

RESEARCH ARTICLE

Vegetation characteristics control local sediment and nutrient retention on but not underneath vegetation in floodplain meadows

Lena Kretz^{1,2*}, Elisabeth Bondar-Kunze^{3,4}, Thomas Hein^{3,4}, Ronny Richter^{1,5,6}, Christiane Schulz-Zunkel², Carolin Seele-Dilbat^{1,2}, Fons van der Plas⁷, Michael Vieweg², Christian Wirth^{1,5,8}

1 Systematic Botany and Functional Biodiversity, Life science, Leipzig University, Leipzig, Germany, **2** Department of Conservation Biology and Social-Ecological Systems, Helmholtz Centre for Environmental Research (UFZ), Leipzig, Germany, **3** Institute of Hydrobiology and Aquatic Ecosystem Management, University of Natural Resources and Life Sciences, Vienna, Austria, **4** WasserCluster Lunz, Lunz am See, Austria, **5** German Centre for Integrative Biodiversity Research (iDiv) Halle-Jena-Leipzig, Leipzig, Germany, **6** Geoinformatics and Remote Sensing, Institute for Geography, Leipzig University, Leipzig, Germany, **7** Plant Ecology and Nature Conservation, Wageningen University, Wageningen, The Netherlands, **8** Max Planck Institute of Biogeochemistry, Jena, Germany

* lena.kretz@uni-leipzig.de



OPEN ACCESS

Citation: Kretz L, Bondar-Kunze E, Hein T, Richter R, Schulz-Zunkel C, Seele-Dilbat C, et al. (2021) Vegetation characteristics control local sediment and nutrient retention on but not underneath vegetation in floodplain meadows. PLoS ONE 16(12): e0252694. <https://doi.org/10.1371/journal.pone.0252694>

Editor: Michael A. Chadwick, King's College London, UNITED KINGDOM

Received: May 17, 2021

Accepted: November 16, 2021

Published: December 2, 2021

Copyright: © 2021 Kretz et al. This is an open access article distributed under the terms of the [Creative Commons Attribution License](https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Data Availability Statement: Data are available from the Data Repository iData: <https://doi.org/10.25829/idiv.3479-10-1609>.

Funding: C.W. received funding from the German Federal Ministry of Education and Research (BMBF) and the German Federal Agency for Nature Conservation (BfN) for the project: Wilde Mulde - Revitalization of a wild river landscape in Central Germany (Funding label: 01LC1322E). There was

Abstract

Sediment and nutrient retention are essential ecosystem functions that floodplains provide and that improve river water quality. During floods, the floodplain vegetation retains sediment, which settles on plant surfaces and the soil underneath plants. Both sedimentation processes require that flow velocity is reduced, which may be caused by the topographic features and the vegetation structure of the floodplain. However, the relative importance of these two drivers and their key components have rarely been both quantified. In addition to topographic factors, we expect vegetation height and density, mean leaf size and pubescence, as well as species diversity of the floodplain vegetation to increase the floodplain's capacity for sedimentation. To test this, we measured sediment and nutrients (carbon, nitrogen and phosphorus) both on the vegetation itself and on sediment traps underneath the vegetation after a flood at 24 sites along the River Mulde (Germany). Additionally, we measured biotic and topographic predictor variables. Sedimentation on the vegetation surface was positively driven by plant biomass and the height variation of the vegetation, and decreased with the hydrological distance (total $R^2 = 0.56$). Sedimentation underneath the vegetation was not driven by any vegetation characteristics but decreased with hydrological distance (total $R^2 = 0.42$). Carbon, nitrogen and phosphorus content in the sediment on the traps increased with the total amount of sediment (total $R^2 = 0.64, 0.62$ and 0.84 , respectively), while C, N and P on the vegetation additionally increased with hydrological distance (total $R^2 = 0.80, 0.79$ and 0.92 , respectively). This offers the potential to promote sediment and especially nutrient retention via vegetation management, such as adapted mowing. The pronounced signal of the hydrological distance to the river emphasises the importance of a laterally connected floodplain with abandoned meanders and morphological depressions.

no additional external funding received for this study.

Competing interests: The authors have declared that no competing interests exist.

Our study improves our understanding of the locations where floodplain management has its most significant impact on sediment and nutrient retention to increase water purification processes.

Introduction

Worldwide, streams and rivers suffer from large loads of sediment and nutrients, which is predominantly caused by anthropogenic activities [1–3]. Soil erosion and overfertilization, caused by industrial agriculture and forestry, increase the loads of sediment and nutrients in river systems and cause eutrophication and siltation [4–6]. Additionally, the process of sediment transport along the river is often interrupted by hydro-engineering infrastructure [6]. River floodplains, however, can act as a sink for sediment and its associated nutrients by retaining these during floods [7,8], thus providing the important ecosystem function of sediment and nutrient retention [9,10].

Natural floodplains reduce sediment and nutrient transport to downstream areas during inundation. Especially in hydrologically connected systems, a large amount of the annual riverine sediment and nutrient load can be retained in floodplains. The amount increases with the inundation duration and the area of inundation [11]. Further, models predict that the inundation depth increase the floodplain roughness and with that might increase sedimentation on the floodplain [12]. The accumulated nutrients can have a positive effect on the productivity of the floodplain vegetation [13]. However, anthropogenic activities have strongly diminished floodplain areas, due to channelization, embankments, bank stabilization, and river straightening [7,14,15]. Consequently, worldwide floodplains are considered threatened ecosystems [14,15]. As a result, floodplain restoration efforts have increased during the last decades. Many countries started programs emphasizing the river-floodplain reconnection for restoring ecological conditions, but also for flood protection. Furthermore, reconnection measures are expected to affect the retention capacity of floodplains [16], but its drivers still need to be better integrated into river and floodplain restoration and management [17]. However, to manage floodplains for optimal sediment and nutrient retention, we need to understand how vegetation structure, as well as the composition and diversity of plant communities, affect sedimentation and how these biotic drivers interact with the hydromorphological control.

Sediment retention is a complex phenomenon that depends on the flood and different biogeomorphic processes in the floodplain [18,19]. While deposition of coarse sediment is mostly influenced by the topography of the floodplain, the vegetation type and structure influencing fluvial processes and sediment transport [20,21] are most relevant for sedimentation of finer grain sizes [12,18,22]. Communities of herbaceous vegetation were more efficient in accumulating fine sediment compared to shrublands and floodplain forests [23], and reed beds caused more nitrogen and phosphorus deposition than grass and woodlands [13]. Within a flume experiment, we showed in a previous study, that the structural characteristics of the community (biomass, density, height, structural diversity, and leaf pubescence) increase sedimentation under controlled conditions [24]. However, this is the first study that investigates *in situ* measurements of a real flood event by (1) focusing on sedimentation within the vegetation, separating the process of sedimentation on vegetation from the process of sedimentation underneath the vegetation, (2) investigated the role of species diversity, leaf surface structure and community structure, and (3) combined these vegetation characteristics with topographical parameters of the floodplain, thus allowing to quantify the relative importance of vegetation and topography.

The sediment retention capacity of a floodplain is known to vary with different structural parameters of the vegetation, mostly measured around (in front and behind) vegetation patches. Generally, it was found that biomass increases sediment retention [22,25], which was also the case in the flume experiments for sedimentation on the vegetation [24,26] and partly also underneath the vegetation [24]. Dense floodplain vegetation has been suggested to be very efficient in accumulating fine sediment [23,27]. It reduces the flow velocity and thus allows sediment to sink and deposit [28,29]. Here, also the variation of the vegetation height may have an impact on sedimentation, since varying vegetation height cause turbulence and might increase and decrease flow velocities locally. In the flume experiment a negative relationship was found between height variation and sedimentation on the vegetation [24]. It was found that the deposition of finer sediment (silt and clay) is controlled by vegetation height in herbaceous floodplain vegetation [30].

Riparian zones and floodplain meadows are hotspots of biodiversity [15]. At the same time, they are one of the most threatened habitats in the world [31,32]. Despite this, species diversity *per se* is rarely studied in the context of sediment retention on floodplains, even though it is known to determine other ecosystem functions such as productivity and nutrient dynamics [33]. The results of the flume experiment only showed evidence for effects of species richness on sedimentation in the absence of identity effects [26]. Species diversity has also been shown to correlate with structural diversity of vegetation [34], which was found to increase sedimentation [24]. Dedicated biodiversity experiments have revealed that diverse grasslands exploit the growing space in a complementary fashion and thus have a higher density and taller stature than less diverse grasslands [35,36]. While we account for these two variables directly, there may be additional effects that go beyond the mean characteristics of the vegetation. Combining for example tall/sparse with small/dense plant species may be particularly effective for sediment retention. The trait combination might increase the overall sedimentation irrespective of total density or stature. No significant effects of the species diversity of herbaceous vegetation on sediment retention were found in front of, and behind a vegetation patch when comparing monocultures with a three-species mixture in an experiment [37]. However, the investigation of a longer diversity gradient under field conditions could yield another picture.

Besides vegetation structure, leaf surface structure of the vegetation also matters for sedimentation. In particular, leaf pubescence has been shown to positively and leaf area on non-haired leaves negatively drive sediment retention at the level of herbaceous leaf surfaces [24,38,39]. Therefore, the mean expression of these traits in the vegetation may also be important for sedimentation at the level of floodplain vegetation patches, which has rarely been considered in studies on sedimentation in herbaceous vegetation.

