



## Journal Article

# Numerical modeling of plant root controls on gravel bed river morphodynamics

**Author(s):**

Caponi, Francesco; Siviglia, Annunziato

**Publication Date:**

2018-09-16

**Permanent Link:**

<https://doi.org/10.3929/ethz-b-000293837> →

**Originally published in:**

Geophysical Research Letters 45(17), <http://doi.org/10.1029/2018GL078696> →

**Rights / License:**

[In Copyright - Non-Commercial Use Permitted](#) →

This page was generated automatically upon download from the [ETH Zurich Research Collection](#). For more information please consult the [Terms of use](#).

1 **Numerical modeling of plant-root controls on gravel-bed river**  
2 **morphodynamics**

3 **F. Caponi <sup>1</sup> and A. Sivilgia <sup>1</sup>**

4 <sup>1</sup>Laboratory of Hydraulics, Hydrology and Glaciology, ETH Zurich, Switzerland

5 **Key Points:**

- 6 • A model accounting for the biogeomorphic feedbacks between plant roots and  
7 riverbed morphodynamics is presented
- 8 • Uprooting is the primary plant-root biogeomorphic feedback controlling the co-  
9 evolution of gravel-bed river morphodynamics and vegetation
- 10 • The competition between the potential flow erosion and the uprooting depth medi-  
11 ates plant-root controls on riverbed morphodynamics

## Abstract

The role of vegetation in shaping the geomorphology of rivers, deltas, along with tidal and estuarine environments is widely recognized. While mutual interactions between flow, plant canopy and morphodynamics have been extensively investigated, similar studies considering plant roots are limited. Here, we present results from numerical model that quantify the feedbacks of both the above- and below-ground vegetation on gravel-bed river (GBR) morphodynamics. Plant-root biogeomorphic feedbacks, i.e. uprooting and root-enhanced riverbed cohesion, are quantified through the description of the vertical root distribution. By investigating the evolution of the riverbed of a straight gravel channel with a vegetated patch, we show that uprooting is the primary plant-root biogeomorphic feedback determining the evolution of the riverbed and the competing influence of the potential flow erosion versus uprooting depth mediates the plant-root controls on morphodynamics. These findings broaden our understanding on the role played by plant roots on GBR morphodynamics.

## 1 Introduction

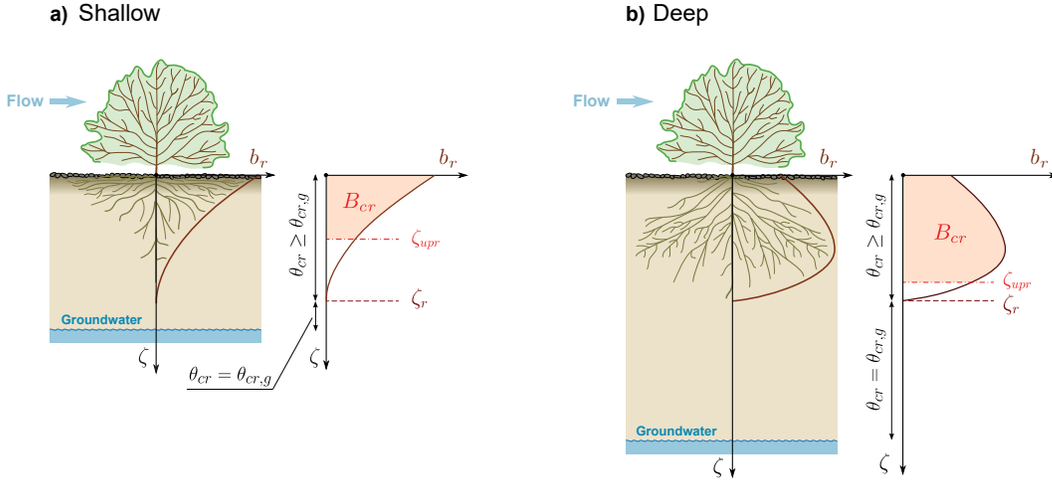
The role of vegetation in shaping the geomorphology of interfaces between water and land surfaces, such as river bars and floodplain, river deltas, along with tidal and estuarine environments, is widely recognized [Corenblit *et al.*, 2015]. Mutual interactions among riparian vegetation, water flow and sediment transport result in a series of biogeomorphic feedbacks [sensu Corenblit *et al.*, 2007] that can affect bar and landform formation in vegetated rivers [e.g. Gurnell, 2014; Bertoldi *et al.*, 2011], determine shifts among alternate stable states [Bertoldi *et al.*, 2014; Bertagni *et al.*, 2018], shape river deltaic marshes [Nardin and Edmonds, 2014] and promote formation of drainage channel networks in tidal systems in the presence of marshes [e.g. Temmerman *et al.*, 2007; Schwarz *et al.*, 2018]. Consequently, development of eco-morphodynamic numerical models [Murray and Paola, 2003; Bertoldi *et al.*, 2014; Oorschot *et al.*, 2016], which quantify such feedbacks, is crucial for predicting the morphodynamics of these areas and for planning sustainable restoration and flood mitigation measures [Wohl *et al.*, 2015].

Although the general importance of vegetation is widely recognized, its precise role in mediating biogeomorphic feedbacks in rivers is not clear. A number of studies indicate that the emergence and strength of vegetation-related feedbacks result from the balance between physical and biological processes [e.g. Corenblit *et al.*, 2007; Tal and Paola, 2007].

44 Modification of sediment supply rates has been suggested as a mechanism responsible for  
45 muting the effects of species-specific plant traits on morphodynamics of sand-bed rivers  
46 [*Manners et al.*, 2015; *Diehl et al.*, 2017] and for altering the vegetation's effects on chan-  
47 nel dynamics in GBRs [*Gran et al.*, 2015]. Changes in the hydrological regime, including  
48 flood frequency and magnitude [*Vesipa et al.*, 2017], as well as water table fluctuations,  
49 have been argued to impact biogeomorphic succession and river channel morphodynamics  
50 [*Bätz et al.*, 2016; *Bertoldi et al.*, 2011; *Bertagni et al.*, 2018]. Subsurface flows and alter-  
51 ation of pore-water pressures in the hyporheic zone may also contribute to mediate the ef-  
52 fects of vegetation on cohesive riverbeds [e.g. *Simon and Collison*, 2001; *Cancienne et al.*,  
53 2008]. Among these processes, survival of riparian vegetation is significantly threatened  
54 by morphological changes, which cause uprooting and scour, limiting plant community  
55 expansion [*Gurnell et al.*, 2012].

56 Most studies examining biogeomorphic feedbacks consider only the above-ground  
57 component of vegetation. Plant canopy, for instance, is known to change turbulence struc-  
58 ture [*Nepf*, 2012] and to significantly increase flow resistance [e.g. *Västilä and Järvelä*,  
59 2014; *Aberle and Järvelä*, 2015]. The reduction of bottom shear stresses in vegetated areas  
60 alters sediment transport, thereby inducing local and reach-scale riverbed changes [*Vargas-*  
61 *Luna et al.*, 2015; *Le Bouteiller and Venditti*, 2015]. However, below-ground vegetation un-  
62 derpins fundamental biogeomorphic feedbacks that are often not included in these studies.  
63 Plant roots contribute to mediate riverbank cohesion and stability, shaping river planform  
64 styles [e.g. *Tal and Paola*, 2010; *Pollen-Bankhead and Simon*, 2010; *Davies and Gibling*,  
65 2011; *Gibling and Davies*, 2012; *Polvi et al.*, 2014], promoting in-channel sediment stabi-  
66 lization and reducing scour [*Pasquale et al.*, 2012; *Pasquale and Perona*, 2014]. Roots pro-  
67 vide resistance to the drag forces exerted by the flow on plant canopy, delaying or possibly  
68 avoiding uprooting [*Edmaier et al.*, 2011, 2015; *Perona and Crouzy*, 2018]. The amount  
69 of roots that anchor plants is of the utmost importance for determining the ability of veg-  
70 etation to withstand erosional events [*Bywater-Reyes et al.*, 2015; *Bankhead et al.*, 2017].  
71 Nonetheless, GBR morphodynamic models mainly describe the effects of roots on vege-  
72 tation anchoring and scouring, adopting lumped approaches that are oversimplified [e.g.  
73 *Murray and Paola*, 2003; *Bertoldi et al.*, 2014].

