brought to you by CORE

Check for updates



ARTICLE

https://doi.org/10.1038/s41467-020-18354-3

OPEN

Global patterns of the leaf economics spectrum in wetlands

Yingji Pan[®] ^{1⊠}, Ellen Cieraad[®] ¹, Jean Armstrong^{2,3}, William Armstrong[®] ^{2,3}, Beverley R. Clarkson[®] ⁴, Timothy D. Colmer³, Ole Pedersen[®] ^{3,5}, Eric J. W. Visser[®] ⁶, Laurentius A. C. J. Voesenek[®] ⁷ & Peter M. van Bodegom[®] ¹

The leaf economics spectrum (LES) describes consistent correlations among a variety of leaf traits that reflect a gradient from conservative to acquisitive plant strategies. So far, whether the LES holds in wetland plants at a global scale has been unclear. Using data on 365 wetland species from 151 studies, we find that wetland plants in general show a shift within trait space along the same common slope as observed in non-wetland plants, with lower leaf mass per area, higher leaf nitrogen and phosphorus, faster photosynthetic rates, and shorter leaf life span compared to non-wetland plants. We conclude that wetland plants tend to cluster at the acquisitive end of the LES. The presented global quantifications of the LES in wetland plants enhance our understanding of wetland plant strategies in terms of resources acquisition and allocation, and provide a stepping-stone to developing trait-based approaches for wetland ecology.

¹ Institute of Environmental Sciences (CML), Leiden University, Leiden, The Netherlands. ² Biological Sciences, University of Hull, Hull, UK. ³ School of Agriculture and Environment, The University of Western Australia, Perth, Australia. ⁴ Manaaki Whenua - Landcare Research, Hamilton, New Zealand. ⁵ Freshwater Biological Laboratory, University of Copenhagen, Copenhagen, Denmark. ⁶ Experimental Plant Ecology, Institute for Water and Wetland Research, Radboud University, Nijmegen, The Netherlands. ⁷ Institute of Environmental Biology, Utrecht University, Utrecht, The Netherlands. ⁸ email: y.pan@cml.leidenuniv.nl

uring the past two decades, trait-based ecology has advanced considerably. The leaf economics spectrum (LES) is an important component thereof. The LES provides convincing evidence of a consistent and continuous relationship among the leaf economics traits, reflecting a gradient of slow (conservative) to fast (acquisitive) strategies in terms of investment and use of nutrients and other resources^{1,2}. The LES has been shown to be present across different plant life forms and varied habitat types at a global scale and to a large extent independent of climate^{1,3}. Along the LES, species with higher leaf mass per area (LMA) generally have a longer leaf life span (LL), but a lower leaf nitrogen content (leaf N, wt/wt), and lower photosynthetic rates, at least on a mass basis (A_{mass}). This conservative strategy usually prevails in less fertile habitats. On the other hand, species with lower LMA, shorter LL, higher leaf N and photosynthetic rate have a faster return on investment of resources, commonly coinciding with nutrient-rich areas. Such trait-trait coordination in LES traits may be caused by underlying physiological and structural trade-offs⁴.

Studies on trait-trait relationships, including those on LES, have focused mainly on non-wetland terrestrial plants from a variety of ecosystems, such as forests or grasslands^{4,5} or on global analyses^{3,6}. However, whether the general LES also exists in global wetland ecosystems still remains unclear. This gap prevails despite the fact that leaf economics traits have been widely measured in wetland plants to study local plant functioning, community structure, growth and competition⁷.

A better understanding of trait-based relationships in wetlands is profoundly needed in light of the important ecosystem services provided by wetlands, including their role as the major carbon sink at a global scale⁸. Important ecological processes in wetlands such as methane emission and denitrification are linked to wetland plant functional traits^{9,10}. LES traits in wetlands are likely to play a role in these ecosystem processes and services^{11,12}. While the wide fertility gradient across different wetland types theoretically provides a natural gradient for the expression of LES from the acquisitive to conservative strategies¹³, additional constraints induced by adverse environmental conditions in wetlands compared to non-wetland systems mean that it cannot be taken for granted that LES traits will show similar patterns.

The varied environmental stressors unique to wetland ecosystems constrain plants that inhabit these systems. For example, intermittent/permanent flooding causes altered biogeochemical processes and the production of phytotoxic compounds such as ferrous iron (Fe²⁺) and sulphide (H₂S, HS⁻, S²⁻) in the substrates, as well as a less efficient way of producing ATP in cells experiencing an O₂ deficit¹⁴. In addition, reactive oxygen species (ROS), which can cause cellular macromolecule and membrane damage, accumulate in plant tissues especially upon return to aerobic conditions after flooding¹⁵. To survive in such an adverse environment, wetland plants have developed a suite of adaptive strategies¹⁵. Whether the LES also exists in wetlands depends, to a large extent, on whether the prevalent adaptive strategies of plants to environmental stressors are generally costly or cheap¹³. If adaptations are cheap, the LES should be unaffected and similar to non-wetland ecosystems. But if adaptive traits are costly, the LES should be shifted along the same axes (or even shifted in trait space entirely) to compensate this cost¹³. Moreover, leaf mass per area (LMA, one of the LES traits) seems to also be directly involved in flooding tolerance of wetland plants¹⁶, which may also lead to deviations within the LES.

Therefore, our research question is: What is the global leaf economics spectrum in wetlands? And how does it differ from that of non-wetland ecosystems? We hypothesize that wetland plants, in general, follow the LES strategies with fast-return species usually having lower LMA with increased leaf nutrient (N and P) content (wt/wt). This would naturally lead to faster photosynthesis in the day and higher dark respiration rate in the night. Assuming the trade-off between LMA and leaf longevity that exists in non-wetland plants³ also applies to wetlands, a lower LMA would result in a shorter leaf life span. Despite this general pattern, we also expect that the cost of developing the adaptive traits might affect the trait-trait relationships of leaf economics traits, and consequently shift the overall LES trait pattern in wetlands.

