The Effects of Added Vegetation on Sand Bar Stability and Stream Hydrodynamics

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Abstract: Vegetation was added to a fully-developed sandy point bar in the meander of a constructed stream. Significant changes in the flow structure and bed topography were observed. As expected, the addition of vegetative resistance decreased the depth-averaged streamwise velocity over the bar and increased it in the open region. In addition, the secondary circulation increased in strength, but became confined to the deepest section of the channel. Over the point bar, the secondary flow was entirely outwards, i.e. towards the outer bank. The changes in flow led to changes in bar shape. Although the region of the bar closest to the inner bank accumulated sediment, erosion of the bar and the removal of plants by scouring was observed at the interface between the planted bar and the open channel.

Subject Headings:
Meandering Streams; Secondary Flow; Stream Restoration; Riverine Bars; Riparian Vegetation; Riverbank Erosion; Sediment Transport

Introduction
The most obvious hydraulic impact of aquatic and riparian vegetation is an increase in flow resistance and a reduction in conveyance capacity (Kouwen and Unny 1973; Kouwen, 1990; Wu et al 1999). Historically, many channels have been straightened and denuded of

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vegetation to accelerate the passage of peak flows, but these anthropogenic modifications can have unintended consequences on water quality, channel stability and stream ecology. For example, many studies have shown that aquatic macrophytes have a positive influence on water quality by utilizing nutrients, producing oxygen and detaining heavy metals and other contaminants (e.g. Chambers and Prepas 1994; Kadlec and Knight 1996; Windham 2003).

Vegetation also promotes biodiversity by creating diverse habitats through spatial heterogeneity in the flow (Kemp et al 2000; Crowder and Diplas 2000, 2002).

Vegetation is also an important ecosystem engineer within channel systems. By reducing near-bed velocity, in-stream and floodplain vegetation can both reduce erosion and promote deposition. Elliott (2000) explains how the aboveground portion of biomass helps increase sedimentation both by reducing the local flow velocities and by providing additional horizontal surface per volume upon which sedimentation can occur. The presence of vegetation can also exert control over river planform (Tal and Paola 2007). Therefore, widespread planting of vegetation is often advocated as a restoration technique for its ability to increase channel stability and for its ability to remove nutrients from the water (Mars et al. 1999; Abernethy and Rutherford 1998; Simon and Collison 2002; Pollen and Simon 2005). While many studies describe the stabilizing effects of vegetation, few have considered, few have considered the relative contributions of the aboveground biomass and the belowground root system. Corenblit et al. (2007) summarizes several experimental studies that indicate that the aboveground biomass in grasses and other channel-lining vegetation is more important than belowground biomass for sediment stability (Gyssels et al. 2005, James et al. 2002, Prosser et al. 1995). These studies suggest the reduction in erosion observed within regions of vegetation is due mostly to the reduction in the turbulent shear stress at the bed, but acknowledge that the root system remains
Recent studies have suggested that the addition of vegetation may create regions of enhanced erosion potential. McBride et al. (2007) observed that the presence of vegetation on a floodplain could elevate the turbulence level at the floodplain-channel interface, relative to unvegetated floodplains. They suggested that the presence of the vegetation created a region of high erosion potential at the channel-floodplain interface. Similarly, Temmerman et al. (2007) found the presence of vegetation to concentrate flow and encourage localized erosion in the growth of channels on tidal flats. Bouma et al (2007) created islands of bamboo shoots on an intertidal flat. Over two-years of monitoring they observed deposition to occur within and downstream of the patch, but erosion occurred at the front and sides of the patches. The observations of Bouma et al. (2007) can be related to previous observations around bridge piers. Bridge piers and abutments generally destabilize the substrate around them, resulting in scour holes. Many studies have recorded both the erosion caused by isolated pilings (Melville 1997, Melville & Chiew 1999), as well as the more complex scouring patterns caused by groups of piles (Ataie-Ashtiani & Beheshti 2006, Bateni & Jeng 2007). This area of research confirms that there is a real possibility for in-stream objects, including vegetation, to promote localized erosion. A finite patch of vegetation may operate in a similar fashion. The divergence of flow away from the region of high-drag within the patch results in accelerated flow at the edges of the patch, which in turn creates the regions of erosion observed by Temmerman et al. (2007) and Bouma et al. (2007).