74 Our goals are to present a simple modeling framework to study key biogeomorphic  
75 feedbacks of plant root and to show the results of model runs that test the importance of  
76 these feedbacks in predicting GBR morphology. We consider vegetation consisting of an



87 **Figure 1.** Vertical root density distributions  $b_r$ , (a) shallow and (b) deep, used to probe the role of uproot-  
 88 ing and root-enhanced riverbed cohesion on the evolution of a GBR.  $B_{cr}$  is the fraction of the entire root  
 89 biomass  $B_r$  that must be exposed to the flow before uprooting occurs,  $\zeta_r$  the rooting depth, i.e. the depth to  
 90 which the roots grow, and  $\zeta_{upr}$  the uprooting depth. Further symbols are reported in the main text.

77 above- and below-ground component. We adopt the stochastic model developed by *Tron*  
 78 *et al.* [2014] and characterize the plant roots by their vertical density distribution, which  
 79 depends on water table dynamics. Then, we model plant-root biogeomorphic feedbacks  
 80 depending on these distributions. This approach allows us to disentangle the role of the  
 81 two vegetation components and to explore how plant root morphology influences biogeo-  
 82 morphic feedbacks. In this study, we examine a simplified GBR morphology, while retain-  
 83 ing the key morphodynamic processes. We investigate the riverbed response to a vegeta-  
 84 tion patch in a straight gravel channel by varying hydromorphological configurations and  
 85 vegetation characteristics.

## 86 2 Modeling Framework

### 91 2.1 Hydromorphodynamics

92 Hydromorphodynamic processes are simulated with the one-dimensional model  
 93 BASEMENT [*Vetsch et al.*, 2017]. Firstly, the hydrodynamic problem is solved by inte-  
 94 grating numerically the Saint-Venant equations and using the Manning-Strickler approach  
 95 for the evaluation of the global flow resistance, whereby the total shear stress is evaluated

96 as

$$\tau = \frac{\rho g u |u|}{K_s^2 h^{1/3}} \quad , \quad (1)$$

98 where  $\rho$  is the water density,  $g$  is the gravitational acceleration,  $u$  is the vertically aver-  
 99 aged flow velocity,  $h$  the water depth and  $K_s$  the Strickler coefficient. Secondly, the Exner  
 100 equation is adopted to describe the time evolution of a cohesionless GBR composed of a  
 101 uniform sediment. It reads:

$$(1 - p) \frac{\partial z_b}{\partial t} + \frac{\partial q_b}{\partial x} = 0 \quad , \quad (2)$$

103 where  $z_b$  is the bed elevation,  $p$  is the sediment porosity and  $q_b$  is the longitudinal bed-  
 104 load flux.  $q_b$  is evaluated as a function of the excess of the Shields shear stress,  $\theta$ , above  
 105 a threshold value  $\theta_{cr}$ , where

$$\theta = \frac{\tau}{(\rho_s - \rho) g d_s} \quad (3)$$

107 and  $\rho_s$  and  $d_s$  are the sediment density and diameter, respectively.

## 108 2.2 Plant Roots

109 Plant roots often display complex architectures [Gregory, 2008], with a maximum  
 110 depth that is mostly limited by groundwater [Fan *et al.*, 2017] and a density that decreases  
 111 with riverbed depth [Jackson *et al.*, 1996]. In riparian ecosystems, however, root growth  
 112 tends to follow water table oscillations [Orellana *et al.*, 2012]. This is particularly relevant  
 113 in GBRs where the large hydraulic conductivity in the hyporheic zone enhances exchanges  
 114 between groundwater and stream flow [Cardenas *et al.*, 2004].

115 To describe the vertical root distribution, we adopt the stochastic model proposed by  
 116 Tron *et al.* [2014], which describes root dynamics driven by water table oscillations. The  
 117 model assumes that roots grow within an optimal zone whose fluctuations follow the wa-  
 118 ter table oscillations, while roots decay otherwise (Figure S1). This zone results from the  
 119 optimal balance between the amount of pore water available for root uptake and dissolved  
 120 oxygen levels needed for root respiration [Gregory, 2008]. The maximum rooting depth  
 121 is limited by the minimum depth reached by this optimal zone (see more details on the  
 122 physical processes underlying the root model in the supporting information).

123 By considering water table fluctuations as a stochastic process [Ridolfi *et al.*, 2011],  
 124 the model produces a probability density distribution (PDF) of the root density,  $b_r$ , over  
 125 the riverbed depth,  $\zeta$ , (downward oriented axis with origin at the riverbed, see Figures 1a,  
 126 1b and S1). This PDF depends on physically-based parameters that define the water table

127 oscillations (characterized by a mean oscillation depth, frequency and decay rate) and the  
 128 plant root characteristics (see details on the mathematical formulation in the supporting  
 129 information). In this study, we describe shallow root profiles (Figure 1a) that result from  
 130 shallow and more variable water table oscillations and deep root profiles (Figure 1b) char-  
 131 acterized by a deep and more stable water table [Tron *et al.*, 2014, 2015].

## 132 **2.3 Biogeomorphic Feedbacks**

### 133 **2.3.1 Canopy feedback on flow resistance and sediment transport**

134 The presence of plant canopy increases the global flow resistance by increasing local  
 135 roughness, modifying flow patterns and providing additional drag [Nepf, 2012]. The addi-  
 136 tional drag varies significantly with morphology and bio-mechanical properties of canopy  
 137 [Aberle and Järvelä, 2015], including stem density, flexibility, presence and type of foliage  
 138 and submerged and emergent conditions [e.g. Västilä and Järvelä, 2014]. In line with pre-  
 139 vious models, which are based on a depth-averaged description of the flow [e.g. Bertoldi  
 140 *et al.*, 2014], we model the global flow resistance (equation 1 to be used for hydrodynamic  
 141 computation) by considering a single Strickler coefficient  $K_{s,v}$  [Kim *et al.*, 2012; Bertoldi  
 142 *et al.*, 2014; Le Bouteiller and Venditti, 2015] that incorporates not only the shear stress  
 143 exerted by the fluid directly on the sediment grain (bottom shear stress), but also the addi-  
 144 tional drag generated by vegetation.

145 Flow pattern changes associated to the presence of vegetation have also profound  
 146 effects on sediment transport [e.g. Yager and Schmeeckle, 2013; Le Bouteiller and Ven-  
 147 ditti, 2015]. Bottom shear stress is reduced in a plant patch, and the decrease is higher for  
 148 denser vegetation [Le Bouteiller and Venditti, 2015] and larger plant frontal areas [Vargas-  
 149 Luna *et al.*, 2015]. Since direct quantification of the bottom shear stress is extremely dif-  
 150 ficult in the presence of vegetation [Le Bouteiller and Venditti, 2015], we model the re-  
 151 duction of bottom shear stress by multiplying the total shear stress  $\tau$  by a factor  $\gamma \leq 1$   
 152 [Le Bouteiller and Venditti, 2015] and compute the sediment flux,  $q_b$ , using the reduced  
 153 Shields stress,  $\gamma\theta$ .