Topographic variables are the main abiotic factors that could explain sediment distribution within the floodplain. Discharge and with it, inundation depth are strongly affected by elevation and with that the location within the floodplain is relevant for sedimentation. Fine sediment is transported farther along the river and into the floodplain than coarse sediment and only settles in areas with reduced flow velocity [29]. In general, sedimentation was found to decrease with increasing distance from the river [27,29,40]. However, a straight line does not necessarily represent the topographic diversity of a dynamic riverine floodplain and the winding path the water travels into the floodplain during floods. Therefore, the length of the shortest path of lowest elevation is a better measure of the ways the river water travels from the river into the floodplain during floods. Such a measure for the true 'hydrological distance' may thus better represent the topography of the floodplain. Some studies used other terms to describe a similar measure such as the flow path [41,42] or the hydrological connectivity [16,43,44].

Flood magnitude and duration are the fundamental controls on the delivery of sediment transported by a river from its channel onto its floodplain [45]. Flood magnitude and duration

also affect the spatio-temporal distribution of depth and velocity of the inundating water as it flows across the floodplain surface [46]. However, the amount and particle size of sediment retained across the floodplain is strongly moderated by two local, natural, factors. First, floodplain topography imposes strong local variations on the depth and velocity of the flood water [20,21,44]. Second, any vegetation cover on the floodplain surface interacts with the flood water, slowing local flow velocities and thus encouraging the settlement of sediment particles from the water [20,21,44]. However, because the structure of herbaceous vegetation is flexible, it can respond to high flow velocities by bending and adopting a more streamlined form, and this leads to changes in its ability to intercept and retain sediment as flow depth and velocity vary through time and across space [47–49]. Understanding of these complex interactions is critical to building a picture of the importance of floodplain topography and vegetation cover for the retention of sediments and associated nutrients. While experiments in laboratory flumes can go a long way towards delivering such understanding, particularly by manipulating flow depth, velocity and transported sediment load across vegetated surfaces displaying different traits, the results of such experiments need to be validated and elaborated within the complexity of real-world situations.

In this paper, we present a set of field observations exploring the detailed influence of natural topographic features and meadow vegetation communities on finer sediment and nutrient retention during a single flood event. Although the results of such field investigations are limited by the magnitude and duration of the investigated event, they can reveal detailed and highly informative spatial patterns that are relevant to floods of similar magnitude at other sites and they can reveal associations that may be applicable to floods of differing magnitude and duration. In the present case, we investigated a flood of relatively low magnitude that occur on average biannually, but others already emphasised the importance of low magnitude, but higher frequent floods for nutrient retention on floodplains [50]. We seek to reveal the detailed impact of topographic setting and different traits of flood meadow vegetation on sediment and nutrient retention during flood events of relatively low magnitude and high frequency.

Thus, the aim of this study was to analyse combined effects of vegetation characteristics and selected topographic features at the scale of vegetation patches on the floodplain. First, we disentangled sedimentation on and underneath the vegetation under *in situ* conditions, second we quantified the relative importance of vegetation characteristics in relation to topographical parameters and third, we investigated the effects of additional vegetation characteristics (species diversity and leaf surface structure) on sedimentation within a vegetation patch. We tested the following hypotheses:

- (H1) Sedimentation on and underneath the vegetation increases with increasing vegetation biomass, cover, vertical density, vegetation height and height variation.
- (H2) Sedimentation on and underneath the vegetation decreases with increasing hydrological distance from the river.
- (H3) Sedimentation on and underneath the vegetation increases with increasing plant species diversity.
- (H4) Sedimentation on the vegetation increases with increasing leaf pubescence and decreasing mean leaf area.
- (H5) Total carbon (C), nitrogen (N) and phosphorus (P) in the sediment on and underneath the vegetation increase with the total amount of sediment deposited.

Material and methods

Study site

The study was located along the Mulde River in Central Germany (Fig 1), close to its mouth into the Elbe River. Along this river section, the river still flows in its natural bed and has been only moderately modified by hydro-engineering infrastructures and bank stabilization in the past. About half of the cut-banks are not embanked. The study took place in the frame of the restoration project 'Wilde Mulde-Revitalisation of a dynamic riverine landscape in Central Germany'. The project area extends between the towns Raguhn and Dessau ($51^{\circ}43'-46' N$, $12^{\circ}17'-18' E$). Within the project area, we defined three floodplains as study areas in 2016 (Fig 1). The Mulde River is dammed around 22 km upstream of the project area and has another

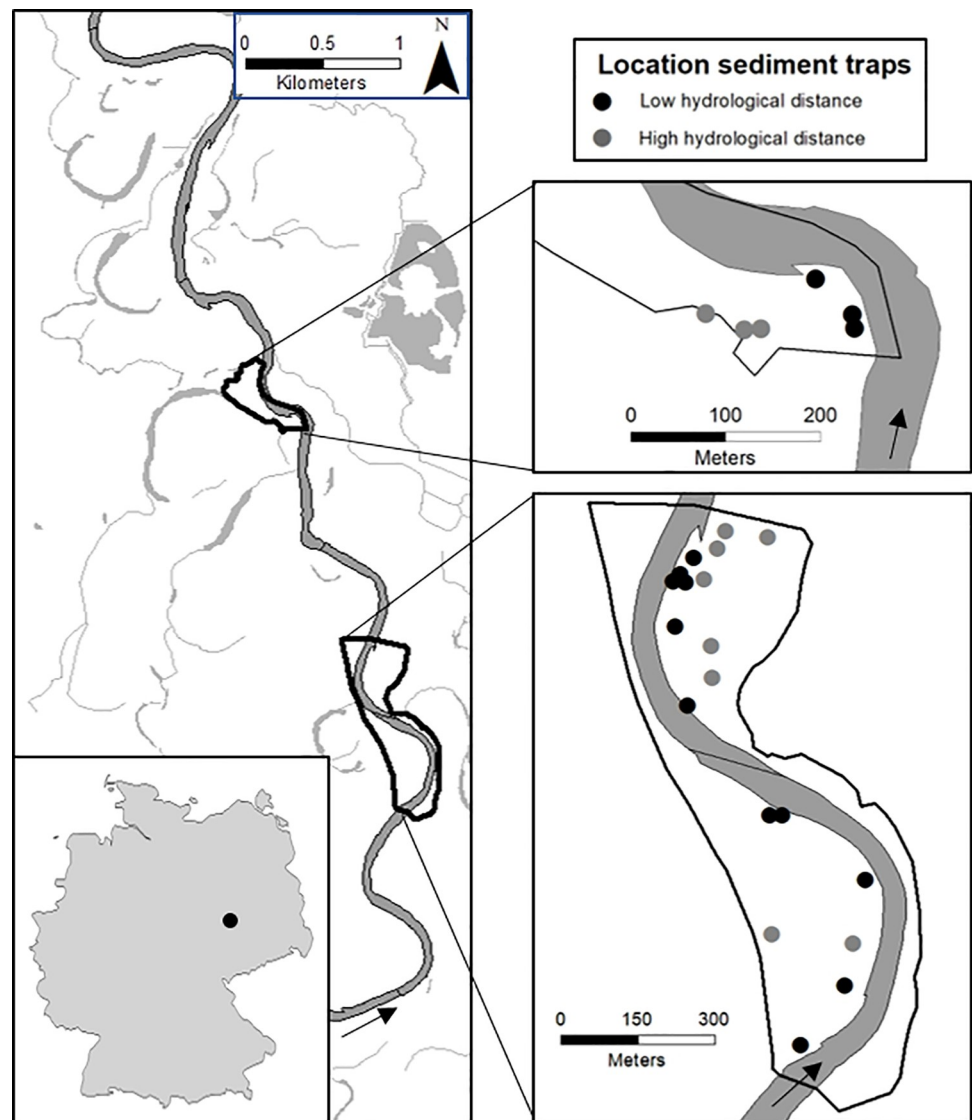


Fig 1. Map of the study site. Map of the three floodplains in two areas along the Mulde River with trap locations. The grey line is the Mulde River and the black frames are the study areas. Low hydrological distance = Short pathway of lowest elevation the water takes to the plot. High hydrological distance = Long pathway of lowest elevation the water takes to the plot.

<https://doi.org/10.1371/journal.pone.0252694.g001>

smaller weir about 5 km upstream of the first study area. Upstream of the study areas, the Mulde River has a mean discharge of $67 \text{ m}^3 \text{ s}^{-1}$ (gauging station ‘Priorau 560090’). In general, the study area is a mosaic of hardwood and softwood floodplain forests and meadows, with our study focusing on the floodplain meadows.

The topography of the floodplain meadows is strongly formed by the river, creating a mosaic of steep slip-off slopes with gravel banks in front, depressions, and abandoned meanders further away from the river that get reconnected during floods. The dominant species in the meadows are, depending on microtopography and management, *Arrhenatherum elatius*, *Bromus inermis*, *Calamagrostis epigejos*, *Elymus repens* and *Phalaris arundinacea*.

In February 2017 a small flood occurred for several days with overbank flow conditions. The peak discharge was $353 \text{ m}^3 \text{ s}^{-1}$, which equals a flood with a discharge occurring on average every second year. However it was the only discharge of such high magnitude that occurred during the 5-years study period (2016–2020).

Vegetation data

In summer 2016, we established a grid of vegetation plots. Within the three study areas, plots were selected to span the elevation gradient of the slip-off slope and the floodplain meadow above mean flow conditions using a stratified random sampling strategy. In autumn 2016 we selected 54 plots (18 plots per study area) for this study using with the following criteria: (i) plots are fully covered by vegetation; (ii) plots span a gradient of vegetation height (ranging from 36 cm to 124 cm); (iii) lower elevation plots were given preference, due to their higher probability to get flooded; (iv) depressions and abandoned meanders at distance to the river were also represented, while ensuring that the selection still represents the whole elevation gradient. With this approach, the plots are representative for the floodplain and at the same time form an observational design by spanning gradients for regression analysis. Within each plot (2 m x 2 m) we identified all vascular plant species and estimated the cover of each species in summer 2016 before the flood. We calculated the Shannon diversity index [51] based on cover. Overall, we inventoried 44 species with the species richness ranging from 2 to 10 species per plot.