To have successful replanting and restoration of channels, we need to understand how the placement and planting density impact the local flow field, which predicts the potential for deposition and erosion near the restored vegetation. This study seeks to provide some insight
into how the addition of vegetation at a specific location on the point bar in a channel bend, alters the flow field, which in turn leads to changes in the bed topography.

Description of Facility

The experiments were conducted in the Outdoor StreamLab (OSL), an experimental facility built on a retired spillway adjacent to the University of Minnesota’s St. Anthony Falls Laboratory in downtown Minneapolis. During 2008, within the facility’s 40-m by 20-m Riparian Basin, a sand-bed stream was constructed with three meander bends that have an average wavelength of 25 m and a sinuosity of 1.3 (Figure 1). In the straight sections in between the meanders, riffles were constructed with coarse-grained sediment and cobbles to mimic the pool-riffle geometry of many natural streams. A concrete headbox at the mouth of the stream was supplied with water from the Mississippi River via two 18 in. diameter steel pipes, which were controlled manually with knife-valves. The base flow rate for the stream was $38 \pm 5$ L/s and was calculated from the height of water above a contracted weir at the upstream end of the stream using an air ultrasonic transducer (Massa Products Corporation, Hingham, MA).

The banks of the channel were fixed in geometry and position with coconut fiber matting, but the bed of the channel was mobile, and consisted of coarse-grained sand (median grain size: $D_{50} = 0.7$ mm). A sediment recirculation system carried bedload sediment collected from the downstream end back to the upstream end of the channel using an adjustable rate auger. During the first flood event, point bars formed from the mobile bed material near the inner bank of the second and third meander bends (see Figure 2). These point bars formed within the first few hours of the first flood event on July 10, 2008, and remained as roughly stable artifacts in the stream during the base flow and subsequent flood events in July (Figure 3). The magnitudes of
the water surface slope, derived from the survey data, were $S = 0.006$ and 0.007 for the flood
level and base level flows, respectively. The magnitude of the average bed slope was 0.007.

Stream Coordinate System and Momentum Equations

It is useful to define a streamline coordinate system that follows the curvature of the
stream, with the downstream coordinate tangent to the stream centerline. This coordinate system
is left-handed, orthogonal and curvilinear, similar to systems defined by Smith and McLean
(1984) and Dietrich and Smith (1983) and consists of an $s$-axis, tangent to the centerline of the
stream and positive in the downstream direction, an $n$-axis, perpendicular to the stream centerline
and positive towards the right bank, and a vertical axis, $z$, positive in the upwards direction with $z
= 0$ at the water surface. The time-average velocity field is denoted $(u, v, w)$ in the directions $(s,$
$n, z)$, respectively. A depth average is denoted by a bracket. The force balance equations in the
downstream and cross-stream directions are then:

$$
\begin{align}
\tau_{s,b} &= -\rho g h \frac{\partial \eta}{\partial s} - \rho \frac{1}{1+N} \frac{\partial}{\partial s} \left\langle u^2 \right\rangle h - \rho \frac{\partial}{\partial n} \left\langle uv \right\rangle h - 2\rho \frac{\left\langle uv \right\rangle h}{(1+N)R} - \frac{1}{2} \rho C_D a h \left\langle \left| u \right| \right\rangle \\
\tau_{n,b} &= \rho g h \frac{\partial \eta}{\partial n} - \rho \frac{\left\langle u^2 \right\rangle h}{(1+N)R} + \rho \frac{1}{1+N} \frac{\partial}{\partial s} \left\langle uv \right\rangle h + \rho \frac{\partial}{\partial n} \left\langle v^2 \right\rangle h + \rho \frac{\left\langle v^2 \right\rangle h}{(1+N)R} - \frac{1}{2} \rho C_D a h \left\langle \left| v \right| \right\rangle
\end{align}
$$