### 154 **2.3.2 Plant-root feedback on riverbed cohesion**

155 Buried roots are known to significantly modify mechanical and biochemical prop-  
 156 erties of riverbed thereby reducing erosion on riverbanks and slope surfaces [Vannoppen

157 *et al.*, 2015]. Studies assessing the reduced root-riverbed erosion in cohesive substrates of-  
 158 ten indicate a negative exponential relation between root density and the bed shear stress  
 159 needed to mobilize sediments. However, this relation might not hold for cohesionless  
 160 substrate, such as gravel, because of the different particle detachment mechanism [*Politti*  
 161 *et al.*, 2018]. Alternatively, we can use a linear relation between root density ( $b_r$ ) and the  
 162 critical Shields parameter ( $\theta_{cr}$ ), as indicated by *Pasquale and Perona* [2014] for GBRs.  
 163 We assume that at the riverbed depth  $\zeta$

$$164 \theta_{cr}(\zeta) = \theta_{cr,g} + (\theta_{cr,v} - \theta_{cr,g})b_r(\zeta), \quad (4)$$

165 where  $\theta_{cr,g}$  and  $\theta_{cr,v}$  ( $> \theta_{cr,g}$ ) represent the threshold values for incipient sediment mo-  
 166 tion on bare and vegetated riverbed, respectively [*Bertoldi et al.*, 2014].

### 167 2.3.3 Uprooting

168 Plant removal by uprooting depends on the balance between drag forces of the wa-  
 169 ter flow acting on the above-ground part of vegetation and resisting forces provided by  
 170 the buried part of the roots [*Edmaier et al.*, 2011]. Resisting forces increase with rooting  
 171 depth [*Edmaier et al.*, 2015; *Bywater-Reyes et al.*, 2015] and the maximum density depth  
 172 [*Pasquale et al.*, 2012], most likely exceeding applied drag forces. Vegetation that devel-  
 173 ops substantial root biomass is, in fact, unlikely to be uprooted by drag forces alone even  
 174 at high flows [*Bywater-Reyes et al.*, 2015; *Bankhead et al.*, 2017]. Uprooting rather oc-  
 175 curs as a consequence of riverbed erosion that gradually exposes part of the roots to the  
 176 flow thus reducing the anchoring resistance of the plant (Type II uprooting as defined by  
 177 *Edmaier et al.* [2011]). Experimental evidence suggests that exposure of only part of the  
 178 entire root biomass might be sufficient to uproot plants [*Edmaier et al.*, 2015]. In light of  
 179 this evidence, we define a critical biomass value,  $B_{cr}$ , as the fraction  $\beta$  of the entire root  
 180 biomass (Figure 1) that must be exposed to the flow before uprooting occurs. We calculate  
 181 this value as follows:

$$182 B_{cr} = \beta \int_0^{\zeta_r} b_r(z) dz = \int_0^{\zeta_{upr}} b_r(z) dz, \quad (5)$$

183 and we assume that uprooting occurs when riverbed scouring reaches the uprooting  
 184 depth  $\zeta_{upr}$ .  $\zeta_{upr}$  increases with the rooting depth (Figure S3) and depends on the value  
 185 of  $\beta$  (Figure S2) and the root distribution (Figures 1a and 1b). Finally, when uprooting  
 186 occurs  $b_r$  is set to 0 and  $K_{s,v}$  to the Strickler value assigned to bare riverbed,  $K_{s,g}$ .

## 2.4 Numerical Simulations

Simulations are conducted under steady flow (constant discharge  $Q = 100 \text{ m}^3/\text{s}$ ) in a straight rectangular channel of length  $L = 1500 \text{ m}$ , 10 m wide and initial constant slope  $S_0$ . The bed is composed of uniform sediments ( $p = 0.4$ , reference grain size  $d_s = 20 \text{ mm}$ ) and the bedload flux  $q_b$  is calculated using the Meyer-Peter and Müller formula [Meyer-Peter et al., 1948] ( $\theta_{cr,g} = 0.047$ ). A vegetation patch of length  $L_{veg} = 500 \text{ m}$ , comprising above- and below-ground vegetation, is placed between the coordinates  $x_{up} = 600 \text{ m}$  and  $x_{dw} = 1100 \text{ m}$  (Figure 2a) covering the entire channel width. We design this configuration to represent reach-scale morphodynamics of a vegetated gravel bar in a simplified way, as similarly done by previous experimental studies on sand-bed substrates [e.g. Manners et al., 2015; Diehl et al., 2017; Le Bouteiller and Venditti, 2014; Le Bouteiller and Venditti, 2015]. We mimic different types of vegetation depending on their impact on global flow resistance ( $K_{s,v}$  in Table 1), while for the bare riverbed we use  $K_{s,g} = 30 \text{ m}^{1/3} \text{ s}^{-1}$ . The bottom stress reduction coefficient,  $\gamma$ , has been chosen within the range reported in Vargas-Luna et al. [2015]. Model runs are grouped into five sets based on the type of roots characterizing the patch. The control set *NR* has no roots. In sets *SR1* and *SR2*, we consider shallow- (Figure 1a) and deep-root distributions (Figure 1b) with different rooting depths, respectively. These root configurations are typically observed in field and laboratory experiments with cuttings and juvenile riparian vegetation growing in GBRs [e.g. Pasquale et al., 2012; Gorla et al., 2015]. The parameter values used in the root model are reported in the supporting information. For all simulations we set  $\beta = 0.9$  in equation 5.

The numerical domain consists of cross-sections that are spaced out evenly ( $2 \text{ m}$ ) and vertical root distributions discretized by using riverbed layers of  $0.1 \text{ m}$ . For all runs, initial conditions are obtained by setting uniform flow conditions both at the inlet and the outlet of the numerical domain and by running fixed-bed simulation until the steady state is reached. Different water surface profiles are obtained for different values of  $K_{s,v}$  and  $S_0$ . We then perform morphodynamic simulations for the five sets until riverbed equilibrium is reached (at  $t = T_{eq}$ ), keeping  $z_b$  fixed both at the inlet and outlet. The key parameter settings and configurations considered are summarized in Table 1.

**Table 1.** Model parameters defining numerical runs<sup>a</sup>

Run	$S_0$ [-]	$F_r$ [-]	$K_{s,v}$ [ $m^{1/3}s^{-1}$ ]	$\gamma$ [-]	$\theta_{cr,v}$ [m]	$S_{eq}$ [-]	$E_{eq}$ [m]	$\zeta_{upr}$ [m]	$\zeta_r$ [m]	Plant root type
NR-EP1	0.005	0.8	25	0.69	$\theta_{cr,g}$	0.0063	0.31	-	-	No roots
NR-EP2	0.005	0.8	20	0.44	$\theta_{cr,g}$	0.008	0.75	-	-	No roots
NR-EP3	0.02	1.5	25	0.69	$\theta_{cr,g}$	0.025	1	-	-	No roots
NR-EP4	0.005	0.8	15	0.25	$\theta_{cr,g}$	0.01	1.3	-	-	No roots
NR-EP5	0.02	1.5	15	0.25	$\theta_{cr,g}$	0.039	3.9	-	-	No roots
SR1-EP1	0.005	0.8	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	0.31	0.45	0.6	Shallow
SR1-EP2	0.005	0.8	20	0.44	$[\theta_{cr,g},0.1,0.2]$	-	0.75	0.45	0.6	Shallow
SR1-EP3	0.02	1.5	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	1	0.45	0.6	Shallow
SR1-EP4	0.005	0.8	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	1.3	0.45	0.6	Shallow
SR1-EP5	0.02	1.5	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	3.9	0.45	0.6	Shallow
SR2-EP1	0.005	0.8	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	0.31	0.55	0.8	Shallow
SR2-EP2	0.005	0.8	20	0.44	$[\theta_{cr,g},0.1,0.2]$	-	0.75	0.55	0.8	Shallow
SR2-EP3	0.02	1.5	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	1	0.55	0.8	Shallow
SR2-EP4	0.005	0.8	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	1.3	0.55	0.8	Shallow
SR2-EP5	0.02	1.5	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	3.9	0.55	0.8	Shallow
DR3-EP1	0.005	0.8	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	0.31	0.75	0.8	Deep
DR3-EP2	0.005	0.8	20	0.44	$[\theta_{cr,g},0.1,0.2]$	-	0.75	0.75	0.8	Deep
DR3-EP3	0.02	1.5	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	1	0.75	0.8	Deep
DR3-EP4	0.005	0.8	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	1.3	0.75	0.8	Deep
DR3-EP5	0.02	1.5	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	3.9	0.75	0.8	Deep
DR4-EP1	0.005	0.8	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	0.31	0.95	1	Deep
DR4-EP2	0.005	0.8	20	0.44	$[\theta_{cr,g},0.1,0.2]$	-	0.75	0.95	1	Deep
DR4-EP3	0.02	1.5	25	0.69	$[\theta_{cr,g},0.1,0.2]$	-	1	0.95	1	Deep
DR4-EP4	0.005	0.8	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	1.3	0.95	1	Deep
DR4-EP5	0.02	1.5	15	0.25	$[\theta_{cr,g},0.1,0.2]$	-	3.9	0.95	1	Deep