To test these hypotheses, we collected the LES traits measured in 365 wetland species of 184 families from 151 studies of both published and unpublished sources from a global scale. These wetland species are mainly from 10 wetland habitat types (including, as adapted from the Ramsar Convention¹⁷, artificial waterbodies, bogs, estuaries, fens, forested/shrub wetlands, mangrove swamps, marsh, rivers and lakes, temporary brackish/saline non-forested wetlands and temporary non-forested wetlands; see details in Appendix S1 of Pan et al.¹⁸). These habitat types occupy different positions along the gradients of two dominant drivers: hydrological regime (flooding depth and duration) and fertility (from oligotrophic to eutrophic)¹⁹. The wetland plant species analyzed in this study represent a full spectrum of plant characteristics and belong to eight life-form categories (emergent, floating-leaved, grass, isoetid, seagrass, sedge, shrub/tree and submerged). To take the effect of submergence on wetland plants into account, we carefully separated traits measured on plants of which only the root-zone or part of the stem was flooded of which tissues emergent above the water table were measured (hereafter called waterlogged wetland plants) vs. traits measured on plant tissues that were submerged (hereafter called submerged wetland plants).

By comparing these trait expressions with an extensive published dataset on non-wetland plants, we examine the trait-trait relationships of LES traits in wetland vs. non-wetland plants. We find that wetland plants, in general, tend to cluster at the acquisitive end of the LES compared to non-wetland plants, with lower leaf mass per area, higher leaf nitrogen and phosphorus, faster photosynthetic rates, and shorter leaf life span. Our global analyses on the LES in wetlands provide a useful perspective on the acquisition and turnover of resources of plants under stressful wetland conditions at a global scale. The results provide a stepping-stone to developing trait-based approaches for wetland ecology. In this way, we can better understand the strategies and functioning of wetland plants from a resource investment/gain perspective¹³. Therefore, studying LES traits in wetlands will not only extend our understanding of global plant strategies on resource acquisition and investment, but also give insight into wetland plant strategies and how these strategies are linked to ecosystem functioning^{11,13}.

Results

Overall bivariate trait-trait relationships. The overall trait-trait relationships of wetland plants showed similar trends as those among non-wetland plants in terms of the slope directions. Among the significant trait-trait relationships, five out of seven relationships of waterlogged plants had a lower R^2 than those of non-wetland plants (such as leaf P vs. leaf N and leaf N vs. LMA), while three out of four relationships for submerged plants had a lower R^2 than those of non-wetland plants (Table 1 and Supplementary Table 1). In addition, the confidence interval of R^2 of four non-significant trait-trait relationships of both plants of waterlogged and of submerged conditions showed no overlap with the (significant) R^2 observed for the corresponding trait-trait relationship of non-wetland plants. In combination, these results indicate weaker trait-trait relationships between wetland plant

61 (-0.68, -0.54) .31 (-1.74, -1.00) 	-1.01 (-1.180.87)	122/15217001	0.0 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4		
			-2.24 (-4.45, -1.12)	1.26 (0.97, 1.65)	Waterlogged
	-0.75 (-1.16, -0.48)	-1.98 (-2.54, -1.55)	0.84 (0.49, 1.44)		Submerged
12 (-0.02, -0.70)	-1.20 (-1.27, -1.13)	-1.30 (-1.38, -1.23)	-1.04 (-1.15, -0.95)	1.61 (1.50, 1.72)	Non-wetland
	1.60 (1.44, 1.77)	2.01 (1.67, 2.40)	2.36 (1.18, 4.71)	-1.89 (-2.96, -1.21)	Waterlogged
	1.53 (1.17, 2.00)	1.30 (0.90, 1.88)	0.59 (0.34, 1.05)	-1.70 (-43.06, -0.07)	Submerged
	1.51 (1.44, 1.58)	1.67 (1.57, 1.77)	1.44 (1.32, 1.57)	-2.10 (-2.25, -1.97)	Non-wetland
31* (P < 0.001; n = 264)		1.26 (1.01, 1.57)	-1.54 (-3.11, -0.76)	-1.31 (-2.13, -0.80)	Waterlogged
31^* ($P < 0.001$; $n = 41$)		1.58 (0.62, 4.05)	0.91 (0.36, 2.30)	-0.91 (-22.96, -0.04)	Submerged
0^* (P < 0.001; $n = 555$)		1.05 (0.91, 1.20)	0.89 (0.75, 1.06)	-1.00 (-1.14, -0.88)	Non-wetland
17^* (P < 0.001; $n = 90$)	0.12^* ($P = 0.003$; $n = 72$)		1.04 (0.54, 2.00)	-0.71 (-1.16, -0.43)	Waterlogged
$P^* (P = 0.001; n = 18)$	0.11 ($P = 0.468$; $n = 7$)		-0.48 (-0.82, -0.28)	I	Submerged
4^* (P < 0.001; $n = 537$)	0.19^* ($P < 0.001$; $n = 171$)		0.90 (0.83, 0.98)	-1.35 (-1.43, -1.28)	Non-wetland
02 ($P = 0.681$; $n = 11$)	0.13 ($P = 0.307$; $n = 10$)	$0.14 \ (P = 0.255; n = 11)$		0.53 (0.08, 3.30)	Waterlogged
27 ($P = 0.083$; $n = 12$)	0.13 ($P = 0.423$; $n = 7$)	$0.05 \ (P = 0.383; n = 16)$		1	Submerged
8^* (P < 0.001; $n = 221$)	0.37^* ($P < 0.001$; $n = 84$)	0.61^* ($P < 0.001$; $n = 220$)		-1.51 (-1.65, -1.38)	Non-wetland
35 (P = 0.015; n = 16)	$0.34 \ (P = 0.028; n = 14)$	$0.40 \ (P = 0.021; n = 13)$	$0.00 \ (P = 0.960; n = 4)$		Waterlogged
(P = 0.949; n = 3)	$0.02 \ (P = 0.917; n = 3)$	ı	ı		Submerged
$45^{*}(P < 0.001; n = 4.89)$	0.27^* (P < 0.001; $n = 173$)	0.69^* (P < 0.001: $n = 382$)	0.62^* (P < 0.001; $n = 187$)		Non-wetland
31* ($P < 0.001$; $n = 264$) 31* ($P < 0.001$; $n = 41$) ($P < 0.001$; $n = 41$) ($P < 0.001$; $n = 555$) ($P < 0.001$; $n = 555$) ($P < 0.001$; $n = 537$) ($P < 0.001$; $n = 18$) ($P < 0.001$; $n = 13$) ($P < 0.001$; $n = 12$) ($P < 0.001$; $n = 12$) ($P < 0.001$; $n = 16$) ($P < 0.001$;	$\begin{array}{c} 0.12^{*} \ (P=0.003; \ n=72)\\ 0.11 \ (P=0.468; \ n=7)\\ 0.19^{*} \ (P<0.001; \ n=171)\\ 0.13 \ (P=0.307; \ n=10)\\ 0.13 \ (P=0.423; \ n=7)\\ 0.37^{*} \ (P<0.001; \ n=84)\\ 0.34 \ (P=0.028; \ n=14)\\ 0.34 \ (P=0.028; \ n=14)\\ 0.20 \ (P=0.017; \ n=3)\\ 0.21 \ (P<0.011; \ n$		$\begin{array}{c} 1.26 (1.01, 1.57) \\ 1.26 (0.62, 4.05) \\ 1.58 (0.62, 4.05) \\ 1.05 (0.91, 1.20) \\ 0.014 (P = 0.255; n = 11) \\ 0.05 (P = 0.383; n = 16) \\ 0.61^* (P < 0.001; n = 220) \\ 0.40 (P = 0.021; n = 13) \\ 0.40^* (P < 0.001; n = -382) \end{array}$	$\begin{array}{c} 1.26 \ (1.01, 1.57) \\ 1.58 \ (0.62, 4.05) \\ 1.58 \ (0.62, 4.05) \\ 1.05 \ (0.91, 1.20) \\ 1.05 \ (0.91, 1.20) \\ 1.04 \ (0.54, 2.00) \\ 0.90 \ (0.83, 0.98) \\ 0.90 \ (0.90 \ $	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