Terms A1 and B1 represent the boundary shear stress, $\eta$ is the super-elevation of the water
surface, $h$ is the total depth of the water column, $\rho$ is the density of water, $g$ is the acceleration
due to gravity, $R$ is the local radius of curvature of the stream and the non-dimensional
coordinate $N = n/R$. Terms A6 and B7 represent the additional hydraulic resistance provided by
vegetation, when present. The vegetation density is represented by the frontal area per unit
volume, $a$, and $C_D$ is the vegetation drag coefficient.

To the leading order, the dominant terms in the cross-stream force balance are often B2
and B3, the cross-stream pressure gradient and the centrifugal force, respectively. Near the
down of the water column, bed friction causes a lower velocity, and thus a lower centrifugal
force. The pressure gradient, which is uniform over depth, is thus unbalanced near the bed,
driving a secondary flow toward the inside of the meander, i.e. toward the point bar. Near the
water surface, the velocity is higher than the depth-average, and the centrifugal force exceeds the
pressure gradient, causing the secondary flow to be outward, away from the point bar. In this
paper we examine how the addition of vegetation to a point bar changes this secondary
circulation, and consider the effects this has on water and sediment supply to the sand bar, as
well as the resulting changes in bed topography.

**Experimental Methods**

Bank-full flood events, representative of the average flood magnitude in natural channels,
were created at approximately weekly intervals. Each flood lasted 9 hours and had a constant
flow rate of $208 \pm 5$ L/s. At the start of each of the experimental floods, the flow rate was
gradually increased from the base flow rate (38 L/s) to the bank-full level (208 L/s) over a period
of ten minutes. The flow rate in the stream was monitored continuously and adjusted manually
so as to be independent of changes in the flow in the Mississippi River. Velocity measurements
were made during each of the repeated 9-hour, bank-full flood events. A 3D, 4-beam, sideways-
looking, fixed-probe, laboratory Nortek Vectrino Acoustic Doppler Velocimeter (ADV), was
used to simultaneously measure velocity in the $s, n$ and $z$ ($u, v$ and $w$) directions at different cross
sections along the length of the stream. The ADV was mounted on a motorized traverse oriented
perpendicular to the local stream direction, i.e. along the local $n$ - axis. At each cross section,
velocity was measured at roughly ten points in the horizontal, $n$ - axis, and a varying number of
points in the vertical, z-axis, due to the varying depth. The highest data points were within two
centimeters of the surface, and the lowest points were within 5 cm of the bed. The velocity was
recorded at each point for between 120 to 240 seconds at 25 Hz. The velocity data was filtered
to exclude erroneous values with low correlation coefficients or low signal-to-noise ratios
(SNRs). The most common cause of erroneous data points was an obstruction in the sampling
volume such as stream debris or a solid surface like the streambed or bank. Each transect was
aligned perpendicularly to the streambanks, and the transect position was marked with stakes and
surveyed to ensure repeatable positioning throughout the summer. Alignment was ensured by
checking the total flow-rate at each transect against the flow-rate delivered from the upstream
headbox.

