example, Miller 2000; Unger and others 2009). These conditions also facilitate encroachment of shrubs and trees which are obliga-tory or facultative mycorrhizal, and can further facilitate the establishment of mycorrhizal fungal communities (for example, Soudzilovskaia and others 2015). Establishment of tussock species and small shrubs may also enhance hummock mosses by providing them scaffolding (Weber 1902). On the other hand, if rich fen development proceeds towards reed beds, the decrease in the proportion of non-mycorrhizal plants reflects the increase in helophytes such as Phragmites, Typha, Glyceria and Cladium. These helophytes are facultative mycor-rhizal, which may be beneficial for them for example, on the floodplains, where high water level fluctuations create alternate conditions en-abling and disabling mycorrhiza (Søndergaard and Lægaard 1977; Braendle and Crawford 1999). Rich fen development towards reed beds may be pre-ceded by an autogenic increase in hummockness, resulting in faster peat decomposition and peat compaction. When water level on such a fen rises due to external factors (river flooding or increased recharge for example, due to catchment defor-estation or changes in the regional climate), the

Figure 2. Rich fen stress indicator (RFSI) for the rich fens which finished in the past. The horizontal axis represents the distance to the rich fen finish (cm) expressed as a difference between peat depth of the first non-rich fen vegetation sample above the rich fen finish in a profile and peat depth of a separate vegetation sample—0 represents the end of rich fen. Rich fen transformation direction: triangle—into poor fen or bog, diamond—into fen woodlands, square—into reeds and tall sedges. Black dots—rich fen vegetation samples; white dots—non-rich fen vegetation samples up to 50 cm above the rich fen finish. Solid lines—linear regression lines for black dots; dashed lines—linear regression lines for black and white dots; only lines with regression coefficient significant at 0.05 probability level are shown.
compacted peat is less capable to buffer water level fluctuations [the mire does not swell in response to water table rise (compare Rydin and Jeglum (2013) for 'mire breathing')]. At the same time, compacted peat impedes groundwater through-flow. In effect, the flooding by river waters takes over groundwater seepage. In fact, the expansion of tall helophytes ('reed bed species') may be a combined effect of eutrophication due to peat decomposition and increased input of (eutrophic) surface water (compare Navrátilová and others 2017). Still, however, one should not exclude that the documented transitions to reed beds were fully externally driven and occurred independently from gradual autogenic processes leading to lowered plant stress.

An increased share of tussock sedges, hummock bryophytes or reeds in mesotrophic rich fen plant communities may cause or mark lower stability. Such features characterize many present-day rich fen systems. Our results suggest that such fens may be in their terminal successional stage, probably enhanced by anthropogenic alleviation of environmental stresses through drainage and/or eutrophication. Mowing management can mask symptoms of this shift; but as soon as it is stopped, an ecosystem shift may occur. Rich fens which are still open and devoid of tussock-forming sedges, hummock-forming bryophytes, shrubs and trees, should deserve special attention of conservationists as possibly rare examples of stable, potentially long-lasting systems that can be maintained by passive protection. Our results are in line with conclusions of Klimkowska and others (2019) from a cross-European meta-analysis of fen restoration projects, that restoration usually fails to re-establish func-

Figure 3. Rich fen stress indicator (RFSI) for three rich fens which are lasting until now (PAL81a) or terminated due to recent drainage (PAL100f, PAL100 h). Solid line—linear regression line for a separate profile; only lines with regression coefficient significant at 0.05 probability level are shown.

Figure 4. Median rich fen stress indicator (RFSI) values for c. 50-cm-thick peat sections in relation to the distance to rich fen finish. Black triangles—median RFSI for the lowest part up to 50 cm thick of each rich fen phase in a profile (excluding sections with less than three vegetation samples and excluding those that did not finish naturally); white triangles—median RFSI for c. 50-cm section of non-rich fen peat lying just above the rich fen phases finish. The horizontal axis represents the distance to the rich fen finish (cm) expressed as a difference between the first non-rich fen sample above the rich fen finish in a profile and the uppermost vegetation sample in the 50-cm section. Solid line—linear regression line for black triangles; dashed line—linear regression line for black and white triangles together; both regressions are significant at 0.05 probability level. Regression coefficient with 95% CI: solid line: \(-0.00043 \pm 0.00041\), dashed line: \(-0.00057 \pm 0.00032\).
tional fens because reference levels of plant stress are not achieved (Klimkowska and others 2019).

The rich fen stress indicator that we proposed is a potential predictive tool supporting conservation decisions in fens. However, it should not be regarded as a ready tool for conservationists, before being validated in contemporary fen communities.

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Compliance with Ethical Standards

Conflict of interest The authors declare that they have no conflict of interest.

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