Vegetation characteristics

We measured the maximum height of the vegetation using two metrics: (i) the maximum inflorescence height (highest inflorescence), which represents the maximum vegetation height, and (ii) the maximum canopy height (highest leaf), which represents the maximum height of the vegetation surface. Both metrics were measured with the help of a meter stick five times per plot (in the middle of the square plot and at arm length inside the plot from each corner). We measured the vegetation height at that time point no matter if the vegetation hung over or not. We did this once in summer 2016 before the flood and once in spring 2017 after the flood, trying to catch the point in time that represents the vegetation structure during the flood best. Additionally, we took images of side views in the form of cross sections of the vegetation in spring 2017 on all flooded plots to estimate the density and height distribution of the vegetation during the flood. To this end, we placed a camera, 1 m with 90° angle in front of the plot (Fig 2). At 50 cm inside the plot we positioned a camera background wall so that every image shows exactly the first 50 cm of the plot (Figs 2 and S1). We carefully pushed down the vegetation outside the plot with a flooring material. Afterwards we analysed the images with the statistical software R [52] for height and density distribution in the same way as done in the flume experiment [24]. From these structural images, we derived the variables vertical density, mean height, median height, and height variation (Table 1, S1 Fig). The images were colour

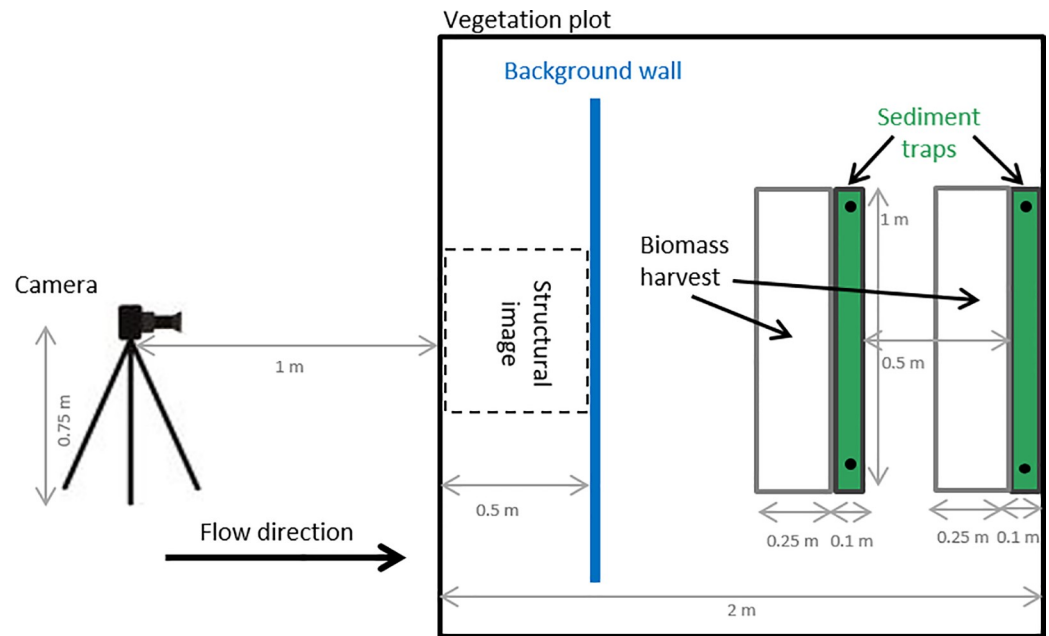


Fig 2. Set-up of different measurements in the vegetation plots (2 m x 2 m). The set-up of the green sediment traps (image in S2 Fig) and the defined area of biomass harvest upstream each plot after the flood event. It also shows the set-up for the structural images of the vegetation with the camera 1 m in front and the blue background wall 0.5 m inside the plot (image in S1 Fig).

<https://doi.org/10.1371/journal.pone.0252694.g002>

normalised and resampled from a resolution of 4000 by 6000 pixels to a resolution of 400 by 600 pixels and afterwards transformed into grey-scale images. In order to perform a binary classification of the image into vegetation and background, we used the otsu-tresholding

Table 1. List of predictor variables.

Hypothesis	Predictor	Unit	Details	Sampling date
H1	Vegetation cover	%	Estimate of vegetation cover	summer before flood 2016
H1	Biomass	g m ⁻²	Dry weight of biomass harvested after the flood	after flood 2017
H1	Vertical density	%	Percent of vegetation pixels on the image of standard size	after flood 2017
H1	Mean height	length*	Mean height of vegetation pixels on the image	after flood 2017
H1	Median height	length*	Median height of vegetation pixels on the image	after flood 2017
H1	Height variation	length*	Standard deviation of vegetation pixel height on the image	after flood 2017
H1	Highest leaf 16	Cm	Mean of 5 point measurements of the highest leaf	summer before flood 2016
H1	Highest inflorescence 16	Cm	Mean of 5 point measurements of the highest inflorescence	summer before flood 2016
H1	Highest leaf 17	Cm	Mean of 5 point measurements of the highest leaf	after flood 2017
H1	Highest inflorescence 17	Cm	Mean of 5 point measurements of the highest inflorescence	after flood 2017
H2	Hydrological distance	M	Length of lowest path the river water takes to the plot	
	Elevation above river	M	Elevation of plot above mean flow conditions of the river: $e_r = e_p - e_{mf}$	
	River kilometre	Km	Location along the river (last tributary used as point 0)	
	Precipitation		Some rainfall while collection of the sediment traps (categorical: no, yes)	after flood 2017
H3	Shannon diversity index		Sum of proportion of species times ln of proportion of species	summer before flood 2016
H4	Leaf pubescence	%	Sum of cover of hairy species	summer before flood 2016
H4	Leaf area	cm ²	Mean leaf area per species times species cover on the plot	summer before flood 2016

Predictor variables with detailed explanations, units and sampling dates.

* the length is standardized between the images, however not calibrated to any unit.

<https://doi.org/10.1371/journal.pone.0252694.t001>

method [53], as implemented in the package EBImage [54]. All variables are described in Table 1.

Study design

For investigating sedimentation on the floodplain, we used artificial lawn (*Kunstrasen Arizona*, Hornbach, 1.05 g m⁻² lawn, 26 cm lawn height, S2 Fig) as sediment traps—a commonly used and established method [29,55]. The material has several advantages: (i) it can be easily cut to the required size; (ii) it can be flexibly and firmly fixed to the ground, and (iii) it exposes a surface with a high capacity to collect and keep sediment. To keep the sward structure as intact as possible, we cut the artificial lawn into narrow strips (10 cm x 100 cm strips), which were carefully inserted into the vegetation. Two sediment traps were fixed at the downstream side of each plot, to well represent the conditions of the vegetation on the plot (Figs 2 and S2). While sediment traps represent a good method to measure sedimentation on a standardized surface (thus only affected by surrounding vegetation and its effects on fluvial processes), a limitation is that it removes the effects of the local fine-scale vegetation structure and composition on sedimentation. Combining measures of sedimentation on the vegetation itself, as well as on sediment traps, may be best to partition the effects of fluvial processes (caused by surrounding conditions) and local vegetation properties on sedimentation. We deployed the sediment traps on all 54 plots in January 2017 and fixed them with tent stakes and steel washers (56 cm outer diameter). During the flood in February 2017, 24 plots were inundated (Fig 1). We collected the sediment traps immediately after the flood retreated. In addition, we also harvested the patch of biomass directly in front (upstream) of the trap, to quantify the vegetation biomass that influence the sedimentation on the trap (Fig 2). In the lab, we quantified the sediment by washing it off the traps with a few litres of water and dried the sediment-rich water in beakers in a compartment drier at 70°C. Afterwards, the dry sediment was weighed. The same was done with the sediment on vegetation and, additionally, we dried and weighed the biomass itself. The two sediment trap samples per plot were pooled together as were the two biomass samples per plot.

Nutrient analysis

All sediment samples on the vegetation and on the traps (except two samples with too little sediment) were sieved (< 2 mm) and analysed for C, N and P. To determine the total C and N concentration, the dried sediment samples were ground to a fine powder in a ball mill (Retsch MM2, Vienna, Austria). The homogenized sample was weighed, placed in tin caps and measured by using the Elemental Analysis Isotope Ratio Mass Spectrometry (EA-IRMS; EA—Thermo Scientific™ FLASH 2000 HT™; IRMS—Thermo Scientific™ Delta V™ Advantage) [56]. To determine the total P concentration, the sediment was also ground to a fine powder in a ball mill (Retsch MM400). The homogenized samples were measured by using the Inductively Coupled Plasma Optical Emission Spectrometry (ICP-OES "Arcos", Spectro, Kleve, D). As indicator for the nutrient quality the N:P ratio was calculated.