Two types of tracer tests, using Rhodamine WT, provided information regarding the
transport at the reach-scale as well as locally around the point bar. First, to measure reach-
averaged parameters, dye was injected as a line source near the mouth of the stream and a
fluorometer was set up near the downstream bridge (Figure 1). We diluted 2 ml of Rhodamine
WT into a 500 ml solution and injected the solution over the stream’s cross-section over a period
of less than 1 second (approximately instantaneous). A submersible recording fluorometer
(SCUFA, Turner Designs) recorded dye concentrations in the water at the downstream bridge at
a rate of 1 Hz. To estimate the reach-scale longitudinal dispersion and the retention time, the
downstream concentration records were analyzed using the method of moments (e.g. Murphy et
al., 2007). Second, to estimate the difference in transport time-scales between the vegetated and
unvegetated regions in the second meander, an identical mixture of dye was injected
instantaneously as a planar source near the mouth of the stream and fluorometers were set up in
Planes C and D (Figure 1). The SCUFA was set at mid-depth in the middle of the vegetated sand
bar (points C1 and D1), and a Seapoint Sensors Fluorometer, sampling at 7.5 Hz, was set up in the same plane at the midpoint of the open region (points C2 and D2 in Figure 1). The two fluorometers were synchronized using a stopwatch.

Floods were started on July 10, 2008 and a steady bathymetry was established during the first flood. Surveys made with a Leica Total Station were used to record channel geometry as it developed over the summer. On August 5, 2008, two reed species, *Juncus effusus* and *Scirpus atrovirens* were planted on the portion of the sand bar in the second meander that was exposed at base flow. This vegetation was planted in a uniform, staggered array that produced a vegetated frontal area per unit volume of $a = 5.2 \text{ m}^{-1}$, where $a = n_{\text{plant}}d_{\text{plant}}$, $n_{\text{plant}} = 69 \text{ m}^{-2}$ is the number of plant plugs per unit area, and $d_{\text{plant (avg)}} = .075 \text{ m}$ is the characteristic diameter of a single plant plug (a close grouping of several individual stems. See Figure 4). Throughout the subsequent floods, the velocities and the reach-scale transport parameters were monitored for changes using the methods described above.

**Results and Discussion**

As expected, a secondary circulation was observed in the meander bends prior to the addition of vegetation. This circulation was most intense near the apex of the meander (Plane B of Figure 1), with a strong lateral outflow near the water surface and a return current near the bed of the stream (Figure 5b). The secondary circulation predominantly occupied the deeper part of the cross section, with smaller lateral, $v$, and vertical velocities, $w$, over the point bar. The depth-averaged streamwise velocity, $\langle u \rangle$, was highest near the outer bank of the meander and smallest over the point bar (Figure 5a).

After nearly one month of flow, including five floods, the point bar in the second
meander was planted with emergent vegetation during base flow conditions. During the first flood event after the planting (August 6, 2008), the cross-sectional geometry changed rapidly due to the flow disturbance created by the plants (Figure 3). The outermost row of plants was scoured away, as well as part of the next outermost row, removing approximately 50 cm of the emergent point bar’s width along with most of the vegetation in this zone. This loss of sediment and resulting loss of point bar area, observed in the early stages of the flood, was confirmed by photographic and survey data. Similar measurements for the unvegetated point bar in meander 3 showed no loss in emergent bar area, confirming that the loss observed in the second point bar was due to the added vegetation. The plants and bar area that were not scoured away in the first hours of the first flood were stable for the remainder of the summer flood sequence.

Both the depth-averaged streamwise velocity and the secondary circulation at the apex of meander 2 changed significantly after the vegetation was added (Figure 6). First, the depth-averaged streamwise velocity decreased over the bar and increased in the open region (Figure 6a). Second, the secondary circulation increased in strength, but was confined to the deepest section of the channel. The depth-averaged centrifugal force, i.e. term B3 (e.g. as in Kitanidis and Kennedy, 1983), which drives the secondary circulation, increased by 30% after the addition of vegetation to the bar (Figure 7). A direct calculation of the average v – component of the velocity at the apex of the bend, calculated by taking the root-mean-square of the velocities at each cross-stream coordinate in the channel, showed approximately a 50% increase after the vegetation was added. Because the strength of the secondary circulation increased significantly and the flow at the outer bank was enhanced, erosion would be likely in a natural channel, and this could accelerate the meander growth. This could not occur in our channel because the banks were fixed in position by buried fiber matting. Finally, over the point bar, a strong outwards
(toward the outer bank) flow extended over the entire depth of the water column.