traits than corresponding relationships among non-wetland plants (Table 1, lower left section), and suggests that wetland plants are less constrained within the LES with larger trait variation. A summary of the results of all standardized major axis (SMA) analyses is given in Table 2.

Bivariate trait relationships between leaf P, N and LMA. Leaf P and leaf N were positively correlated, across non-wetland plants³, waterlogged wetland plants ($R^2 = 0.31$) and submerged wetland plants ($R^2 = 0.31$). The SMA analysis revealed that there was no significant difference in slopes of leaf P-leaf N associations between non-wetland plants and wetland plants (P = 0.30 and P = 0.91 for waterlogged and submerged wetland plants, respectively). However, the parallel slopes of both waterlogged and submerged wetland plants were elevated compared to nonwetland plants (both P < 0.001), which indicates that at a given leaf N, wetland plants tended to have a higher leaf P than nonwetland plants. Moreover, there was a significant shift along the common slope towards higher values in wetland plants (both P < 0.001; Fig. 1a). This suggests that the proportional change of leaf P with leaf N of wetland plants was similar to non-wetland plants, while wetland plants generally had higher leaf N and leaf P than non-wetland plants.

Leaf N and LMA were negatively correlated in non-wetland and wetland plants (Table 1). The waterlogged wetland plants had a significantly flatter slope (P < 0.001), while submerged wetland plants had a significantly steeper slope (P < 0.001). Thus, as LMA decreases, the increase in leaf N was less pronounced in waterlogged wetland plants, while such increase of leaf N was steeper in submerged wetland plants, compared to non-wetland plants (Fig. 1b).

Leaf P and LMA were negatively correlated in both wetland and non-wetland plants with similar slopes (P = 0.04 and P =0.03 for waterlogged and submerged wetland plants, respectively, Fig. 1c). Wetland plants had a parallel slope which is shifted towards the upper left corner (P < 0.001) compared with nonwetland plants. This indicates that even though leaf P and LMA maintained similar relationships in non-wetland and wetland plants, wetland plants maintained a higher value of leaf P but a lower value of LMA (Fig. 1c).

Bivariate trait relationships with photosynthetic rate. The slopes of photosynthetic rate-leaf N in wetland plants were similar to those of non-wetland plants (P = 0.06 and P = 0.18 for waterlogged and submerged wetland plants, respectively, Fig. 2a). However, waterlogged wetland plants were significantly shifted along a common slope towards a higher photosynthetic rate and leaf N values (P < 0.001) and had an elevated parallel slope (P < 0.001) compared to non-wetland plants, indicating that at given leaf N, waterlogged wetland plants had a higher photosynthetic rate. This suggests that waterlogged wetland plants had a higher nitrogen use efficiency (photosynthesis per unit investment of leaf N). No significant shift along the common slope nor elevation differences among parallel slopes were detected for submerged wetland plants (P = 0.61 and P = 0.20, respectively).

There were no significant differences in slopes of photosynthetic rate-leaf P between wetland plants and non-wetland plants (P = 0.16 and P = 0.36 for waterlogged and submerged wetlandplants, respectively Fig. 2b). However, wetland plants of both conditions showed a significant shift along the common slope towards higher photosynthetic rate and leaf P values (both P < 0.001). This suggests a similar proportional change between leaf P and photosynthetic rate of both wetland plants and nonwetland plants, while wetland plants had higher values of photosynthetic rate and leaf P than non-wetland plants. No Ľ

80

ı.

₹ **6**

٩

z

<u>log</u> ı

og LMA <0.00</pre> 0.001 0.058

> 80 80 80 80

n.a. n.a.

n.a. n.a. **0.001 0.001** n.a. 0.056 0.005 0.005

000 0.032 0.429 0.052

		1				0.695	.p.r
·			·	·	·	-	-
						m	
						.29	Ъ.
1 1	1		1	1	1	0	
						.217	ъ.
1 1 1	1 1	I		1	1	o.	Ċ.
84 0 84	84 2 66	8 2 84	2 8 8 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	5 6	66		
0.0 0.0	0.0 0.7	0.7	0.7 0.0	0.0	0	<u>с</u>	n.d.
-					-		
				~		\sim	
				073	821	300	ъ.
1 1	I		I	o.	o.	o.	Ċ
8	8	88	88	2	30	1	
 0.5	0.5	- 0.5	0.5	!	0.0	0.0	n.d.
5	5	5	5	5	-		
)25 762 727	727	727	727	2	2	040	34
0.0	0.0	0.0	0.7	Ś	2	0.0	0.1
555	555	5 5	5	5	.	~	~
0.00	0.00	0.0	0.0		046	487	379
, ¥ ¥ ¥	∀ ∀ ₹	∀ ₹	v	1	o.	0.	Ö.
22		5	2	5	8	52	74
0.13	0.13	5	1 2 1 2 1 0	0.10	0.9(0.56	0.8.
	-			_	_	-	
	_						
60 19 19 19	12 2 0	12 16	-16			02	04
0 .0 0.0 0.7	0 .0 0.7 0.4	0.2 0.4	0.4		D.a	0.0	0.1
5 5	5						
0.0	00. 0	568		381	<i></i>	725	719
₽₽ o o	v öö	0 0	Ö.		n.ĉ	0	0
<u>6</u> 00 0	6	c	ת	6	4	8	4
).89 30.0	0.08		0.14	0.15	No.	0.49	0.85
		C	'	0	0	J	0
3 36	536		Ë.	472	50	231	÷
0 0 - 0	0. L 0	цо Ц	C	5	ŝ.	0.1	n.c
•					-		

differences in slopes (first column of each tra

n.d.