<sup>a</sup>  $S_0$  = initial riverbed slope,  $S_{eq}$  = equilibrium riverbed slope (only for runs NR),  $F_r$  = Froude number of the uniform flow,  $E_{eq}$  = erosion potential,  $K_{s,v}$  = Strickler friction coefficient incorporating the effect of the drag generated by the vegetation  $\gamma$  = bottom stress reduction coefficient,  $\theta_{cr,v}$  = critical Shields parameter incorporating the increase of cohesion due to the presence of roots,  $\zeta_{upr}$  = riverbed depth at which uprooting occurs and  $\zeta_r$  = rooting depth.

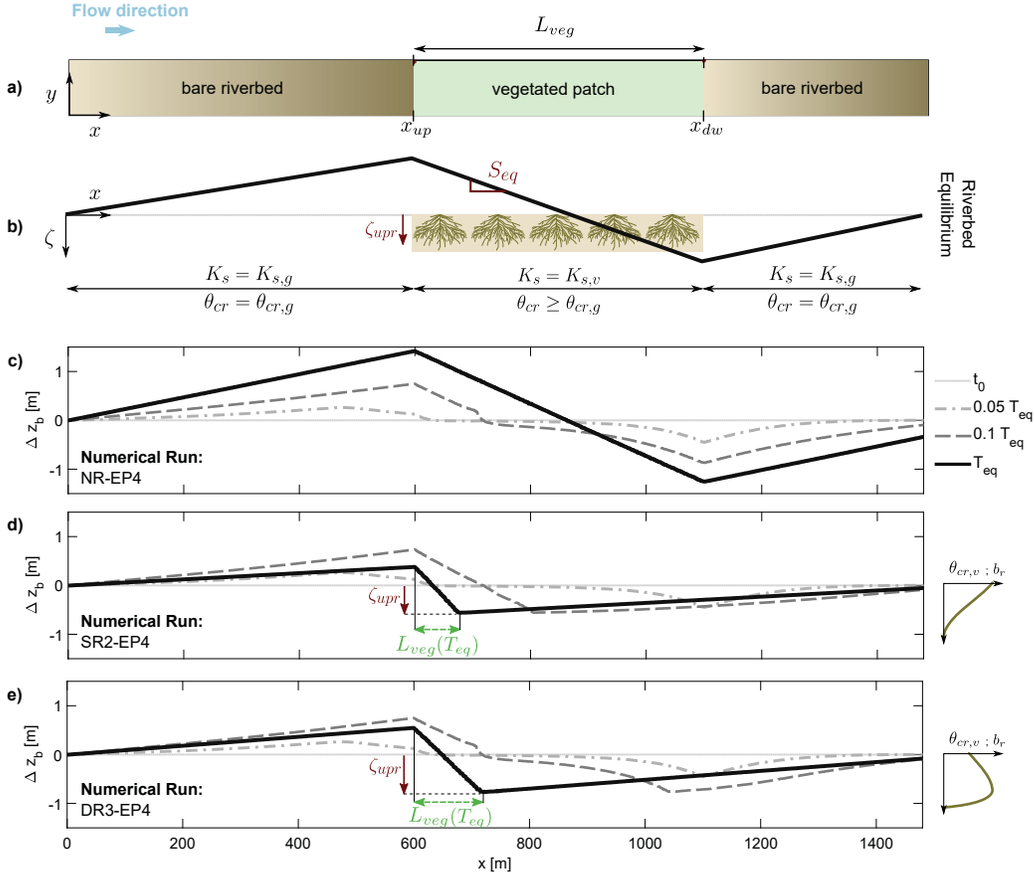
### 3 Results

#### 3.1 The Role of Above-ground Vegetation

Numerical results from runs with no roots, in which we varied the roughness of the above-ground vegetation and the flow characteristics (set NR in Table 1), quantify the riverbed changes due to the interactions between flow and above-ground biomass. The final riverbed equilibrium (Figure 2b), which is common to all runs NR (Table 1), is characterized by an increased riverbed slope within the vegetated patch, a deposition in the upstream part, and a scour at  $x_{dw}$  (Figure 2b). Riverbed evolution starts with a deposition process from upstream the patch and erosion downstream ( $t=0.05 T_{eq}$ ) (dash-dotted line in Figure 2c). Over time, while deposition advances downstream within the patch, erosion proceeds upstream ( $t=0.1 T_{eq}$ ) (dashed lines in Figure 2c), progressively increasing the riverbed slope across the whole vegetated patch. The maximum bed level changes in our experiments occur at the interface between the vegetated and bare riverbed (at  $x = \{x_{up}, x_{dw}\}$ , Figure 2c). Riverbed steepening in vegetated areas has been previously reported [Le Bouteiller and Venditti, 2015]. Such configurations are a direct consequence of the friction exerted by the vegetation and the consequent reduction of the bed shear stress. Simulated flow velocities and bed shear stresses are reduced within and upstream from the vegetated patch. In addition, vegetation obstructs the flow increasing flow velocity and bed shear stress downstream from the patch. This, in turn, reduces the sediment transport capacity,  $q_b$ , within the vegetated patch, generating a sediment transport imbalance throughout the channel [Le Bouteiller and Venditti, 2014]. At equilibrium, the slope within the patch,  $S_{eq}$  (Table 1 and Figure 2b), depends on the difference between  $K_{s,v}$  and  $K_{s,g}$  and on flow characteristics (measured through the Froude number). We measure the strength of the erosion process, resulting from the interaction between flow and above-ground vegetation, through the *erosion potential*  $E_{eq}$ , i.e. the maximum scour at  $x = x_{dw}$ . Numerical runs of set NR give five different values of  $E_{eq}$  which are reported, in increasing order from EP1 to EP5, in Table 1.