The difference in the velocity field before and after the insertion of vegetation occurred because the vegetation increased the hydraulic resistance over the point bar. Defining the bed stress by a bed drag coefficient, \( \tau_{zs} = -\rho C_f \langle u \rangle \langle u \rangle \), we can compare the hydraulic resistance provided by the bed (term A1) with that provided by the vegetation (term A6), by comparing the terms \( C_f \approx 0.002 \) (for a sand bed) and \( C_D a h \approx 0.8 \), assuming \( C_D = O(1) \). This comparison indicates that the addition of vegetative drag (term A6) increased the total drag on the bar by two orders of magnitude, significantly retarding the flow and causing a lateral diversion toward the open channel. As the flow was diverted away from the region of high drag, the downstream velocity accelerated near the edge of the vegetation, causing the observed scour. Specifically, the velocity at the vegetation edge \( (y = 50 \text{ cm}) \) increased from 45 cm/s before the addition of vegetation (Figure 5a) to 55 cm/s after the addition of vegetation (Figure 6a).

These changes in the secondary circulation can be explained by consideration of the spatial acceleration terms in the cross-stream momentum balance. As water decelerated over the point bar, an effect that was magnified by the presence of vegetation, the downstream slope of the water surface was reduced. A concomitant acceleration of the flow in the deeper portion of the channel increased the downstream slope in this region of flow. The combined effect of these changes in the downstream surface slope was a reduction in the cross-stream surface slope (i.e. the cross stream pressure gradient) near the apex. This phenomenon can also be observed where flow shoals over a bare point bar, but here it was greatly magnified by the two orders of magnitude increase in resistance introduced with the vegetation. For a further discussion of the effects of spatial accelerations on the surface slope and a very helpful diagram, see Dietrich and Smith (1983). The result is that after the vegetation was added, the centrifugal force exceeded
the cross-stream pressure gradient over the entire depth over point bar, and caused a lateral flow
toward the open channel and outer bank that extended over the entire water depth in the
vegetated region, i.e. there was no return flow at the bed (Figure 6b). Importantly, the return
current near the bed was then limited to only the deepest parts of the channel (Figure 6b), in
contrast to the conditions before the vegetation (Figure 5b), in which the return flow extended
onto the bar. Importantly, these observations suggest that the addition of vegetation changed the
secondary flow in such a way as to cut off sediment supply from the open channel to the bar.

Finally, measurements of the bed geometry taken before and after the vegetation was
added show how the depth-profile changed (Figures 5b, 6b and 8b). Approximately 5 cm of
sediment was deposited within the vegetation and between 0 and 4 cm of erosion occurred near
the edge of the vegetation. Erosion (0 to 5 cm) also occurred in the deeper parts of the cross
section near the outer bank. The new bed geometry was recorded on August 26, 2008, four
floods after the addition of vegetation, but the changes were observed to take place primarily
during the first flood following the addition of the vegetation. The pattern of deposition within
the vegetated patch and erosion at its edge is similar to that observed by Bouma et al (2007)
within and around a circular patch of bamboo reeds.

Implications for Erosion

Several studies suggest that plant growth can be inhibited by flow. Chambers et al.
(1991), observed a strong negative correlation between macrophyte biomass and current
velocities, with little vegetation present above a threshold of 1 m/s. Similarly, Nilsson (1987)
found the percentage of bare ground along a reach increased with increasing current velocities in
the free stream. Bouma et al. (2007) saw similar sediment patterns within an artificial patch of
vegetation with erosion near the edges and deposition deeper within the patch. These studies imply that particular planting strategies can be less successful if they lead to locally enhanced velocities. To gain insight into this problem, we consider whether the observed loss of vegetation introduced in our study is consistent with our physical understanding of plant and sediment stability.