80

The differences in slopes (first column of each t and vs. submerged wetland plants (second row

conditions, the shift along the common

those

(indicated as n.d. = not determined)

vation differences among parallel slopes were detected (P =0 and P = 0.65 for waterlogged and submerged wetland plants, pectively), suggesting that wetland plants and non-wetland nts have a similar photosynthetic rate per unit leaf P.

The photosynthetic rate-LMA associations were similar ween waterlogged wetland plants and non-wetland plants, sept for a significant shift (P < 0.001) along the common slope vards the corner of lower LMA values but higher photosyntic rates. This suggests that waterlogged wetland plants erally had lower LMA, but a higher photosynthetic rate. For omerged wetland plants, the photosynthetic rate-LMA slope s significantly steeper than for non-wetland plants (P < 0.01). is shows that the decrease of photosynthetic rate with an rease per unit of LMA was stronger in submerged wetland nts, indicating that the effect of changed leaf structure on the otosynthesis was bigger in submerged wetland plants. In other rds, the photosynthetic rate of submerged wetland plants was en more reduced by an increase of LMA (Fig. 2c). The nificantly different slopes of submerged plants also imply a ft in trait space.

variate trait relationships with dark respiration rate. For the ationship between dark respiration rate vs. leaf N, submerged tland plants showed a significantly flatter slope (P < 0.01) than n-wetland plants. This suggests that submerged wetland plants intained their respiration rate to a lower level as leaf N reases than non-wetland plants (Fig. 2d). Moreover, subrged wetland plants tended to have a lower dark respiration e at a given leaf P (P < 0.001), LMA (P < 0.001) or photothetic rate (P < 0.01) (Fig. 2e–g). In combination, this suggests t for dark respiration, trait relationships are substantially difent for submerged plants than for non-wetland plants.

For waterlogged wetland plants, the patterns of dark respiran rate were less apparent. For dark respiration rate vs. leaf N, found no significant difference in the slopes (P = 0.15), nor a ft along the common slope (P = 0.06), nor elevation ferences among parallel slopes (P = 0.42) between the waterged wetland plants and non-wetland plants (Fig. 2d). For leaf no significant slope (P = 0.13) was observed, but waterlogged tland plants showed a significant shift along the common slope vards higher dark respiration rate and leaf P values (P < 0.001) g. 2e). In both cases, the small sample size for dark respiration e could have reduced the statistical power to detect differences slope, and particularly in the case of dark respiration rate vs. f N the differences in slope was substantial (Supplementary ble 2). Also for dark respiration rate vs. LMA, a P-value of 0.03 combination with the low sample size n = 11, suggested the sence of a different slope for wetland vs. non-wetland plants g. 2f). Finally, for photosynthetic rate vs. dark respiration rate, significant differences were observed for waterlogged vs. nontland plants (Fig. 2g).

variate trait relationships with leaf life span. How leaf traits co-vary with the leaf life span (LL) in submerged wetland plants remains uncertain, because of the limited number of data points (n = 3 for LL-leaf N and LL-leaf P, and the absence of data linking LL-LMA, LL-photosynthetic rate and LL-dark respiration rate). For waterlogged wetland plants, we found a significant lower parallel slopes between LL-leaf N than non-wetland plants (P < 0.01, Fig. 3a), suggesting that at a given leaf N, waterlogged wetland plants had a shorter leaf life span. We found no significant differences in the relationships between leaf life span and other traits for waterlogged wetland plants (P > 0.01, Fig. 3b–e).

In summary, compared with non-wetland plants, significantly different slopes were detected in the relationship between leaf

Table 2 Comparison of the bivariate relationships in wetland vs. non-wetland plants.



Fig. 1 The bivariate trait relationships between leaf P, N and LMA. a Leaf phosphorus (leaf P) vs leaf nitrogen (leaf N). **b** Leaf N vs. leaf dry mass per unit area (LMA). **c** Leaf P vs. LMA. The waterlogged and submerged wetland plants are shown in light blue squares and dark blue triangles, respectively. The non-wetland plant data from GLOPNET³ are shown in red circles with a solid red line. If the slope for wetland plants differs significantly from that of non-wetland plants, this is indicated by a solid dark or light blue line, for waterlogged and submerged plants, respectively. Dashed lines with the notation of **shift* and/or **ele*. identify a significant shift along the common slope, and/or significant elevation differences among parallel slopes, respectively. Note that graph axes are log₁₀ scaled.

N-LMA in both waterlogged and submerged wetland plants (Fig. 1b), and between photosynthetic rate-LMA (Fig. 2c) and dark respiration rate-leaf N (Fig. 2d) in submerged wetland plants only. This suggests that submerged wetland plants have even more trait deviations from non-wetland plants than waterlogged wetland plants. In general, wetland plants tended to have a lower LMA with higher leaf N and leaf P contents, and consequently higher photosynthetic rate and shorter leaf life span. For submerged wetland plants, the photosynthetic rate was constrained by an increase in LMA. However, this increase was compensated by a much more gradual increase in dark respiration rate with increasing leaf N, than was evident for non-wetland plants.

Discussion

We compared leaf economics spectrum (LES) trait associations of wetland and non-wetland plants and found that the LES does exist in wetland plants, but with weaker and often deviating/ shifting trait-trait associations relative to the non-wetland LES. The weaker trait-trait associations (as indicated by the lower coefficients of determination (R^2) of trait-trait relationships) suggest that alternative strategies exist among wetland plants to deal with the complex and adverse wetland conditions with specific stressors. It may also suggest that besides nutrients and light, other limitations in wetlands also influence the LES and require alternative strategies and consequently the special leaf structure and function of wetland plants. This would cause a higher variation in LES traits. Besides habitat N and P fertility, leaf N can be driven by various factors, including potassium (K), temperature, phytotoxins, or the plants' intrinsic maximal growth rate⁷. Habitat wetness may also drive leaf N through two indirect

mechanisms. On the one hand, denitrification caused by prolonged soil flooding may decrease nitrate availability, thus reducing leaf N²⁰. On the other hand, species living in wet habitats usually have a lower LMA, and thus tend to have a higher leaf N^{21,22}. The more variable leaf N may further affect the expression of trait-trait associations in wetland plants, such as the leaf Nphotosynthetic rate associations²³ and the leaf N-dark respiration rate associations²⁴.