#### 3.2 The Role of Uprooting

Uprooting should play a fundamental role in the riverbed's response to erosion processes and deep roots by offering more anchoring resistance than shallow roots to erosional events. In addition, a sufficiently intense erosional event could completely remove

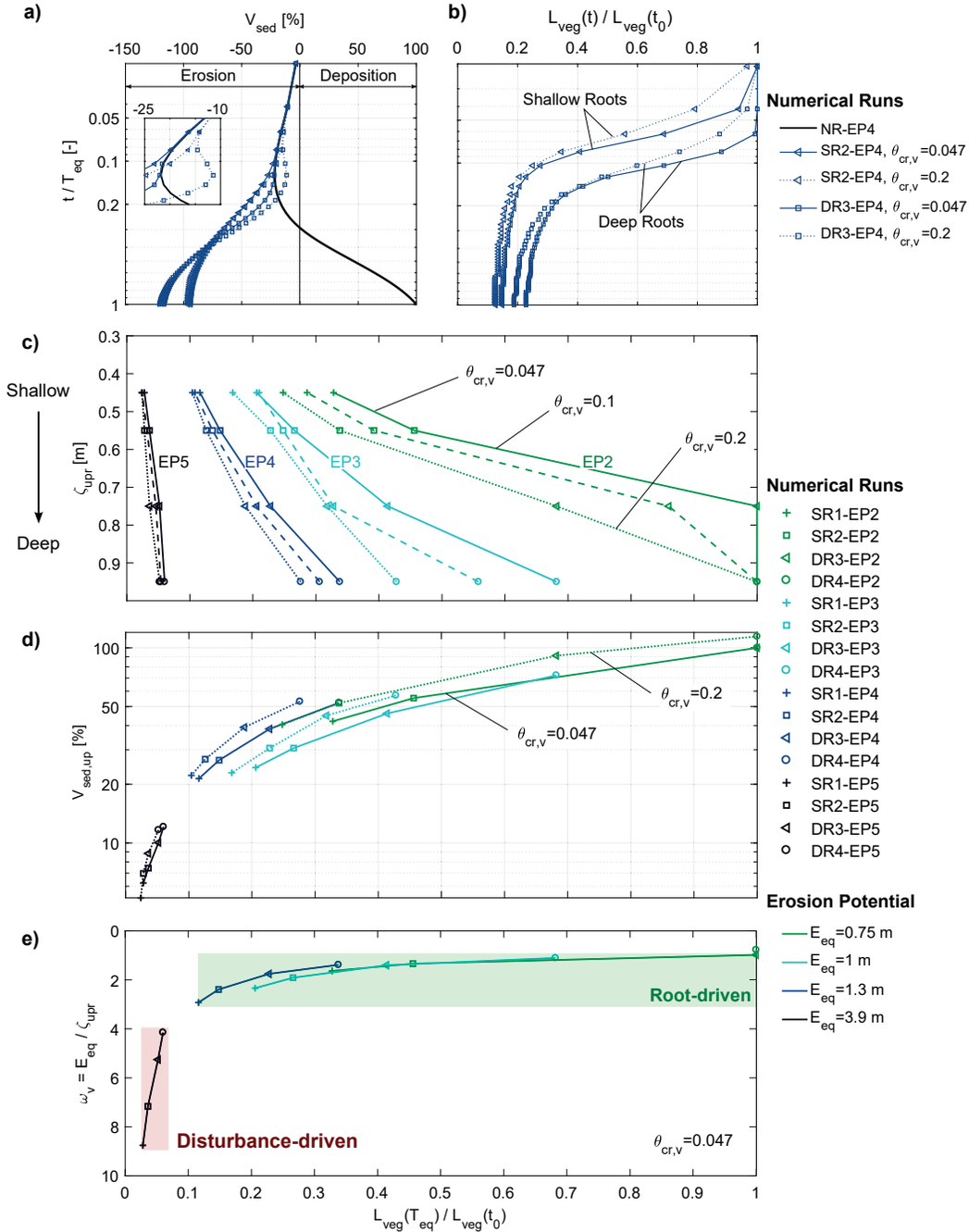


244 **Figure 2.** Effect of plant roots on riverbed evolution of a straight gravel channel with a vegetated patch  
 245 characterized by the same above-ground vegetation and different vertical root distributions. Schematic illus-  
 246 tration of (a) the channel used for the simulations and (b) bed level changes at equilibrium (black-thick line).  
 247 Time evolution of the riverbed, evaluated in terms of bed level changes ( $\Delta z_b(x, t) = z_b(x, t) - z_b(x, t_0)$ ),  
 248 for the case with (c) no roots (run NR-EP4), (d) shallow (run SR2-EP4) and (e) deep (run SR2-EP4) root  
 249 distributions. In (d) and (e)  $\zeta_{upr}$  is the uprooting depth and  $L_{veg}(T_{eq})$  the patch length at equilibrium.

254 vegetation, regardless of the root distribution type. To test these hypotheses, we perform  
 255 simulations of the sets SR and DR assuming  $\theta_{cr,v} = \theta_{cr,g}$  (see Table 1) and leaving out  
 256 the root-enhanced riverbed cohesion effect. Despite the presence of shallow (run SR2-  
 257 EP4) or deep (run DR3-EP4) roots, bed evolution follows the same dynamics as the case  
 258 with no roots (NR-EP4, Figure 2c) as long as the erosion at  $x_{dw}$  is smaller than  $\zeta_{upr}$   
 259 (compare solutions at  $t=0.05 T_{eq}$  in Figures 2d-e). If riverbed changes exceed  $\zeta_{upr}$ , up-  
 260 rooting triggers an upward cascade-like mechanism that converts part of the vegetated  
 261 patch to bare riverbed (Figures 2d-e). The erosion process stops when sediment balance  
 262 is reached throughout the entire channel.

263 For a given erosion potential (EP4 in Figures 3a and 3b), uprooting leads to shorter  
 264 patch lengths that are higher for shallow root profiles (runs SR2-EP4 and DR3-EP4 in  
 265 Figure 3b). The time-to-uprooting is shorter for shallow roots (Figure 3b), since smaller  
 266 scour depths, and therefore shorter times, are required to remove vegetation with shallow  
 267 roots. In order to characterize the cumulative riverbed dynamics, in Figure 3a, we plot the  
 268 time evolution of the integral of the normalized net eroded/deposited volume throughout  
 269 the whole channel ( $V_{sed}$ ). Compared to the case with no roots (run NR-EP4), the reduc-  
 270 tion of the patch length causes significant morphological changes while small differences  
 271 are observed at equilibrium between shallow and deep root distributions.

272 The ratio between  $L_{veg}$  reached at equilibrium and its initial value ( $L_{veg}(t_0)$ ) is  
 273 shown for all the runs from series EP2 to EP5 in Figure 3c. Each line corresponds to runs  
 274 characterized by the same erosion potential but different root distributions and therefore  
 275 different  $\zeta_{upr}$ . Results show that, for a given  $\zeta_{upr}$ , the residual biomass (i.e. length of  
 276 the patch at equilibrium) decreases as the strength of the erosion process increases (from  
 277 EP2 to EP5). Shallow roots (small  $\zeta_{upr}$ ) (sets SR1 and SR2) lead to a residual biomass  
 278 smaller than 0.4, regardless of the strength of the erosion process, whereas the length  
 279 of deep rooted patches (sets DR3 and DR4) ranges between 0.05 and 1. Moreover, large  
 280 values of the erosion potential result in shorter vegetated patches, smaller differences be-  
 281 tween shallow and deep roots (Figure 3c) and smaller deposition  $V_{sed,up}$  upstream from  
 282 the patch (black solid line in Figure 3d). As an example, in runs EP5, the vegetated patch  
 283 is reduced by about 95% and  $V_{sed,up}$  remains around 10% regardless of the root distribu-  
 284 tion type.



285 **Figure 3.** Plant-root controls on GBR morphodynamics. Time evolution of (a) normalized net  
 286 eroded/deposited volume along the channel,  $V_{sed}$  (%), and (b) normalized length of the vegetated patch  
 287 for shallow (SR2-EP4) and deep (DR3-EP4) roots. (c) Uprooting depth vs residual biomass. (d) Upstream  
 288 normalized deposited volume,  $V_{sed,up}$  (%), versus residual biomass. (e) Influence of the relative strength  
 289 of erosion process,  $\omega_v$  (equation 6), on the residual biomass. In all plots, solid lines refer to  $\theta_{cr,v} = \theta_{cr,g}$ ,  
 290 while dashed and dotted lines to  $\theta_{cr,v} = 0.1$  and  $0.2$ , respectively. For a given run, the normalized net  
 291 eroded/deposited volume through the whole channel is calculated as  $V_{sed} = \int_0^L \Delta z_b(t, x) / \Delta z_b^{NR}(t =$   
 292  $T_{eq}, x) dx$ , where  $\Delta z_b^{NR}$  refers to the same run with no roots.  $V_{sed,up}$  is calculated as  $V_{sed}$  but in the range  
 293  $x \in [0, x_{up}]$ .