There are two physical mechanisms that limit the invasion and propagation of vegetation into a stream channel. First, for a given unconsolidated sandy bed, the substrate becomes mobile above a certain bed shear stress, defined by the Shields Parameter. But sediment motion alone does not govern the presence of plant life. A plant may be able to survive an area with weak sediment motion, but a rapidly scouring bed will cause plants to uproot and will preclude the growth of aquatic vegetation that depends on the substrate for stability (Fonseca et al., 1983).

Second, plants have an inherent lodging velocity that defines the flow speed at which the plant material fails. This value is a function of the stem flexural stiffness, geometry and natural roughness of the plant (Duan et al. 2002). For the mobile sand bed found in the OSL, the scouring threshold appeared to be reached before the lodging velocity because the plants lost during the flood came out as intact plugs, with no observed damage to the plant material. This implies that the plants dislodged because the substrate around them eroded, so that to understand this loss we must consider the changes in sediment stability.

The Shields Parameter, \( \psi \), describes the ratio of destabilizing (drag) and stabilizing forces (settling) for cohesionless sediment. This parameter is defined as

\[
\psi = \frac{\tau_b}{(\rho_s - \rho)gd} = \frac{\rho C_f \langle u^2 \rangle}{(\rho_s - \rho)gd}
\]  

Here, \( \rho_s \) is the sediment density, \( C_f \) is the coefficient of friction of the bed and \( d \) is the sediment grain diameter. We can evaluate the changing stability of the bed by comparing the Shields
Parameter before and after the vegetation was added. Because the sediment is unchanged, we can assume that both the settling forces and the bed friction coefficient do not change. It is then convenient to form the following ratio, to describe the changes in bed stability:

$$\frac{\psi_{\text{veg}}}{\psi_{\text{unveg}}} = \frac{\left( u_{\text{veg}} \right)^2}{\left( u_{\text{unveg}} \right)^2}$$

The areas in the vegetated cross section where erosion was observed corresponded to a Shields Parameter ratio of greater than 1 (Figure 8). Similarly, the areas in which deposition occurred corresponded to a Shields Parameter ratio of less than 1. The correspondence between the Shields Parameter ratio and the observed erosion/deposition patterns suggests that the vegetation changed the stability of the bed by altering the local flow speed and thus the local bed stress. This is consistent with the observation above, that the plants lost were removed intact, i.e. the sediment eroded away around the plug. Further, this set of experiments reinforces the theory that high flow energy and rapid sediment scour can preclude vegetative growth and propagation. The recorded changes in the bed profile agree well with the areas where vegetation was lost from the planted array. It is important to note that this ratio of the Shields Parameters indicates tendencies only. It does not suggest that certain areas will erode indefinitely and other areas will continue to accrete.

A more developed root system may have better anchored the plants in the sediment, but research shows that erosion of the sediment near the edge of the vegetation is representative of the behavior in real systems. In Bouma et al. (2007), the added vegetation was anchored 30 cm into the sediment, and despite the fact that none of the plants were lost during the experiment, significant erosion was still observed near the edges of the artificial patch. This indicates that the
diversion of flow would have caused similar erosion of the sediment whether the plants were able to remain anchored or not.

Furthermore, in this set of experiments the addition of vegetation to the stream channel is not directly analogous to a natural scenario where locally present vegetation slowly colonizes the bank via hydrochory or other processes. This experiment more closely represents a restoration scenario, where vegetation is added to an emergent point bar and the resulting adjustments caused by bank-full flow are observed, measured and documented. These experiments would be similar to a scenario in which vegetation propagated into a channel during an extended period of low flow, then was forced to adjust upon the return of bank-full flow levels, potentially showing similar patterns of erosion and deposition of suspended sediment.