Our results indicate that the general directions of relationships among LES traits are maintained in wetland plants, which suggests that the principal ecological links behind the trait-trait associations have similarities with those in non-wetland systems³. However, our study also reveals differences and these support previous suggestions that wetland plants might possess a unique functional behaviour in photosynthesis-related activities due to their specific adaptation to wetland conditions^{25,26}. There are five key aspects in which the LES of wetland plants seems to differ profoundly from the non-wetland LES.

Firstly, in general, wetland plants have a lower LMA, higher leaf N and leaf P content, and a higher photosynthetic rate than non-wetland plants. The waterlogged wetland plants show a shorter leaf life span. Unfortunately, the pattern of submerged wetland plants is uncertain for leaf life span due to a limited number of data points. We conclude that wetland plants comply with a fast-return strategy in resource acquisition among the majority of the LES trait-trait associations²⁷. Thus, while nutrient and carbon cycling rates in wetland soils are generally slower compared with non-wetland systems¹¹, the aboveground carbon and nutrient cycles in wetlands are expected to be faster.

Secondly, a major deviation in LES trait-trait relationships of wetland plants compared to non-wetland plants occurs in the leaf N-LMA relationship (Fig. 1b). The different behaviour of LMA



Fig. 2 The bivariate associations with photosynthetic rate and dark respiration rate. a Photosynthetic rate (A_{mass}) vs. leaf nitrogen (leaf N). **b** A_{mass} vs. leaf phosphorus (leaf P). **c** A_{mass} vs. leaf dry mass per unit area (LMA). **d** Dark respiration rate (R_{mass}) vs. leaf N. **e** R_{mass} vs. leaf P. **f** R_{mass} vs. LMA. **g** A_{mass} vs. R_{mass} . The waterlogged and submerged wetland plants are shown in light blue squares and dark blue triangles, respectively. The non-wetland plant data from GLOPNET³ are shown in red circles with a solid red line. If the slope for wetland plants differs significantly from that of non-wetland plants, this is indicated by a solid dark or light blue line, for waterlogged and submerged plants, respectively. Dashed lines with the notation of **shift* and/or **ele*. identify a significant shift along the common slope, and/or significant elevation differences among parallel slopes, respectively. Note that graph axes are log₁₀ scaled.



Fig. 3 The bivariate relationships with leaf life span. a Leaf life span vs. leaf nitrogen (leaf N). **b** Leaf life span vs. leaf phosphorus (leaf P). **c** Leaf life span vs. leaf dry mass per unit area (LMA). **d** Leaf life span vs. photosynthetic rate (A_{mass}). **e** Leaf life span vs. dark respiration rate (R_{mass}). The waterlogged and submerged wetland plants are shown in light blue squares and dark blue triangles, respectively. The non-wetland plant data from GLOPNET³ are shown in red circles with a solid red line. The dashed light blue line with the notation of **ele*. identifies significant elevation differences among parallel slopes respectively. Note that graph axes are log_{10} scaled and the absence of leaf life span data coupled to LMA, A_{mass} , or R_{mass} for submerged plants.

highlights the different functional role of LMA in wetland plants^{16,28}. This complies with experimental studies that have found some low-LMA leaves of hydrophytic wetland plants to be functionally highly acquisitive^{21,22}. However, in addition to further stimulating the acquisition of nutrients, we also expect that a lower LMA is essential to deal with the lower CO₂ and O₂ availabilities to the leaves in (partially) submerged conditions²⁹. Therefore, besides its leaf economics aspect, LMA should be considered also as a key wetland trait.

Thirdly, a lower LMA may have important implications for the functioning of the remainder of the LES in wetland conditions. In non-wetland low nutrient conditions, plants tend to conserve their nutrients by increasing their LMA to protect the leaves against herbivory and other damages³⁰. Our results suggest that such protection of leaves is not feasible in wetlands. In addition, the higher leaf N in wetland plants may also cause an increased risk of herbivory³¹. Together, these processes partially explain the higher herbivory rates in wetland ecosystems compared to nonwetland terrestrial ecosystems³¹. One way to compensate for the higher losses is to become more acquisitive. Such a strategy is supported by the shift in LES traits along the common slope, but may also relate to the elevated leaf P at a given LMA. The results on the leaf N to leaf P relationships suggest that leaf P is even more elevated in wetland plants than leaf N (Fig. 1a). Through these changes in leaf nutrient economics, wetland plant species may raise their photosynthetic capacity in order to create faster growth dynamics (and concomitant higher turnover).

Fourthly, wetland plant species seem to go even further in stimulating photosynthetic capacity. The photosynthetic rate of waterlogged plants was elevated at a given leaf N compared to the photosynthesis-leaf N relationships in non-wetland plants. Leaf N (and leaf P) expresses the combination of photosynthesis-related active nutrients and those nutrients used for storage and protection³². If wetland plants indeed invest less energy in the protection of their leaves, the fraction of nutrients involved in photosynthesis increases⁴, which in turn would explain the elevated photosynthetic rate of waterlogged plants that we observed. The lower LMA itself may also influence the leaf Nphotosynthetic rate relationships, thus increasing the leaf N efficiency of photosynthesis²³. Finally, some submerged aquatic plants are able to enhance their photosynthesis with special leaf structure, such as thin cuticles and oriented chloroplasts towards the epidermis^{22,25}.

Lastly, leaves of submerged wetland plants have a lower dark respiration rate (mass basis) than expected from a comparison with the non-wetland LES. Oxygen can decline to hypoxic levels during submergence, and especially in shallow water bodies during the night³³. Low oxygen can restrict aerobic respiration, both in roots³⁴ and in leaves³⁵. The relatively low dark respiration rate in leaves of wetland plants may be due to a lower investment of resources in leaf construction and maintenance, and related reductions of energy requirements and respiration during the night²⁴. The lower respiratory demand allows to more readily face hypoxia when leaves become submerged. In addition, leaves with porous tissues will enhance the oxygen status of the innermost cells. Note that, although the adaptive formation of aerenchyma will significantly decrease the cell oxygen consumption on a tissue volume basis³⁶, the data analysed here are measurements expressed on a tissue mass basis. Hence, aerenchyma formation per se does not explain the patterns found in this study.