Topographical variables

The elevation and position of the single plots were measured with a Differential GPS (R8, Trimble Inc.) The mean elevation of the four plot corners e_p was expressed as elevation above the river e_r (as $e_r = e_p - e_{mf}$). Mean water level height e_{mf} was calculated using the closest gauging station (Priorau, 560090; daily measurements 1995–2015). The elevation differences between the study areas were derived from the digital elevation model (DEM, © GeoBasis-DE, LVerm-Geo LSA, [m.E. 2016, C22-7009893-2016]). The hydrological distance was defined as the

length of the shortest path of lowest elevation that the river water takes to a single plot in the floodplain. It was derived using the flow accumulation approach on the DEM of the floodplain area and calculated using the TopoToolbox 2 [57] in MATLAB [58]. We included longitudinal stream distances as river kilometre in the study to account for the plot location along the river, since we visually observed lower flow velocity at the study area further downstream. The river kilometre was measured along the middle line of the river starting from the last tributary to the river upstream of the study area. We chose this tributary as the zero point because it is the last major tributary. Precipitation occurrence was included as a categorical variable, because some of the traps experienced rainfall after the flood, before all traps could be collected.

Leaf surface traits

We also included two leaf surface traits, leaf pubescence and leaf area (at plot-level—see below), as predictors of sedimentation, because we showed, with an earlier flume experiment, that, in controlled settings, pubescence can increase leaf surface sedimentation and that sedimentation increases with decreasing leaf area on leaves with no or just a few hairs [39]. Out of the 44 species, we classified five as pubescent species (*Carex hirta*, *Galium aparine*, *Urtica dioica*, *Verbascum densiflorum* and *Veronica maritima*). We quantified plot-level pubescence as the summed cover of these five species. Data about the mean area of individual leaves were obtained from TRY—a global database of plant traits [59] TRY version 5.0; data used of [60–71]. Three species were not included in the leaf area calculation, since they either had no leaves (*Cuscuta europaea* and *Equisetum pratense*) or because there were not data available in the TRY database (*Carex praecox*). All three species occurred on a maximum of two plots, and in these, with densities below 5% cover. For an estimate of the leaf area per plot, we summed the cover-weighted leaf areas of all species per plot.

Data analysis

All statistical analyses were done with the statistical software R [52]. We ran two separate linear models to investigate which factors drove sedimentation on the vegetation and on the sediment traps. We also calculated the ratio of sedimentation on the vegetation to the sedimentation on the traps and run a separate linear model to explain it. Further, we ran six linear models to explain total amount of C, N and P in the sediment on the vegetation and in the sediment on the traps. We used the candidate variables listed in Table 1 as explanatory variables; in the ratio model we additionally included the interaction of the river kilometre and the hydrological connectivity, while in the C, N and P models, we additionally used the sediment amount as an explanatory variable. To meet model requirements regarding the normality of the error distribution, the two variables, “sedimentation on traps” (except for the ratio of sediment on vegetation to on traps) and “hydrological distance”, were natural log-transformed. We scaled all continuous variables to ensure comparability of the model estimates. To avoid multicollinearity, we removed explanatory variables with a variation inflation factor above 5.0 (vif function, car library, [72]). With the remaining variables, we ran a backward model selection procedure and selected the final model with best model fit based on Akaike’s Information Criterion (stepAIC function, MASS library, [73]). We tested the differences of the N:P ratios close and far from the river using paired two-sample t-tests. Therefore, the plots were separated by the mean of the hydrological distance.

Results

General results

The median sedimentation on the vegetation was 28.60 g m⁻², while on the traps the median sedimentation was about double (60.55 g m⁻², Table 2). Both, sedimentation on the vegetation

Table 2. Descriptive statistics.

Variables	Unit	Min	Max	Mean	Median	Sd
Sediment on vegetation	g m^{-2}	10.36	105.56	37.33	28.60	25.96
Sediment on traps	g m^{-2}	4.25	4955.50	832.57	60.55	1440.33
C in sediment on vegetation	g m^{-2}	0.82	18.79	4.67	3.76	3.88
N in sediment on vegetation	g m^{-2}	0.05	1.00	0.37	0.36	0.22
P in sediment on vegetation	g m^{-2}	0.01	0.28	0.10	0.09	0.07
C in sediment on traps	g m^{-2}	0.56	178.49	26.09	3.98	42.68
N in sediment on traps	g m^{-2}	0.04	12.88	1.88	0.30	3.06
P in sediment on traps	g m^{-2}	0.02	3.78	0.87	0.16	1.09
Vegetation cover	%	7.90	90.20	50.77	52.61	21.31
Biomass	g m^{-2}	30.12	499.16	239.51	219.36	116.20
Vertical density	%	0.08	0.35	0.20	0.19	0.05
Mean height	length*	0.09	0.55	0.25	0.20	0.10
Median height	length*	0.09	0.55	0.24	0.21	0.10
Height variation	length*	0.01	0.18	0.06	0.03	0.05
Highest leaf 16	cm	36.00	124.00	72.08	73.00	26.00
Highest inflorescence 16	cm	0.00	141.00	66.17	75.50	43.87
Highest leaf 17	cm	16.00	72.00	31.25	23.00	16.37
Highest inflorescence 17	cm	0.00	91.00	14.67	0.00	29.61
Hydrological distance	M	2.83	586.13	142.53	91.82	156.82
Elevation above river	M	0.26	1.71	1.24	1.31	0.37
River kilometre	km	3.64	6.98	5.15	4.99	1.08
Shannon diversity index		0.14	1.73	1.12	1.16	0.44
Leaf pubescence	%	0.00	37.50	6.90	2.50	9.27
Leaf area	cm^2	234.29	3906.17	1487.25	1602.99	879.92

Descriptive statistic of all continuous variables. Min = minimum, Max = maximum, Sd = Standard deviation.

* the length is standardized between the images, however not calibrated to any unit.

<https://doi.org/10.1371/journal.pone.0252694.t002>

and on the traps were highly variable. Sedimentation on vegetation ranged from 10.36 to 105.56 g m^{-2} and sedimentation on traps even ranged from 4.25 to 4955.50 g m^{-2} , where some sediment traps that were heavily packed with sediment (Table 2). Descriptive statistics for C, N and P and for the explanatory variables are shown in Table 2.

Sedimentation on and underneath the vegetation

Sedimentation on the vegetation was influenced most strongly by the amount of vegetation biomass, but also by log hydrological distance and the height variation of the vegetation as well as the river kilometre ($R^2 = 0.56$, Table 3). The amount of sediment on the vegetation increased with increasing biomass ($p < 0.01$; Fig 3A) and decreased with increasing height variation of the vegetation ($p = 0.03$; Fig 3B). In addition, sedimentation on the vegetation decreased with log hydrological distance from the river ($p = 0.01$; Fig 3C), while it increased with the river kilometre ($p = 0.02$; Fig 3D).

The sedimentation on the sediment traps was driven by a single topographic variable, the log hydrological distance to the river ($R^2 = 0.42$, Table 3). Sediment traps with a short hydrological distance (close to the river) collected more sediment, and sedimentation decreased with a larger hydrological distance ($p < 0.01$, Fig 4).

Additionally, the ratio of sedimentation on the vegetation to sedimentation on the traps was driven by the hydrological distance and, the river kilometre as well as their interaction (R^2

Table 3. Model results. Final multiple regression model results of the sedimentation on the vegetation and on the traps.

	Sediment on vegetation				
	Estimate	Std. Error	t value	Pr(> t)	Sig
(Intercept)	37.3320	3.5080	10.6420	0.0000	***
River kilometre	9.5700	3.8320	2.4970	0.0231	*
log Hydrological distance	-12.0610	4.4330	-2.7210	0.0145	*
Biomass	14.4820	3.9990	3.6220	0.0021	**
Highest inflorescence I6	-6.8990	5.0780	-1.3590	0.1920	
Vertical density	7.4390	3.8380	1.9380	0.0694	.
Height variation	-9.6850	4.0560	-2.3880	0.0288	*
	Sediment on trap				
	Estimate	Std. Error	t value	Pr(> t)	Sig
(Intercept)	5.7200	0.5990	9.5490	6.83E-09	***
River kilometre	-0.7547	0.4264	-1.7700	0.0920	.
log Hydrological distance	-1.4044	0.3458	-4.0610	0.0006	***
Precipitation	-1.4622	0.8481	-1.7240	0.1001	

<https://doi.org/10.1371/journal.pone.0252694.t003>

= 0.62, S1 Table). The ratio was low with short hydrological distance, meaning that relatively more sediment settled on the traps close to the river, and decrease with increasing hydrological distance ($p < 0.01$, S3A Fig). There was also relatively more sediment on the traps at the upstream study sites, while sedimentation on the biomass relatively increased downstream the river ($p < 0.01$, S3B Fig). The interaction of river kilometre and hydrological distance was also significant ($p < 0.01$, S3B Fig), showing that with increasing river kilometre (i.e. more downstream), the relative increase of sedimentation on the vegetation is stronger with hydrological distance than at more upstream sites.

Carbon, nitrogen and phosphorus content in the sediment

Carbon, nitrogen and phosphorus content in the sediment strongly increased with the total amount of sediment on the vegetation (Fig 5) and log sediment on the traps ($p < 0.01$ for all models, S2 Table). In addition, N on the vegetation increased with vegetation biomass ($p = 0.01$) and with log hydrological distance ($p < 0.01$, $R^2 = 0.79$, Fig 5, S2 Table). Carbon and P on the vegetation additionally increased with log hydrological distance (both $p < 0.01$, $R^2 = 0.80$ and 0.92 , respectively, Fig 5, S2 Table). Carbon and N content in the sediment on the traps increased with the river kilometre (both $p = 0.02$, $R^2 = 0.64$ and 0.62 , respectively, S2 Table), while P content in the sediment on the traps was only explained by the amount of sediment on the trap ($R^2 = 0.84$, S2 Table).

The N:P ratio in the sediment on the vegetation for sites closer to the river and further away from the river did not differ significantly ($p = 0.095$). However, there was a trend towards a higher N:P ratio further away from the river. The same comparison (close and far away from the river) for the N:P ratio in the sediment on the traps showed a significantly higher N:P ratio for the sites further away from the river ($p = 0.001$).