### 3.3 The Role of Root-enhanced Riverbed Cohesion

We investigate the effect of the root-enhanced riverbed cohesion by running simulations of sets SR and DR and considering  $\theta_{cr,v} > \theta_{cr,g}$  (see Table 1). Root-enhanced cohesion affects riverbed dynamics only slightly (compare dashed and dotted lines vs solid lines in Figures 3a, 3c and 3d). Furthermore, it decreases the time-to-uprooting (see an example in Figure 3b), which reduces the length of the vegetated patch at equilibrium (Figure 3c). Such reduction is larger for deep roots and smaller for large values of  $E_{eq}$  (Figure 3c) and can be explained as follows. The added cohesion, i.e. higher critical shear stress, reduces sediment mobility and further decreases the transport capacity within the patch, which was already diminished by the above-ground effect of the vegetated patch. This causes a larger difference in transport capacity and hence an increase of the scour at the interface between the vegetated patch and the bare riverbed (see Figure S4 and Table S1), which in turn favors uprooting.

## 4 Discussion and Implications

Our modeling study demonstrates that the competition between the potential flow erosion and the uprooting depth mediates plant-root controls on GBR morphodynamics. The results of numerical runs show that the strength of the erosion process is primarily the result of the interactions between flow and above-ground vegetation, while vegetation anchoring resistance depends on the vertical root distribution. This competition can be measured by introducing the nondimensional parameter  $\omega_v$  (a similar parameter has been previously used by *Perona and Crouzy* [2018]) defined as:

$$\omega_v = \frac{E_{eq}}{\zeta_{upr}} = \frac{\text{Strength of Erosion}}{\text{Vegetation Anchoring Resistance}} \quad , \quad (6)$$

where the strength of the erosion process is measured through the erosion potential,  $E_{eq}$ , representing the maximum erosion occurring in absence of any resisting force. Conversely, the resistance opposed by vegetation to uprooting is measured through the depth  $\zeta_{upr}$ . If we plot  $\omega_v$  against the normalized vegetated patch length, two different regions can be identified in Figure 3e: disturbance-driven and root-driven. Our results indicate that for  $1 < \omega_v < 4$  (root-driven region) riverbed dynamics greatly depends on  $\omega_v$ , whereas, for  $\omega_v > 4$  (disturbance-driven region), changes in  $\omega_v$  only slightly influence the length of the vegetated patch. The extension of these two regions is primarily controlled by the uprooting mechanism and is marginally affected by the root-enhanced riverbed cohesion.

325 Changes in the erosion rate might also influence the threshold values defining these re-  
 326 gions [Perona and Crouzy, 2018]. In analogy to the nondimensional parameter,  $T^*$ , pro-  
 327 posed by Tal and Paola [2007], defined as the ratio between characteristic timescales of  
 328 riverbed reworking and vegetation encroachment,  $\omega_v$  can provide information about mor-  
 329 phological trajectories and may offers insights on the role played by the above- and below-  
 330 ground vegetation components. For instance, in the disturbance-driven region, where ero-  
 331 sional processes dominate, the riverbed will likely evolve towards a configuration with low  
 332 vegetation cover and high sediment mobility, marginally affected by the root distribution.  
 333 On the contrary, in the root-driven region ( $E_{eq} \sim \zeta_{upr}$ ), erosion and plant anchoring resis-  
 334 tance are balanced, producing conditions more suitable for vegetation development. In this  
 335 region, the type of root distribution (deep or shallow), as well as plant canopy characteris-  
 336 tics, can play a fundamental role in mediating the evolution of GBRs. Deep groundwater  
 337 environments could favor the development of deep roots (Figure 1b) [Bertoldi et al., 2011;  
 338 Bätz et al., 2016], which would enhance vegetation resistance to uprooting [Bywater-Reyes  
 339 et al., 2015]. Whereas, shallow and highly variable water tables (Figure 1a) may limit  
 340 rooting depths and favor vegetation less resistant to erosional events [Tron et al., 2014;  
 341 Pasquale et al., 2012]. The novelty of this model is the ability to take into account dif-  
 342 ferent environmental conditions, such as changes in water table dynamics, that can help  
 343 interpreting observed vegetation-morphology dynamics [Bertoldi et al., 2011; Bätz et al.,  
 344 2016].

345 In this study we explore simplified conditions to reduce the inherent complexity of  
 346 the problem and disentangle the contribution of each feedback, independently. We exam-  
 347 ine the morphodynamics of a straight cohesionless gravel channel covered by a vegetation  
 348 patch. This configuration does not target to capture vegetation dynamics and the associ-  
 349 ated development of fluvial landform over time [Gurnell, 2014], but rather to investigate  
 350 the underlying processes occurring in GBRs at the event scale. However, the condition an-  
 351 alyzed simplifies the topography of a real river gravel bar. The modeling framework we  
 352 developed can be easily extended to take into account both flow unsteadiness and more  
 353 complex morphologies, such as alternate bar patterns [Serlet et al., 2018]. Moreover, we  
 354 could investigate erosion processes related to changes in sediment supply rate [Gran et al.,  
 355 2015; Diehl et al., 2017] and bar migration [Bertoldi et al., 2014], which are not consid-  
 356 ered here. These processes might change the strength of erosion (i.e.  $E_{eq}$ ) and thus the  
 357 value of  $\omega_v$ , possibly shifting a system from one region to another of Figure 3e. We use

358 a rather simple treatment of canopy effect on flow and sediment transport. For instance,  
359 local effects on scour and deposition pattern resulting from alteration of turbulence struc-  
360 tures around vegetation patches can not be captured by the model [e.g. *Kim et al.*, 2015].  
361 However, the model qualitatively captures the key features of riverbed adjustment in pres-  
362 ence of vegetated patch observed in laboratory experiments [*Le Bouteiller and Venditti*,  
363 2014; *Diehl et al.*, 2017].

364 The present results can have significant implications on the prediction of the co-  
365 evolution between vegetation and river morphology. Firstly, they suggest that a detailed  
366 description of uprooting in eco-morphodynamic models [*Solari et al.*, 2016] is a key ingre-  
367 dient needed for quantification. This is crucial for determining the effect of flood events  
368 on vegetation survival and development and on planning sustainable strategies for river  
369 restoration projects [e.g. *Bankhead et al.*, 2017; *Vesipa et al.*, 2017]. Secondly, model re-  
370 sults suggest that further investigations linking plant roots, groundwater and river mor-  
371 phology are necessary. The proposed modeling framework can be extended to include  
372 above- and below-ground vegetation dynamics to predict morphological trajectories in re-  
373 lation to changes in water table dynamics. How vegetation allocates biomass to its above-  
374 and below-ground component might play a fundamental role on mediating biogeomorphic  
375 feedbacks. Finally, further investigations should examine the role of natural stochasticity of  
376 uprooting [*Perona and Crouzy*, 2018], which could be introduced by a stochastic represen-  
377 tation of the critical root biomass (defined by  $\beta$  in our model).

## 378 **Acknowledgments**

379 We wish to thank W. Bertoldi for helpful discussions and for commenting the manuscript.  
380 The support of the Swiss National Science Foundation (Grant No. 159813) is gratefully  
381 acknowledged. We would like also to thank the Editor V. Yvanov and two anonymous re-  
382 viewers for their constructive comments that greatly improved the manuscript. The numer-  
383 ical model BASEMENT can be freely downloaded at <http://www.basement.ethz.ch>. The  
384 results of numerical simulations used in the paper can be found at <https://polybox.ethz.ch/index.php/s/dcewDQq6KHz36jj>.