Some of these mechanisms may be further amplified at submerged conditions, where we additionally observed that the altered leaf structure may also affect the photosynthetic rate through a deviating A_{mass} -LMA relationship (Fig. 2c), and through influencing the respiration rate by deviating R_{mass} -leaf N associations (Fig. 2d). We found a significant reduction of the photosynthetic rate at a given LMA in submerged wetland plants. The additional limitation to photosynthesis of submerged wetland plants can be due to the much lower light availability with water depth and turbid water²⁹. However, the unique adaptive traits evolved in wetland plants such as leaf gas films and aerenchyma tissues should enhance the gas exchange/flux in plant tissues^{15,35}, and therefore partially compensate the costs posed by the adverse wetland conditions. This may explain the observed pattern that the photosynthetic rate at a given leaf N and leaf *P*-value was not affected (Fig. 2a, b).

All of the described significant changes in the slope of trait-trait relationship, in the position along the slope or due to shifted parallel slopes were detected based on a rather conservative *P*-value threshold (P < 0.01) in this study. This threshold was chosen to help ensure that the most ecologically relevant relationships were detected in relatively large datasets (e.g. a relationship with an R^2 of only 0.05 is already significant at P = 0.05 at a sample size of n = 77). However, for those relationships with smaller sample sizes (in particular in relation to dark respiration rate and leaf life span), this approach may have resulted in overly conservative interpretation. This indicates that deviations in the LES of wetland plants may include even more trait-trait relationships than identified here. Future research could serve to quantify whether these patterns are robust in the face of more data.

Altogether, our analysis suggests that the direct link between photosynthetic rate and dark respiration rate, as evidenced from non-wetland plants to complement N-rich enzymatic and other metabolic components that lead to a higher respiration cost when maintaining a high photosynthetic rate^{24,37}, also exists in wetland plants. However, such a relationship is expressed differently in wetland plant species compared with non-wetland plants. The results from our analysis show that submerged wetland plants are capable of having lower dark respiration rate at a given photosynthetic rate than non-wetland plants.

When upscaling the findings to wetland ecosystem functioning, we ascribe the generally high productivity in wetland ecosystems globally to the adaptation of wetland plants by having generally fast-return strategies and a higher payback rate. In this way, the adverse wetland conditions may have very limited impact on the wetland plant functioning in terms of resource accumulation. The assumed trade-offs between the cost of adaptation to wetlands and plant function from the leaf economics spectrum perspective are therefore not profound in general¹³. In addition, there are some environmental stressors that rarely happen in wetlands. For example, drought stress, which is a common problem in terrestrial ecosystems, is less constraining in most wetlands, and might move LES traits of wetland plants to the optimum end with lower LMA with higher leaf nutrient content^{16,38}. The combination of the high productivity in wetlands and the retarded biochemical cycling rate in the anoxic environments of the substrates together make wetlands the largest contributor to the terrestrial biological carbon pool⁸.

Methods

Data compilation. We defined wetland plants as plants that mainly occur in (or are exposed to) wetland habitats as described by the Ramsar Convention¹⁷. We summarized the 3 major groups including 42 sub-groups wetland habitat types in Ramsar Convention to be 12 categories (as estuary, intertidal wetland, mangrove swamps, rivers and lakes, brackish and saline inland wetlands, permanent non-forested wetlands, temporary non-forested wetlands, permanent forested wetlands, artificial waterbodies, marsh, bog, and fen). We collected leaf economics traits for wetland plants on a global scale including those plants exposed to intermittent/ permanent wetland conditions (waterlogged or flooded) from both field and experiment measurements. The wetland plant leaf economics trait dataset was compiled based on a systematic search in Web of Science and Google Scholar (last updated on the 5th June 2018). The literature search included permutations of the following keywords: wetland plants, marsh plant, bog plant, isoetid, aquatic plants,

ARTICLE

macrophytes, submerged plants, floating-leaved plants, emergent plants, mangroves, leaf economics traits, leaf economics spectrum, leaf nitrogen, leaf phosphorus, SLA, LMA, leaf life span, photosynthetic rate, underwater photosynthetic rate, dark respiration rate. Additionally, our network of wetland experts from around the world contributed recommendations for possible literature that we had overlooked. Finally, we added unpublished data of our own and of our network. We did not include data from other trait databases that are dominated by terrestrial records, including TRY, because the few records available for wetland plants in these databases do not have a sufficiently detailed habitat description that would allow the differentiation between waterlogged and submerged required for our analysis.

We followed the nomination system in The Plant List (http://www.theplantlist. org) to unify all plant synonyms names from the original references to a unique and consistent accepted name.

We supplemented the trait observations in our database with Ellenberg moisture indicator values. The Ellenberg moisture indicator is a classic index which generally reflects the plants' adaptation/acclimation to habitat wetness. Plant species can be categorized into 12 levels from those occupying very dry habitats (level 1) to strictly aquatic plants (level 12)³⁹. For the current meta-analysis, we selected plant species with Ellenberg moisture value > 7 to represent wetland plants, as described in detail in Supplementary Methods. For these species, we selected records of the six LES traits (leaf nitrogen, leaf phosphorus, leaf dry mass per unit area, leaf life span, photosynthetic rate, and dark respiration rate). We took trait values for the same six traits for non-wetland plant traits (of 1569 species) from the GLOPNET database for comparison³. For a consistent analysis of trait-trait tradeoffs, we expressed all leaf economics traits on a mass basis. Mass-based and areabased traits can be interconverted via a division by LMA. The mean value for each trait of each species was used (using the median did not alter the interpretation of the general pattern, as shown in Supplementary Figs. 1-3 and Supplementary Tables 3 and 4.). We used species-mean values to attain a sufficient number of trait-trait combinations for a given species. We assume that the trait observations used for calculating the species-mean values were representative for the environmental/growth conditions in which the species occurs. Possible uncertainty in species trait mean values (for example due to intra-specific variation) will then result in noise in trait-trait relationships. In total, 365 wetland species of 184 families from 151 studies were compiled and analyzed. A map of the sampling sites with accurate spatial location information can be found in Supplementary Fig. 4. The species are from varied life forms, including grasses, sedges, seagrasses, shrubs/ trees, emergent, floating-leaved, isoetid, and submerged plants. Traits of most (308) species had been measured at waterlogged conditions, with submerged measurements being available for 75 species. The leaf trait data of wetland and non-wetland plant species analyzed in this paper can be found in the DRYAD repository (https://doi.org/10.5061/dryad.v6wwpzgsq).