Discussion

With this study, we disentangled *in situ* measurements of sedimentation on and underneath the vegetation on a floodplain and quantifying its relative importance in relation to topographic drivers. Biomass and height variation increase sedimentation on the vegetation, while vegetation characteristics did not explain sedimentation underneath the vegetation. The

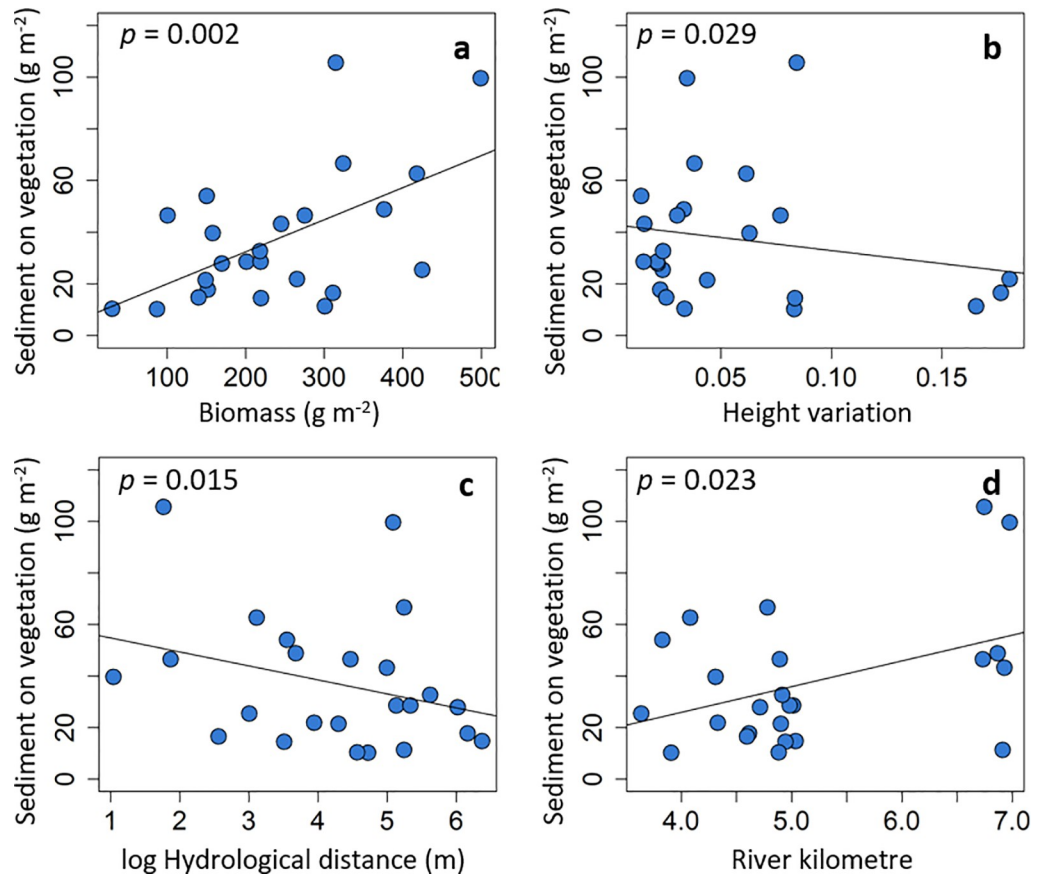


Fig 3. Sedimentation on the vegetation. Sedimentation on the vegetation explained by (a) plant biomass, (b) height variation, (c) log hydrological distance, and (d) river kilometre.

<https://doi.org/10.1371/journal.pone.0252694.g003>

hydrological distance was a key variable explaining sediment and nutrient retention on and underneath the vegetation. Carbon, N and P on the vegetation increased with hydrological distance from the river in spite of the decreasing amount of sediment with increasing hydrological distance. We could not find evidence that species diversity and leaf surface structure affect the amount of sediment and nutrient retention.

The current study was conducted on three floodplains in one river. The magnitude of the flood was relatively low, but of high frequency (biannual), so that 24 sites were inundated and thus available for the analysis. The explanatory power of the results reveals the more general importance of these factors in floodplain sedimentation processes.

Vegetation characteristics

Regarding hypothesis (H1), we found evidence that sedimentation on the vegetation increased with increasing plant biomass and decrease with height variation. More vegetation biomass is able to provide a larger surface for sediment to settle, and thus increase sedimentation on the biomass, as it was found in the flume experiments [24,26]. However, we also expected that the sedimentation on the ground underneath the vegetation would increase with increasing biomass as a consequence of a stronger reduction in flow velocity, as it was found in the flume experiment [24], but this was not supported by our findings. Three reasons might explain this: (1) it is likely that larger grain sizes (sand) accumulated underneath the vegetation, which

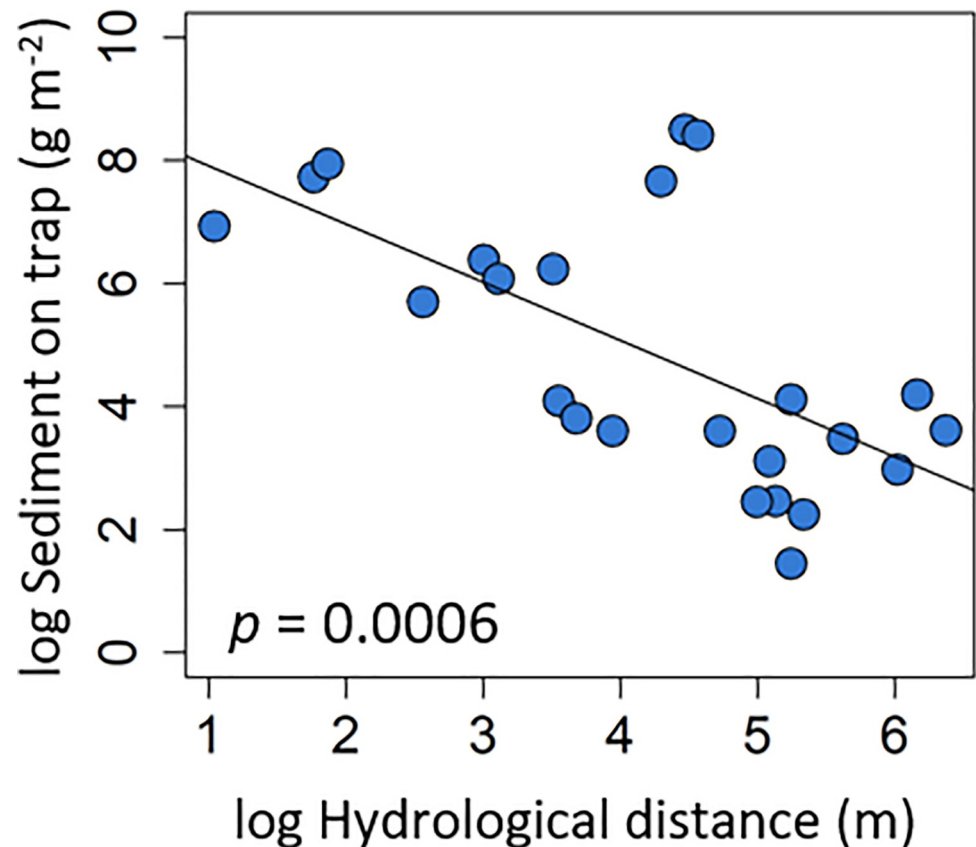


Fig 4. Sedimentation on traps. Sedimentation on traps explained by log hydrological distance.

<https://doi.org/10.1371/journal.pone.0252694.g004>

might be less affected by the biomass above; (2) the effect of the hydrological distance on the sedimentation underneath the vegetation overrides the effects of the vegetation structure; and (3) during the winter decomposition of the plant biomass started and might change the vegetation structure compared to the flume experiment conducted at the biomass peak. Other studies found positive or non-significant relations between standing biomass and trapped sediment on the ground [22,25,38]. Sediment on the vegetation surface is likely to be finer grained (silt

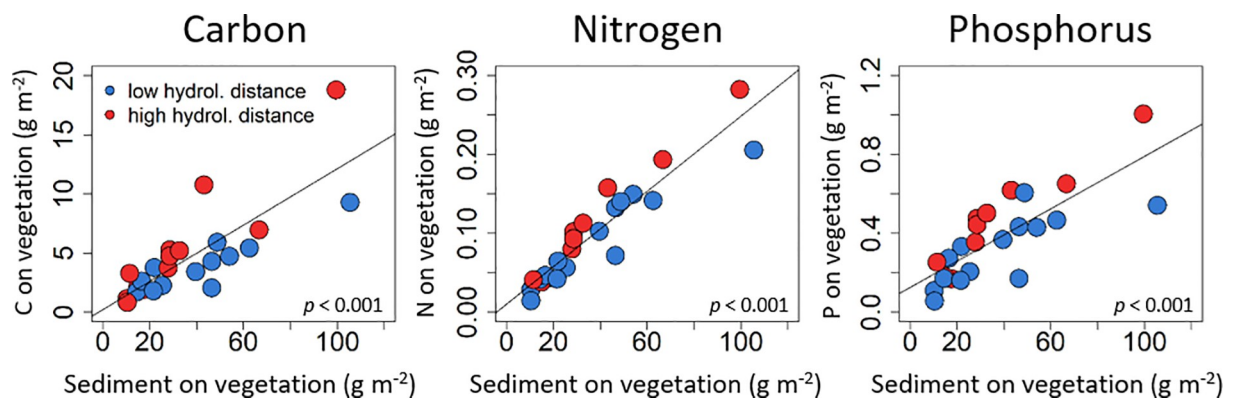


Fig 5. Nutrients on the vegetation. Carbon, nitrogen and phosphorus on the vegetation explained by the amount of sediment on the vegetation, and grouped by the mean for low and high hydrological distances from the river.