## 385 **References**

386 Aberle, J., and J. Järvelä (2015), Hydrodynamics of vegetated channels, in *Rivers–*  
387 *Physical, Fluvial and Environmental Processes*, pp. 519–541, Springer.

- 388 Bankhead, N. L., R. E. Thomas, and A. Simon (2017), A combined field, laboratory and  
389 numerical study of the forces applied to, and the potential for removal of, bar top veg-  
390 etation in a braided river, *Earth Surface Processes and Landforms*, 42(3), 439–459, doi:  
391 10.1002/esp.3997.
- 392 Bätz, N., P. Colombini, P. Cherubini, and S. N. Lane (2016), Groundwater controls on  
393 biogeomorphic succession and river channel morphodynamics, *Journal of Geophysical*  
394 *Research F: Earth Surface*, 121(10), 1763–1785, doi:10.1002/2016JF004009.
- 395 Bertagni, M., C. Camporeale, and P. Perona (2018), Parametric transitions between bare  
396 and vegetated states in water-driven patterns, *Proceedings of the National Academy of*  
397 *Sciences*, doi:10.1073/pnas.1721765115.
- 398 Bertoldi, W., N. A. Drake, and A. M. Gurnell (2011), Interactions between river flows and  
399 colonizing vegetation on a braided river: exploring spatial and temporal dynamics in  
400 riparian vegetation cover using satellite data, *Earth Surface Processes and Landforms*,  
401 36(11), 1474–1486, doi:10.1002/esp.2166.
- 402 Bertoldi, W., A. Siviglia, S. Tettamanti, M. Toffolon, D. Vetsch, and S. Francalanci  
403 (2014), Modeling vegetation controls on fluvial morphological trajectories, *Geophys-  
404 ical Research Letters*, 41(20), 7167–7175, doi:10.1002/2014GL061666.
- 405 Bywater-Reyes, S., Andrew C. Wilcox, J. C. Stella, and A. F. Lightbody (2015), Flow and  
406 scour constraints on uprooting of pioneer woody seedlings, *Water Resources Research*,  
407 pp. 2760–2772, doi:10.1002/2014WR016632.Received.
- 408 Cancienne, R. M., G. A. Fox, and S. Andrew (2008), Influence of seepage undercutting  
409 on the stability of root-reinforced streambanks, *Earth Surface Processes and Landforms*,  
410 33(11), 1769–1786, doi:10.1002/esp.1657.
- 411 Cardenas, M. B., J. Wilson, and V. A. Zlotnik (2004), Impact of heterogeneity, bed forms,  
412 and stream curvature on subchannel hyporheic exchange, *Water Resources Research*,  
413 40(8).
- 414 Corenblit, D., E. Tabacchi, J. Steiger, and A. M. Gurnell (2007), Reciprocal interactions  
415 and adjustments between fluvial landforms and vegetation dynamics in river corridors:  
416 A review of complementary approaches, *Earth-Science Reviews*, 84(1-2), 56–86, doi:  
417 10.1016/j.earscirev.2007.05.004.
- 418 Corenblit, D., A. Baas, T. Balke, T. Bouma, F. Fromard, V. Garófano-Gómez,  
419 E. González, A. M. Gurnell, B. Hortobágyi, F. Julien, et al. (2015), Engineer pioneer  
420 plants respond to and affect geomorphic constraints similarly along water–terrestrial in-

- 421 terfaces world-wide, *Global Ecology and Biogeography*, 24(12), 1363–1376.
- 422 Davies, N. S., and M. R. Gibling (2011), Evolution of fixed-channel alluvial plains  
423 in response to Carboniferous vegetation, *Nature Geoscience*, 4(9), 629–633, doi:  
424 10.1038/ngeo1237.
- 425 Diehl, R. M., A. C. Wilcox, J. C. Stella, L. Kui, L. S. Sklar, and A. Lightbody (2017),  
426 Fluvial sediment supply and pioneer woody seedlings as a control on bar-surface topog-  
427 raphy, *Earth Surface Processes and Landforms*, 42(5), 724–734, doi:10.1002/esp.4017.
- 428 Edmaier, K., P. Burlando, and P. Perona (2011), Mechanisms of vegetation uprooting by  
429 flow in alluvial non-cohesive sediment, *Hydrology and Earth System Sciences*, 15(5),  
430 1615–1627, doi:10.5194/hess-15-1615-2011.
- 431 Edmaier, K., B. Crouzy, and P. Perona (2015), Experimental characterization of vegeta-  
432 tion uprooting by flow, *Journal of Geophysical Research: Biogeosciences*, 120(9), 1812–  
433 1824.
- 434 Fan, Y., G. Miguez-Macho, E. G. Jobbágy, R. B. Jackson, and C. Otero-Casal (2017), Hy-  
435 drologic regulation of plant rooting depth, *Proceedings of the National Academy of Sci-*  
436 *ences*, p. 201712381.
- 437 Gibling, M. R., and N. S. Davies (2012), Palaeozoic landscapes shaped by plant evolution,  
438 *Nature Geoscience*, 5(2), 99–105, doi:10.1038/ngeo1376.
- 439 Gorla, L., C. Signarbieux, P. Turberg, A. Buttler, and P. Perona (2015), Transient response  
440 of salix cuttings to changing water level regimes, *Water Resources Research*, 51(3),  
441 1758–1774.
- 442 Gran, K. B., M. Tal, and E. D. Wartman (2015), Co-evolution of riparian vegetation and  
443 channel dynamics in an aggrading braided river system, mount pinatubo, philippines,  
444 *Earth Surface Processes and Landforms*, 40(8), 1101–1115, doi:10.1002/esp.3699.
- 445 Gregory, P. J. (2008), *Plant roots: growth, activity and interactions with the soil*, John Wi-  
446 ley & Sons.
- 447 Gurnell, A. (2014), Plants as river system engineers, *Earth Surface Processes and Land-*  
448 *forms*, 39(1), 4–25.
- 449 Gurnell, A. M., W. Bertoldi, and D. Corenblit (2012), Changing river channels: The  
450 roles of hydrological processes, plants and pioneer fluvial landforms in humid tem-  
451 perate, mixed load, gravel bed rivers, *Earth-Science Reviews*, 111(1), 129 – 141, doi:  
452 <https://doi.org/10.1016/j.earscirev.2011.11.005>.