Statistical analysis. The slope and its associated coefficient of determination (R^2) of each trait pair within the six LES traits of waterlogged and submerged wetland plants at the species level was calculated by a standardized major axis (SMA) analysis⁴⁰. The slopes and R^2 -values were compared to those of trait-trait relationships of non-wetland plants as derived from the GLOPNET³. The evaluation was based on the comparison between waterlogged wetland plants and submerged wetland plants, with non-wetland plants, respectively.

We tested each trait-trait relationship within the above-mentioned six LES traits for deviations between wetland and non-wetland plants. No test was run for the associations between leaf life span and LMA, photosynthetic rate and dark respiration rate of submerged wetland plants due to too few data points. In our SMA analysis, we conducted three tests, one to evaluate differences in slopes (i.e. steeper or shallower trait-trait relationships between wetland vs. non-wetland plants), a second to assess shift along the common slope (i.e. a more predominant position of wetland plants on either the conservative or acquisitive end of LES), and a third to assess whether trait associations of wetland and non-wetland plants can be characterized as having elevation differences among parallel slopes (suggesting a specific trait would be more -or less- costly in wetland conditions)⁴⁰:

Test A: sma(y~x*groups) tests for differences in slopes fitted for

different groups

Test B: $sma(y \sim x+groups, shift = T)$ tests for a shift along the common slope Test C: $sma(y \sim x+groups)$ tests for elevation differences among parallel slopes between groups

A significant difference in slope (Test A) implies a difference in the direction and location of the relationship in trait space. Since the location and direction of lines with different slopes are not comparable⁴¹, tests B and C were only run if there was no significantly different slope detected in Test A. If all three tests were non-significant, we conclude that wetland and non-wetland plants have similar trait-trait relationships.

The *P*-value is strongly depended on sample size, and it does not measure the size of an effect or the importance of a result⁴². In this study, we set a rather conservative *P*-value threshold (P < 0.01) for our tests. This was done to help reducing type I errors and to ensure that the most ecologically relevant relationships (with a reasonable effect size) were detected in these datasets⁴³.

The statistical analysis used R software⁴⁴. The major axes analysis was conducted with the sma() and ma() function in the smatr package (version 3.4-8)⁴⁰.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The plant trait data that support the findings of this study and underlie its figures and tables can be downloaded from the DRYAD repository (https://doi.org/10.5061/dryad. v6wwpzgsq). The GLOPNET³ data from which our non-terrestrial data were derived are available from: https://www.nature.com/articles/nature02403#Sec15.

Code availability

Source code file is available in the Supplementary Software.

Received: 25 January 2020; Accepted: 19 August 2020; Published online: 09 September 2020

References

- Reich, P. B., Walters, M. B. & Ellsworth, D. S. From tropics to tundra: Global convergence in plant functioning. *Proc. Natl Acad. Sci. USA* 94, 13730–13734 (1997).
- Shipley, B. et al. Reinforcing loose foundation stones in trait-based plant ecology. *Oecologia* 180, 923–931 (2016).
- Wright, I. J. et al. The worldwide leaf economics spectrum. Nature 428, 821–827 (2004).
- Onoda, Y. et al. Physiological and structural tradeoffs underlying the leaf economics spectrum. N. Phytol. 214, 1447–1463 (2017).
- Dray, S. et al. Combining the fourth-corner and the RLQ methods for assessing trait responses to environmental variation. *Ecology* 95, 14–21 (2014).
- Diaz, S. et al. The global spectrum of plant form and function. Nature 529, 167–171 (2016).
- Güsewell, S. Variation in nitrogen and phosphorus concentrations of wetland plants. Perspect. Plant Ecol. Evol. Syst. 5, 37–61 (2002).
- Page, S. E. & Baird, A. J. Peatlands and global change: response and resilience. Annu. Rev. Environ. Resour. 41, 35–57 (2016).
- Sutton-Grier, A. E. & Megonigal, J. P. Plant species traits regulate methane production in freshwater wetland soils. Soil Biol. Biochem. 43, 413–420 (2011).
- Alldred, M. & Baines, S. B. Effects of wetland plants on denitrification rates: a meta-analysis. *Ecol. Appl.* 26, 676–685 (2016).
- Moor, H. et al. Towards a trait-based ecology of wetland vegetation. J. Ecol. 105, 1623–1635 (2017).
- Sutton-Grier, A. E., Wright, J. P. & Richardson, C. J. Different plant traits affect two pathways of riparian nitrogen removal in a restored freshwater wetland. *Plant Soil* 365, 41–57 (2013).
- Pan, Y., Cieraad, E. & van Bodegom, P. M. Are ecophysiological adaptive traits decoupled from leaf economics traits in wetlands? *Funct. Ecol.* 33, 1202–1210 (2019).
- Lambers, H., Chapin, F. S. & Pons, T. L. Plant Physiological Ecology. (Springer New York, 2008). https://doi.org/10.1007/978-0-387-78341-3.
- Colmer, T. D. & Voesenek, L. A. C. J. Flooding tolerance: suites of plant traits in variable environments. *Funct. Plant Biol.* 36, 665–681 (2009).
- Douma, J. C., Bardin, V., Bartholomeus, R. P. & van Bodegom, P. M. Quantifying the functional responses of vegetation to drought and oxygen stress in temperate ecosystems. *Funct. Ecol.* 26, 1355–1365 (2012).
- Ramsar Convention Secretariat. The Ramsar Convention Manual: a guide to the Convention on Wetlands (Ramsar, Iran, 1971). https://doi.org/10.1007/ 978-94-007-0551-7 (2013).
- Pan, Y. et al. Drivers of plant traits that allow survival in wetlands. Funct. Ecol. https://doi.org/10.1111/1365-2435.13541 (2020).
- Keddy, P. A. Wetland ecology: principles and conservation. (Cambridge University Press, UK, 2010).
- Ordoñez, J. C. et al. Leaf habit and woodiness regulate different leaf economy traits at a given nutrient supply. *Ecology* 91, 3218–3228 (2010).
- Mommer, L., Lenssen, J. P. M., Huber, H., Visser, E. J. W. & De Kroon, H. Ecophysiological determinants of plant performance under flooding: a comparative study of seven plant families. *J. Ecol.* 94, 1117–1129 (2006).
- Pierce, S., Brusa, G., Sartori, M. & Cerabolini, B. E. Combined use of leaf size and economics traits allows direct comparison of hydrophyte and terrestrial herbaceous adaptive strategies. *Ann. Bot.* 109, 1047–1053 (2012).
- Reich, P. B., Ellsworth, D. S. & Walters, M. B. Leaf structure (specific leaf area) modulates photosynthesis-nitrogen relations: Evidence from within and across species and functional groups. *Funct. Ecol.* 12, 948–958 (1998).
- Reich, P. B. et al. Relationships of leaf dark respiration to leaf nitrogen, specific leaf area and leaf life-span: a test across biomes and functional groups. *Oecologia* 114, 471–482 (1998).