<https://doi.org/10.1371/journal.pone.0252694.g005>

and clay), since larger grains might be too heavy to be held by single leaves. Many important plant nutrients occur in or are associated with fine sediment [40,74]. Thus, this clearly shows (1) the relevant role of standing biomass for sediment retention during the flood season, and (2) emphasizes the importance of the vegetation surface for fine sedimentation and nutrient retention.

In the flume experiment it was found that density increases sedimentation on the vegetation [24], which only showed a marginally significant increase in the present study. We did not find any statistical evidence that the vegetation height explains sedimentation, but other studies did [23,24,30]. However, we found that variation of vegetation height explained sedimentation on the vegetation, even though most of the vegetation was not fully inundated. The stronger the height variation, the lower was the sediment retention on the vegetation, meaning that a more even vegetation surface collected more sediment on the vegetation. The same was found in the flume experiment [24]. Others found that the intercepted biovolume calculated by the vegetation cover times the inundation depth explained a large fraction of the sedimentation on the ground [75]. We could not measure the inundation depth (water level above the ground per plot), which we expected that it would increase the importance of the vegetation height and density.

Topography

Regarding topographic parameters, we found support for hypothesis (H2) that sedimentation on the vegetation as well as underneath the vegetation decreased with increasing hydrological distance to the river. In contrast, C, N and P on the vegetation increased with the hydrological distance.

With increasing distance from the river, the flow velocity is likely to decrease and more sediment has already settled, thereby reducing the potential sedimentation on plots with longer water paths. Even though decreasing sedimentation on and underneath the vegetation was observed with hydrological distance, the three plots farthest away from the river did not have the lowest sedimentation rates; they were more than 400 m (413–586 m) away, while all other plots were in the range of 300 m to the river. In the same three plots the sedimentation, especially underneath the vegetation, was still reasonably high (19.65–66.85 mg m⁻² [overall median 60.55 mg m⁻²]), which is in contrast with other studies that found exponential decreasing sedimentation rates on horizontal lines in the floodplain [76,77]. Also other studies found decreasing amounts of sediment with increasing straight distance from the river [27,29,40,78], with increasing flow path [42] and with decreasing hydrological connectivity [16,43,44]. Our results show the substantial role of shallow sites, such as abandoned meander and depression within the floodplain for sediment retention. We additionally found that the ratio of sedimentation on vegetation and on the traps increased with hydrological distance. Thus, our results emphasize the crucial role of vegetation for floodplain sedimentation, although the ratio needs to be interpreted with caution, since the sedimentation on the ground underneath the vegetation is represented by artificial traps, even though they are known to best represent ground surface conditions [55].

With increasing river kilometre sediment on the vegetation and C and N underneath the vegetation increased. We expected that all three study areas receive comparable amounts of sediment with respect to quality and quantity. However, it is possible that the sites further downstream (further away from the last tributary) receive less sediment with larger grain size than the ones further upstream. We also know that during the flood peak the channel width of the downstream site was at least 10% wider than the channel at the upstream site, which causes reduced flow velocity and with that might increase sedimentation. We also visually observed

lower flow velocities at the downstream site, at least for those plots close to the river, which might cause higher amounts of fine grained sediment, and thus, higher C and N retention with increasing river kilometre. For a better understanding of the key drivers, more hydraulic and hydromorphological parameters, such as discharge, inundation duration and flow velocity need to be included in the analysis [77]. Still, while results could have been different for e.g. more extreme floods, our study helps to improve our general understanding of the mechanisms and processes causing sedimentation on floodplains.

Carbon, nitrogen and phosphorus on the vegetation

Our results further support the hypothesis (H5) that nutrients (C, N and P) in the sediment increased with the amount of sediment. In addition to that, this study shows that C, N and P on the vegetation increased with greater hydrological distance. Thus, we observed relatively more nutrients on the vegetation far away from the river even though there is less total amount of sediment. Carbon and P are commonly bound to fine grained sediment, while nitrogen is only partially associated with sediment, but it still follows similar distribution patterns [40,79]. Thus, we can derive that the vegetation primarily captures finer sediment fraction (silt, clay, and organic material), which probably also decreases in size with distance from the river, but has more nutrients bound to it [80]. With this result, our study emphasized again the crucial role of shallow sites far inside the floodplain, such as abandoned meander and depression, for fine sediment and nutrient retention during floods.

In addition, we found an increasing N:P ratio for sites further away from the river. These changes in elemental ratios provided evidence of changes in the nutrient composition of the sediment with distance to the river main channel. A higher N:P ratio indicated a higher N availability compared to P, which suggests that N is relatively more limiting for plant growth close to the river channel, and that P is relatively more limiting for plant growth further away from the river main channel. Subsequent mineralization processes could provide additional nutrient sources for plant growth and stimulate nutrient uptake in terrestrial parts of the floodplain, as well as it might also affect community composition due to changed availability of plant nutrients [81].

Diversity and leaf surface structure

We did not find any evidence for our hypotheses regarding species diversity (H3). The flume experiment also only showed effects of species richness on sedimentation, when species identity effects were not considered [26]. Similarly, others did not find any significant differences in sediment capture capacity between monocultures and a three-species mixture in an experiment [37]. Nevertheless, it is known that species diversity can correlate with vegetation structure [34], and in the flume experiment it was found that structural diversity increase sedimentation on patches [24]. From grassland experiments we know that more diverse vegetation is denser and taller than low diverse vegetation [35,36].

We also did not find evidence for the importance of the leaf pubescence and leaf area in this study (H4), even though in previous studies both have been found to represent relevant traits for sedimentation [38,39]. Three reasons might explain that: (1) Pubescent species were rather poorly represented within our floodplain (five species with a cover mean of 6.9%), so that we had limited statistical power to test for its potential effects. (2) Including stem density and mean number of leaves per individual seems likely to allow a more precise estimation of the pubescence and the leaf area effect at the plot level [38]. (3) Especially for leaf pubescence the seasonality of the flood could be relevant, since decomposition processes might already have diminished the leaf hairs.

Outlook

Overall we would suggest some further research to improve our understanding of the mechanisms. First, a monitoring of similar floods of the same magnitude to validate our findings. Second, the monitoring of a larger flood, which would have the potential to introduce the effects of inundation depth and vegetation deformation. Third, increasing the size, length or number of sediment traps per plot to ensure to capture enough sediment to additionally investigate particle size.

Conclusion

With our *in situ* measurements, we improve the understanding of sediment and nutrient retention in floodplains by providing insights on the vegetation structure besides the floodplain topography and simultaneously disentangling sedimentation on and underneath the vegetation. Our findings will help at the small scale to improve existing model approaches to predict sediment and nutrient retention on floodplains [12,20,50,82]. Notably, we found that more biomass increases sediment and nutrient retention on the vegetation. Sedimentation decreases with hydrological distance to the river, even though it is still reasonably high beyond distances of 400 m. Nutrients (C, N, and P) in the sediment on the vegetation, however, increase with distance to the river. Based on the results about sediment and nutrient retention, we can recommend the following management practices: First, reduced mowing for more standing vegetation biomass during the flood season would increase sediment and nutrient retention via the total biomass. Especially, for nutrient retention, this counts for shallow areas with high hydrological distance to the river. The present study shows, that the mowing regime might be less important, if the focus is on maximal sediment retention, which on a mass basis happens more strongly underneath the vegetation without clear effects of the vegetation structure. Of course, trade-offs between sediment retention and other management goals, such as biodiversity conservation, should be taken into consideration when making decisions about floodplain management. Second, the strong importance of the topographical variable 'hydrological distance' for sediment and nutrient retention emphasizes the high value of laterally connected river-floodplain systems, including long abandoned meanders and depressions, which also has recently been found for nutrient retention in other studies [83,84]. Thus, our study suggests (1) an improvement of lateral connectivity to be able to use the potential retention hotspots far inside the floodplain, and in accordance with that (2) an adapted mowing regime on the floodplain to achieve the management regarding sediment and nutrient retention, and therefore the ecosystem function of water purification of the river. Overall, our findings are transferable in a mechanistic perspective. We are not able to predict any quantitative sedimentation for other floodplains, floods or river catchments, but the mechanistic approach allows to tackle the key characteristics that in general cause the process of sedimentation on floodplain meadows.

Supporting information

S1 Fig. Structural image. a) Original image with blue background wall and blue flooring material in front. b) Automatically analyzed images for the following predictors as explained in Table 1: Vertical density = percent of vegetation pixels on the image of standard size, Mean height = mean height of vegetation pixels on the image, Median height = median height of vegetation pixels on the image, and Height variation = standard deviation of vegetation pixel height on the image.

(TIF)

S2 Fig. Sediment traps. Picture of a sediment trap in the field.
(TIF)

S3 Fig. Sedimentation ratio. Ratio of sediment on vegetation to sediment on traps.
(TIF)

S1 Table. Model results. Statistical model results of the ratio sediment on the vegetation to sediment on the traps.
(PDF)

S2 Table. Model results. Statistical model results of carbon, nitrogen and phosphorus on the vegetation and on the traps.
(PDF)

S1 File.
(DOCX)

S2 File.
(DOCX)

Acknowledgments

We thank the TRY-initiative—a global database of plant traits and all contributors for providing the data. We also thank our student assistants (Antonia Ludwig and Georg Rieland) for supporting the fieldwork and Katie Barry for English proof reading.