- 453 Jackson, R. B., J. Canadell, J. R. Ehleringer, H. A. Mooney, O. E. Sala, and E. D. Schulze  
454 (1996), A global analysis of root distributions for terrestrial biomes, *Oecologia*, *108*(3),  
455 389–411, doi:10.1007/BF00333714.
- 456 Kim, H. S., I. Kimura, and Y. Shimizu (2015), Bed morphological changes around a fi-  
457 nite patch of vegetation, *Earth Surface Processes and Landforms*, *40*(3), 375–388, doi:  
458 10.1002/esp.3639, eSP-13-0354.R3.
- 459 Kim, J., V. Y. Ivanov, and N. D. Katopodes (2012), Hydraulic resistance to overland flow  
460 on surfaces with partially submerged vegetation, *Water Resources Research*, *48*(10).
- 461 Le Bouteiller, C., and J. Venditti (2015), Sediment transport and shear stress partitioning  
462 in a vegetated flow, *Water Resources Research*, *51*(4), 2901–2922.
- 463 Le Bouteiller, C., and J. G. Venditti (2014), Vegetation-driven morphodynamic ad-  
464 justments of a sand bed, *Geophysical Research Letters*, *41*(11), 3876–3883, doi:  
465 10.1002/2014GL060155.
- 466 Manners, R. B., A. C. Wilcox, L. Kui, A. F. Lightbody, J. C. Stella, and L. S. Sklar  
467 (2015), When do plants modify fluvial processes? Plant-hydraulic interactions under  
468 variable flow and sediment supply rates, *Journal of Geophysical Research: Earth Sur-  
469 face*, *120*(2), 325–345.
- 470 Meyer-Peter, E., and R. Müller (1948), Formulas for bed-load transport, in *2nd Meeting*,  
471 *Int. Assoc. of Hydraul. Eng. and Res.*, Stockholm.
- 472 Murray, A. B., and C. Paola (2003), Modelling the effect of vegetation on channel pattern  
473 in bedload rivers, *Earth Surface Processes and Landforms*, *28*(2), 131–143.
- 474 Nardin, W., and D. A. Edmonds (2014), Optimum vegetation height and density for inor-  
475 ganic sedimentation in deltaic marshes, *Nature Geoscience*, *7*(10), 722.
- 476 Nepf, H. M. (2012), Flow and Transport in Regions with Aquatic Vegetation, *Annual Re-  
477 view of Fluid Mechanics*, *44*(1), 123–142, doi:10.1146/annurev-fluid-120710-101048.
- 478 Oorschot, M. v., M. Kleinhans, G. Geerling, and H. Middelkoop (2016), Distinct patterns  
479 of interaction between vegetation and morphodynamics, *Earth Surface Processes and  
480 Landforms*, *41*(6), 791–808.
- 481 Orellana, F., P. Verma, S. P. Loheide, and E. Daly (2012), Monitoring and modeling  
482 water-vegetation interactions in groundwater-dependent ecosystems, *Reviews of Geo-  
483 physics*, *50*(3).
- 484 Pasquale, N., and P. Perona (2014), Experimental assessment of riverbed sediment rein-  
485 forcement by vegetation roots, *River Flow 2014*, pp. 553–561.

- 486 Pasquale, N., P. Perona, R. Francis, and P. Burlando (2012), Effects of streamflow variabil-  
487 ity on the vertical root density distribution of willow cutting experiments, *Ecological*  
488 *Engineering*, 40, 167–172, doi:10.1016/j.ecoleng.2011.12.002.
- 489 Perona, P., and B. Crouzy (2018), Resilience of riverbed vegetation to uprooting by flow,  
490 *Proceedings of the Royal Society A: Mathematical, Physical and Engineering Sciences*.
- 491 Politti, E., W. Bertoldi, A. Gurnell, and A. Henshaw (2018), Feedbacks between the ri-  
492 parian salicaceae and hydrogeomorphic processes: A quantitative review, *Earth-Science*  
493 *Reviews*, 176, 147 – 165, doi:https://doi.org/10.1016/j.earscirev.2017.07.018.
- 494 Pollen-Bankhead, N., and A. Simon (2010), Hydrologic and hydraulic effects of ripar-  
495 ian root networks on streambank stability: Is mechanical root-reinforcement the whole  
496 story?, *Geomorphology*, 116(3-4), 353–362.
- 497 Polvi, L. E., E. Wohl, and D. M. Merritt (2014), Modeling the functional influence of veg-  
498 etation type on streambank cohesion, *Earth Surface Processes and Landforms*, 39(9),  
499 1245–1258, doi:10.1002/esp.3577.
- 500 Ridolfi, L., P. D’Odorico, and F. Laio (2011), *Noise-induced phenomena in the*  
501 *environmental sciences*, xii, 313 p. pp., Cambridge University Press, doi:  
502 10.1017/CBO9780511984730.
- 503 Schwarz, C., O. Gourgue, J. van Belzen, Z. Zhu, T. J. Buoma, J. van de Koppel,  
504 G. Ruessink, N. Claude, and S. Temmerman (2018), Self-organization of a biogeomor-  
505 phic landscape controlled by plant life-history traits, *Nature Geoscience*.
- 506 Serlet, A. J., A. M. Gurnell, G. Zolezzi, G. Wharton, P. Belleudy, and C. Jourdain (2018),  
507 Biomorphodynamics of alternate bars in a channelized, regulated river: an integrated  
508 historical and modelling analysis, *Earth Surface Processes and Landforms*.
- 509 Simon, A., and A. J. C. Collison (2001), Pore-water pressure effects on the detachment of  
510 cohesive streambeds: seepage forces and matric suction, *Earth Surface Processes and*  
511 *Landforms*, 26(13), 1421–1442, doi:10.1002/esp.287.
- 512 Solari, L., M. Van Oorschot, B. Belletti, D. Hendriks, M. Rinaldi, and A. Vargas-Luna  
513 (2016), Advances on modelling riparian vegetation-hydromorphology interactions, *River*  
514 *Research and Applications*, 32(2), 164–178.
- 515 Tal, M., and C. Paola (2007), Dynamic single-thread channels maintained by the interac-  
516 tion of flow and vegetation, *Geology*, 35(4), 347, doi:10.1130/G23260A.1.
- 517 Tal, M., and C. Paola (2010), Effects of vegetation on channel morphodynamics: results  
518 and insights from laboratory experiments, *Earth Surface Processes and Landforms*,

- 519 35(9), 1014–1028, doi:10.1002/esp.1908.
- 520 Temmerman, S., T. Bouma, J. Van de Koppel, D. Van der Wal, M. De Vries, and P. Her-  
521 man (2007), Vegetation causes channel erosion in a tidal landscape, *Geology*, 35(7),  
522 631, doi:10.1130/G23502A.1.
- 523 Tron, S., F. Laio, and L. Ridolfi (2014), Effect of water table fluctuations on phreato-  
524 phytic root distribution, *Journal of Theoretical Biology*, 360, 102–108, doi:  
525 10.1016/j.jtbi.2014.06.035.
- 526 Tron, S., P. Perona, L. Gorla, M. Schwarz, F. Laio, and L. Ridolfi (2015), The signature  
527 of randomness in riparian plant root distributions, *Geophysical Research Letters*, 42(17),  
528 7098–7106, doi:10.1002/2015GL064857.
- 529 Vannoppen, W., M. Vanmaercke, S. De Baets, and J. Poesen (2015), A review of the me-  
530 chanical effects of plant roots on concentrated flow erosion rates, *Earth-Science Re-*  
531 *views*, 150, 666–678.
- 532 Vargas-Luna, A., A. Crosato, and W. S. Uijttewaal (2015), Effects of vegetation on flow  
533 and sediment transport: comparative analyses and validation of predicting models, *Earth*  
534 *Surface Processes and Landforms*, 40(2), 157–176.
- 535 Västilä, K., and J. Järvelä (2014), Modeling the flow resistance of woody vegetation using  
536 physically based properties of the foliage and stem, *Water Resources Research*, 50(1),  
537 229–245, doi:10.1002/2013WR013819.
- 538 Vesipa, R., C. Camporeale, and L. Ridolfi (2017), Effect of river flow fluctuations on  
539 riparian vegetation dynamics: processes and models, *Advances in Water Resources*,  
540 110(July), 29–50, doi:10.1016/j.advwatres.2017.09.028.
- 541 Vetsch, D., A. Siviglia, D. Ehrbar, M. Facchini, S. Kammerer, A. Koch, S. Peter, L. Von-  
542 willer, M. Gerber, C. Volz, D. Farshi, R. Mueller, P. Rousselot, R. Veprek, and R. Faeh  
543 (2017), *System Manuals of BASEMENT, Version 2.7*, Laboratory of Hydraulics, Glaciol-  
544 ogy and Hydrology (VAW). ETH Zurich.
- 545 Wohl, E., S. N. Lane, and A. C. Wilcox (2015), The science and practice of river restora-  
546 tion, *Water Resources Research*, 51(8), 5974–5997.
- 547 Yager, E., and M. Schmeeckle (2013), The influence of vegetation on turbulence and bed  
548 load transport, *Journal of Geophysical Research: Earth Surface*, 118(3), 1585–1601.