- Mommer, L., Pedersen, O. & Visser, E. J. W. Acclimation of a terrestrial plant to submergence facilitates gas exchange under water. *Plant, Cell Environ.* 27, 1281–1287 (2004).
- Herzog, M. & Pedersen, O. Partial versus complete submergence: Snorkelling aids root aeration in Rumex palustris but not in R. acetosa. *Plant, Cell Environ.* 37, 2381–2390 (2014).
- 27. Reich, P. B. The world-wide 'fast-slow' plant economics spectrum: a traits manifesto. *J. Ecol.* **102**, 275–301 (2014).
- Violle, C. et al. Plant functional traits capture species richness variations along a flooding gradient. *Oikos* 120, 389–398 (2011).
- Colmer, T. D., Winkel, A. & Pedersen, O. A perspective on underwater photosynthesis in submerged terrestrial wetland plants. *AoB Plants* 11, 1–15 (2011).
- Westoby, M., Falster, D. S., Moles, A. T., Vesk, P. A. & Wright, I. J. Plant ecological strategies: some leading dimensions of variation between species. *Annu. Rev. Ecol. Syst.* 33, 125–159 (2002).
- Cyr, H. & Face, M. L. Magnitude and patterns of herbivory in aquatic and terrestrial ecosystems. *Nature* 361, 148–150 (1993).
- Hikosaka, K. & Shigeno, A. The role of Rubisco and cell walls in the interspecific variation in photosynthetic capacity. *Oecologia* 160, 443–451 (2009).
- Pedersen, O., Colmer, T. D., Borum, J., Zavala-Perez, A. & Kendrick, G. A. Heat stress of two tropical seagrass species during low tides—impact on underwater net photosynthesis, dark respiration and diel in situ internal aeration. N. Phytol. 210, 1207–1218 (2016).
- 34. Armstrong, W. & Beckett, P. M. Experimental and modelling data contradict the idea of respiratory down-regulation in plant tissues at an internal [O2] substantially above the critical oxygen pressure for cytochrome oxidase. N. Phytol. 190, 431-441 (2011).
- Colmer, T. D. & Pedersen, O. Underwater photosynthesis and respiration in leaves of submerged wetland plants: gas films improve CO₂ and O₂ exchange. *N. Phytol.* 177, 918–926 (2008).
- Jackson, M. B. & Armstrong, W. Formation of aerenchyma and the processes of plant ventilation in relation to soil flooding and submergence. *Plant Biol.* 1, 274–287 (1999).
- 37. Reich, P. B. et al. Scaling of respiration to nitrogen in leaves, stems and roots of higher land plants. *Ecol. Lett.* **11**, 793-801 (2008).
- Pagter, M., Bragato, C. & Brix, H. Tolerance and physiological responses of Phragmites australis to water deficit. *Aquat. Bot.* 81, 285–299 (2005).
- Ellenberg, H. H. Vegetation ecology of central Europe. (Cambridge University Press, 1988).
- Warton, D. I., Duursma, R. A., Falster, D. S. & Taskinen, S. smatr 3- an R package for estimation and inference about allometric lines. *Methods Ecol. Evol.* 3, 257–259 (2012).
- Warton, D. I., Wright, I. J., Falster, D. S. & Westoby, M. Bivariate line-fitting methods for allometry. *Biol. Rev. Camb. Philos. Soc.* 81, 259–291 (2006).
- 42. Wasserstein, R. L., Schirm, A. L. & Lazar, N. A. Moving to a World Beyond "p < 0.05". Am. Stat. **73**, 1–19 (2019).
- Nakagawa, S. & Cuthill, I. C. Effect size, confidence interval and statistical significance: a practical guide for biologists. *Biol. Rev.* 82, 591–605 (2007).
- R. Core Team. R: A language and environment for statistical computing. R Foundation Statistical Computing, Vienna, Austria. URL https://www.rproject.org/ (2018).

Acknowledgements

The establishment of the wetland trait database was first discussed and started in 2008 at the Vegfunction WG39 which was funded by ARC-NZ Research Network for Vegetation Function. We would like to thank all additional contributors to this original workshop, including Paul Adam (U New South Wales, Sydney, AU), Margaret Brock (U New England, Armidale, USA), George Ganf (U Adelaide, Adelaide, AU), Irving A. Mendelssohn (Louisiana State U, Baton Rouge, USA), Eliska Rejmánkova (U California, Davis, USA), Brian Sorrell (Aarhus U, Aarhus, DK), and Evan Weiher (U Wisconsin, Eau Claire, USA), Yingji Pan is grateful to support from the China Scholarship Council (Grant No. 201606140037).

Author contributions

P.v.B. initialized this research. Y.P., P.v.B. and E.C. designed and planned the research. Y.P. and P.v.B. compiled the data with inputs from J.A., W.A., B.R.C., T.D.C., O.P., E.J. W.V. and L.A.C.J.V. All analyses were done by Y.P. with inputs from all co-authors. Y.P., P.v.B. and E.C. wrote the first drafts of the paper, which was further improved by inputs from all co-authors.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41467-020-18354-3.

Correspondence and requests for materials should be addressed to Y.P.

Peer review information *Nature Communications* thanks Helen Moor, Ian Wright and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Peer reviewer reports are available.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/ licenses/by/4.0/.

© The Author(s) 2020