Author Contributions

Conceptualization: Lena Kretz, Carolin Seele-Dilbat, Christian Wirth.

Formal analysis: Lena Kretz, Ronny Richter, Fons van der Plas.

Funding acquisition: Christian Wirth.

Investigation: Lena Kretz.

Methodology: Lena Kretz, Carolin Seele-Dilbat, Fons van der Plas, Christian Wirth.

Project administration: Christian Wirth.

Resources: Elisabeth Bondar-Kunze, Christiane Schulz-Zunkel, Michael Vieweg.

Writing – original draft: Lena Kretz.

Writing – review & editing: Elisabeth Bondar-Kunze, Thomas Hein, Ronny Richter, Christiane Schulz-Zunkel, Carolin Seele-Dilbat, Fons van der Plas, Michael Vieweg, Christian Wirth.

References

1. Hunter HM, Walton RS. Land-use effects on fluxes of suspended sediment, nitrogen and phosphorus from a river catchment of the Great Barrier Reef, Australia. *J Hydrol.* 2008; 356(1–2):131–46.
2. Sharma P, Rai SC. Streamflow, sediment and carbon transport from a Himalayan watershed. *J Hydrol.* 2004; 289(1–4):190–203.
3. Tockner K, Pusch M, Borchardt D, Lorang MS. Multiple stressors in coupled river-floodplain ecosystems. *Freshw Biol.* 2010; 55:135–51.
4. Turnpenny AWH, Williams R. Effects of sedimentation on the gravels of an industrial river system. *J Fish Biol.* 1980; 17(6):681–93.

5. Carpenter SR, Caraco NF, Correll DL, Howarth RW, Sharpley AN, Smith VH. Nonpoint pollution of surface waters with phosphorus and nitrogen. *Ecol Appl.* 1998; 8(3):559–68.
6. Habersack H, Hein T, Stanica A, Liska I, Mair R, Jäger E, et al. Challenges of river basin management: Current status of, and prospects for, the River Danube from a river engineering perspective. *Sci Total Environ.* 2016; 543:828–45. <https://doi.org/10.1016/j.scitotenv.2015.10.123> PMID: 26589137
7. Naiman RJ, Décamps H. The ecology of interfaces: Riparian zones. *Annu Rev Ecol Syst.* 1997; 28:621–58.
8. Noe GB, Hupp CR. Carbon, nitrogen, and phosphorus accumulation in floodplains of Atlantic Coastal Plain rivers, USA. *Ecol Appl.* 2005; 15(4):1178–90.
9. Conte M, Ennaanay D, Mendoza G, Walter MT, Wolny S, Freyberg D, et al. Retention of nutrients and sediment by vegetation. In: Kareiva P, Tallis H, Ricketts TH, Daily GC, Polasky S, editors. *Natural Capital: Theory and Practice of Mapping Ecosystem Services.* Oxford: Oxford University Press; 2011.
10. Hopkins KG, Noe GB, Franco F, Pindilli EJ, Gordon S, Metes MJ, et al. A method to quantify and value floodplain sediment and nutrient retention ecosystem services. *J Environ Manage.* 2018; 220(December 2017):65–76. <https://doi.org/10.1016/j.jenvman.2018.05.013> PMID: 29758400
11. Noe GB, Hupp CR. Retention of riverine sediment and nutrient loads by coastal plain floodplains. *Ecosystems.* 2009; 12(5):728–46.
12. Manners R, Schmidt J, Wheaton JM. Multiscalar model for the determination of spatially explicit riparian vegetation roughness. *J Geophys Res Earth Surf.* 2013; 118(1):65–83.
13. Olde Venterink H, Vermaat JE, Pronk M, Wiegman F, van der Lee GEM, van der Hoorn MW, et al. Importance of sediment deposition and denitrification for nutrient retention in floodplain wetlands. *Appl Veg Sci.* 2006; 9(2):163–74.
14. Steiger J, Tabacchi E, Dufour S, Corenblit D, Peiry JL. Hydrogeomorphic processes affecting riparian habitat within alluvial channel-floodplain river systems: A review for the temperate zone. *River Res Appl.* 2005; 21(7):719–37.
15. Tockner K, Stanford JA. Riverine flood plains: Present state and future trends. *Environ Conserv.* 2002; 29:308–30.
16. Wolf KL, Noe GB, Ahn C. Hydrologic connectivity to streams increases nitrogen and phosphorus inputs and cycling in soils of created and natural floodplain wetlands. *J Environ Qual.* 2013; 42(4):1245–55. <https://doi.org/10.2134/jeq2012.0466> PMID: 24216376
17. Wohl E, Bledsoe BP, Jacobson RB, Poff NL, Rathburn SL, Walters DM, et al. The natural sediment regime in rivers: Broadening the foundation for ecosystem management. *Bioscience.* 2015; 65(4):358–71.
18. Corenblit D, Baas ACW, Bornette G, Darrozes J, Delmotte S, Francis RA, et al. Feedbacks between geomorphology and biota controlling Earth surface processes and landforms: A review of foundation concepts and current understandings. *Earth-Science Rev.* 2011; 106(3–4):307–31.
19. Gurnell AM, Bertoldi W, Corenblit D. Changing river channels: The roles of hydrological processes, plants and pioneer fluvial landforms in humid temperate, mixed load, gravel bed rivers. *Earth-Science Rev.* 2012; 111(1–2):129–41.
20. Camporeale C, Perucca E, Ridolfi L, Gurnell AM. Modeling the interaction between river morphodynamics and riparian vegetation. *Rev Geophys.* 2013; 51(2012):379–414.
21. Steiger J, Gurnell AM. Spatial hydrogeomorphological influences on sediment and nutrient deposition in riparian zones: Observations from the Garonne River, France. *Geomorphology.* 2002; 49:1–23.
22. Elliott AH. Settling of fine sediment in a channel with emergent vegetation. *J Hydraul Eng.* 2000; 126(8):570–7.
23. Corenblit D, Steiger J, Gurnell AM, Naiman RJ. Plants intertwine fluvial landform dynamics with ecological succession and natural selection: A niche construction perspective for riparian systems. *Glob Ecol Biogeogr.* 2009; 18(4):507–20.
24. Kretz L, Koll K, Seele-Dilbat C, van der Plas F, Weigelt A, Wirth C. Vegetation structure alters fine sediment retention on and underneath herbaceous vegetation in a flume experiment. *PLoS One.* 2021;1–16.
25. Burylo M, Rey F, Bochet E, Dutoit T. Plant functional traits and species ability for sediment retention during concentrated flow erosion. *Plant Soil.* 2012; 353(1–2):135–44.
26. Kretz L, Koll K, Seele-Dilbat C, van der Plas F, Weigelt A, Wirth C. Effects of plant species identity overrides diversity effects in explaining sedimentation within vegetation in a flume experiment. *Int Rev Hydrobiol.* 2021.
27. Västilä K, Järvelä J. Characterizing natural riparian vegetation for modeling of flow and suspended sediment transport. *J Soils Sediments.* 2018; 18(10):3114–30.

28. Akram S, Yu B, Ghadiri H, Rose C, Hussein J. The links between water profile, net deposition and erosion in the design and performance of stiff grass hedges. *J Hydrol.* 2014; 510:472–9.
29. Asselman NEM, Middelkoop H. Floodplain sedimentation: Quantities, patterns and processes. *Earth Surf Process Landforms.* 1995; 20(1 1995):481–99.
30. Västilä K, Järvelä J, Koivuor H. Flow-vegetation-sediment interaction in a cohesive compound channel. *J Hydraul Eng.* 2016; 142(1).
31. Nilsson C, Berggren K. Alterations of riparian ecosystems caused by river regulation. *Bioscience.* 2000; 50(9):783.
32. Olson DM, Dinerstein E. The global 200: A representation approach to conserving the earth's most biologically valuable ecoregions. *Conserv Biol.* 1998; 12(3):502–15.
33. Tilman D, Isbell F, Cowles JM. Biodiversity and ecosystem functioning. *Annu Rev Ecol Evol Syst.* 2014; 45:471–93.
34. Proulx R, Roca IT, Cuadra FS, Seiferling I, Wirth C. A novel photographic approach for monitoring the structural heterogeneity and diversity of grassland ecosystems. *J Plant Ecol.* 2014; 7(6):518–25.
35. Lorentzen S, Roscher C, Schumacher J, Schulze ED, Schmid B. Species richness and identity affect the use of aboveground space in experimental grasslands. *Perspect Plant Ecol Evol Syst.* 2008; 10(2):73–87.
36. Wacker L, Baudois O, Eichenberger-Glinz S, Schmid B. Effects of plant species richness on stand structure and productivity. *J Plant Ecol.* 2009; 2(2):95–106.
37. Kervroëdan L, Armand R, Saunier M, Faucon MP. Effects of plant traits and their divergence on runoff and sediment retention in herbaceous vegetation. *Plant Soil.* 2019;511–24.
38. Kervroëdan L, Armand R, Saunier M, Ouvry J, Faucon M. Plant functional trait effects on runoff to design herbaceous hedges for soil erosion control. *Ecol Eng.* 2018; 118(December 2017):143–51.
39. Kretz L, Seele C, van der Plas F, Weigelt A, Wirth C. Leaf area and pubescence drive sedimentation on leaf surfaces during flooding. *Oecologia.* 2020; 193(3):535–45. <https://doi.org/10.1007/s00442-020-04664-2> PMID: 32419047
40. Sival FP, Makaske B, Maas GJ, Runhaar J. Floodplain sedimentation regulating vegetation productivity on small rivers? In: *Research for Managing Rivers: Present and Future Issues.* 2005. p. 82–3.
41. Hupp CR. Hydrology, geomorphology and vegetation of costal plain rivers in the south-eastern USA. *Hydrol Process.* 2000; 14(August 1999):2991–3010.
42. Middelkoop H, Van der Perk M. Modelling spatial patterns of overbank sedimentation on embanked floodplains. *Geogr Ann.* 1998; 80:95–109.
43. Tockner K, Pennetzdorfer D, Reiner N, Schiemer F, Ward JV. Hydrological connectivity, and the exchange of organic matter and nutrients in a dynamic river-floodplain system (Danube, Austria). *Freshw Biol.* 1999; 41(3):521–35.
44. Rodríguez M, Steiger J, Rosales J, Laraque A, López JL, Castellanos B, et al. Multi-annual contemporary flood event overbank sedimentation within the vegetated lower Orinoco floodplain, Venezuela. *River Res Appl.* 2019; 35(8):1241–56.
45. Jeffries R, Darby SE, Sear DA. The influence of vegetation and organic debris on flood-plain sediment dynamics: Case study of a low-order stream in the New Forest, England. *Geomorphology.* 2003; 51(1–3):61–80.
46. Scown MW, Thoms MC, De Jager NR. Floodplain complexity and surface metrics: Influences of scale and geomorphology. *Geomorphology.* 2015; 245:102–16.
47. Paul M, Rupperecht F, Möller I, Bouma TJ, Spencer T, Kudella M, et al. Plant stiffness and biomass as drivers for drag forces under extreme wave loading: A flume study on mimics. *Coast Eng.* 2016; 117:70–8.
48. Järvelä J. Flow resistance of flexible and stiff vegetation: A flume study with natural plants. *J Hydrol.* 2002; 269:44–54.
49. Ortiz AC, Ashton A, Nepf H. Mean and turbulent velocity fields near rigid and flexible plants and the implications for deposition. *J Geophys Res Earth Surf.* 2013; 118(4):2585–99.
50. Natho S, Venohr M. Active versus potential floodplains—The effect of small flood events on nutrient retention along the river Elbe corridor (Germany). *Aquat Sci.* 2014; 76(4):633–42.
51. Magurran AE. *Ecological diversity and its measurement.* Press PU, editor. Princeton; 1988.
52. R Core Team. *R: A language and environment for statistical computing.* R Foundation for Statistical Computing. Vienna, Austria; 2020.

53. Aureliano Netto AF, Nogueira Martins R, Aquino de Souza GS, Araújo GDM, Hatum de Almeida SL, Agnolette Capelini V. Segmentation of Rgb Images Using Different Vegetation Indices and Thresholding Methods. *Nativa*. 2018; 6(4):389.
54. Pau G, Fuchs F, Sklyar O. EBImage—an R package for image processing with applications to cellular phenotypes. *Bioinformatics*. 2010; 26(7):979–81. <https://doi.org/10.1093/bioinformatics/btq046> PMID: 20338898
55. Steiger J, Gurnell AM, Goodson JM. Quantifying and characterizing contemporary riparian sedimentation. *River Res Appl*. 2003; 19(4):335–52.
56. Brodie CR, Leng MJ, Casford JSL, Kendrick CP, Lloyd JM, Yongqiang Z, et al. Evidence for bias in C and N concentrations and $\delta^{13}\text{C}$ composition of terrestrial and aquatic organic materials due to pre-analysis acid preparation methods. *Chem Geol*. 2011; 282(3–4):67–83.
57. Schwanghart W, Scherler D. Short Communication: TopoToolbox 2—MATLAB-based software for topographic analysis and modeling in Earth surface sciences. *Earth Surf Dyn*. 2014; 2(1):1–7.
58. MathWorks. MATLAB and Statistics Toolbox Release, Inc., Natick, Massachusetts, United States; 2016.
59. Kattge J, Bönsch G, Diaz S. TRY plant trait database—enhanced coverage and open access. *Glob Chang Biol*. 2020; 26(Data requested between 06th of August and 27th of September 2019.):119–188. <https://doi.org/10.1111/gcb.14904> PMID: 31891233
60. Campetella G, Botta-Dukát Z, Wellstein C, Canullo R, Gatto S, Chelli S, et al. Patterns of plant trait-environment relationships along a forest succession chronosequence. *Agric Ecosyst Environ*. 2011; 145(1):38–48.
61. Dalke IV, Novakovskiy AB, Maslova SP, Dubrovskiy YA. Morphological and functional traits of herbaceous plants with different functional types in the European Northeast. *Plant Ecol*. 2018; 219(11):1295–305.
62. Price CA, Enquist BJ. Scaling mass and morphology in leaves: An extension of the wbe model. *Ecology*. 2007; 88(5):1132–41. <https://doi.org/10.1890/06-1158> PMID: 17536400
63. Wright IJ, Dong N, Maire V, Prentice IC, Westoby M, Díaz S, et al. Global climatic drivers of leaf size. *Science (80-)*. 2017; 357(6354):917–21. <https://doi.org/10.1126/science.aal4760> PMID: 28860384
64. Fitter AH, Peat HJ. The Ecological Flora Database. *J Ecol*. 1994; 82(2):415–25.
65. Herz K, Dietz S, Haider S, Jandt U, Scheel D, Bruelheide H. Predicting individual plant performance in grasslands. *Ecol Evol*. 2017;(April):8958–65. <https://doi.org/10.1002/ece3.3393> PMID: 29177035
66. Herz K, Dietz S, Haider S, Jandt U, Scheel D, Bruelheide H. Drivers of intraspecific trait variation of grass and forb species in German meadows and pastures. *J Veg Sci*. 2017; 28(4):705–16.
67. Kleyer M, Bekker RM, Knevel IC, Bakker JP, Thompson K, Sonnenschein M, et al. The LEDA Traitbase: A database of life-history traits of the Northwest European flora. *J Ecol*. 2008; 96(6):1266–74.
68. Laughlin DC, Leppert JJ, Moore MM, Sieg CH. A multi-trait test of the leaf-height-seed plant strategy scheme with 133 species from a pine forest flora. *Funct Ecol*. 2010; 24(3):493–501.
69. Lhotsky B, Cseceserits A, Kovacs B, Botta-Dukat Z. New plant trait records of the hungarian flora. *Acta Bot Hung*. 2016; 58(3–4):397–400.
70. Milla R, Reich PB. Multi-trait interactions, not phylogeny, fine-tune leaf size reduction with increasing altitude. *Ann Bot*. 2011; 107(3):455–65. <https://doi.org/10.1093/aob/mcq261> PMID: 21199835
71. Ordoñez JC, Van Bodegom PM, Witte J-PM, Bartholomeus RP, Van Hal JR, Aerts R. Plant strategies in relation to resource supply in mesic to wet environments: Does theory mirror nature? *Am Nat*. 2010; 175(2):225–39. <https://doi.org/10.1086/649582> PMID: 20030563
72. Fox J, Weisberg S. An {R} companion to applied regression. Third. Sage: Thousand Oaks CA; 2019.
73. Venables WN, Ripley BD. Modern applied statistics with S. Fourth. New York: Springer; 2002.
74. Liu X, Zhang X, Zhang M. Major factors influencing the efficacy of vegetated buffers on sediment trapping: A review and analysis. *J Environ Qual*. 2008; 37(5):1667–74. <https://doi.org/10.2134/jeq2007.0437> PMID: 18689727
75. Corenblit D, Steiger J, Gurnell AM, Tabacchi E, Roques L. Control of sediment dynamics by vegetation as a key function driving biogeomorphic succession within fluvial corridors. *Earth Surf Process Landforms*. 2009; 34:1790–810.
76. Walling DE, He Q. The spatial variability of overbank sedimentation on river floodplains. *Geomorphology*. 1998; 24(2–3):209–23.
77. Steiger J, Gurnell AM, Petts GE. Sediment deposition along the channel margins of a reach of the middle River Severn, UK. *Regul Rivers-Research Manag*. 2001; 17:443–60.

78. Klaus VH, Sintermann J, Kleinebecker T, Hölzel N. Sedimentation-induced eutrophication in large river floodplains—An obstacle to restoration? *Biol Conserv.* 2011; 144(1):451–8.
79. Olde Venterink H, Wiegman F, Van der Lee GEM, Vermaat JE. Role of active floodplains for nutrient retention in the River Rhine. *J Environ Qual.* 2003; 32(4):1430–5. <https://doi.org/10.2134/jeq2003.1430> PMID: 12931899
80. River M, Richardson CJ. Particle size distribution predicts particulate phosphorus removal. *Ambio.* 2018; 47(s1):124–33. <https://doi.org/10.1007/s13280-017-0981-z> PMID: 29164536
81. Antheunisse AM, Loeb R, Lamers LPM, Verhoeven JTA. Regional differences in nutrient limitation in floodplains of selected European rivers: Implications for rehabilitation of characteristic floodplain vegetation. *River Res Appl.* 2006; 22:1039–55.
82. James LA. Ten conceptual models of large-scale legacy sedimentation—A review. *Geomorphology.* 2018; 317:199–217.
83. Schulz-Zunkel C, Baborowski M, Ehlert T, Kasperidus HD, Krüger F, Horchler P, et al. Simple modelling for a large-scale assessment of total phosphorus retention in the floodplains of large rivers. *Wetlands.* 2021; 41(6). <https://doi.org/10.1007/s13157-021-01463-0> PMID: 34393321
84. Natho S, Tschikof M, Bondar-Kunze E, Hein T. Modeling the effect of enhanced lateral connectivity on nutrient retention capacity in large river floodplains: How much connected floodplain do we need? *Front Environ Sci.* 2020; 8(June).