Pathway of Surface Water over the Vegetated Bar

The availability of suspended sediment to the bar, as well as the water quality on the bar, both depend upon the supply of new water to this region. The tracer measurements made near the vegetated bar allowed us to draw conclusions about the advective and diffusive transport near the bar. Figure 9 shows the residence time distributions at two longitudinal positions for dye passing through the vegetation (fluorometer positions C1 and D1 in Figure 1) and dye passing through the adjacent open channel (fluorometer positions C2 and D2 in Figure 1). The arrival time of the peak dye concentration is delayed in the vegetation compared to the arrival time of the peak concentration in the open channel. The delays observed at cross-sections C and D (Figure 1) are $\Delta T_{\text{Plane } C} = 18 \pm 10 \text{ sec}$ and $\Delta T_{\text{Plane } D} = 27 \pm 3 \text{ sec}$. If the vegetated region was a distinct advection zone, with little lateral diffusive exchange between the open channel and the vegetated region, then the delay in the dye passage, $\Delta T_{\text{Plane } C}$, should be approximately two thirds
of $\Delta T_{Plane D}$, because Plane C is 2/3 the distance between Plane A (leading edge of the vegetation) and Plane D. Assuming the velocities remained close to constant, the observed timescales agreed with this hypothesis, suggesting that dye entered the sand bar at the upstream edge and advected in streamlines roughly parallel to the bank with little lateral supply from the open channel. Further, we can estimate the lateral diffusivity, $D$, from photographs of the dye evolution along the channel (e.g. as in Nappo and Hiscox, 2008). Then, using the width of the vegetation, $b = 0.7$ m, the lateral diffusive velocity, $D/b = 0.004$ m/s, was found to be much smaller than the measured lateral velocity, $v = -0.12$ m/s. Although some mixing was observed near the edge of the vegetation, the turbulent diffusion was not large enough to offset the significant outward lateral advection, such that diffusion provides a negligible scalar flux to the region of water over the bar. Therefore, longitudinal advection from the upstream portion of the vegetated sand bar was the dominant source of channel water to the region above the bar, and the only potential sediment supply as well. These findings were confirmed by photographs of dye streamlines within the vegetation, which showed little lateral mixing across the boundary (See Figure 10).

The reach-scale tracer tests indicated that the longitudinal dispersion coefficient ($K_x = 5.1 \pm 0.2 \times 10^{-4}$ m$^2$s$^{-1}$) did not change (within uncertainty) after the addition of vegetation on the bar. There was also no significant change in the residence time distribution following the addition of the vegetation. Although the vegetated sand bar creates a large slow-zone, only about 10% of the flow encounters this region, with the remaining flow diverted around it. In order to observe an impact on reach-scale dispersion, the flow needs to encounter multiple slow-zones, so that a larger fraction of the total flow experiences at least one such zone.
Conclusions

Vegetation was added to a fully-developed sandy point bar near the convex bank of a stream meander. Both the flow field and the bed topography changed dramatically after the addition of the vegetation. Before the vegetation was added, the secondary circulation included the shallow areas above the sand bar, with the return current near the bed acting as a supply of surface water and sediment to the bar region. After the addition of vegetation, the secondary circulation was present only in the deepest section of the meander, near the outer bank, and the flow above the sand bar was outward over the entire water column. Importantly, the vegetation altered the secondary circulation sufficiently to cut off a source of water and sediment to the bar.

Deposition occurred within the patch of vegetation, near the inner stream bank, illustrating the positive feedback through which vegetation can stabilize landforms (e.g. Tal and Paola 2007). However, Erosion occurred near the lateral edge of the vegetation, resulting in a 30% loss of emergent bar width at the apex at base flow. This included the removal of some newly planted vegetation. The positions of erosion and deposition were consistent with the observed changes in the Shields Parameter. Areas where the Shields Parameters ratio was greater than unity corresponded to regions of erosion and areas where the ratio was less than unity corresponded to areas of deposition. The observations suggest that the spatial accelerations caused by the presence of the vegetation shifted the sand bar area to a new geometric equilibrium. More generally, these observations exemplify why the edges of a vegetation patch can be regions of enhanced erosion, as was observed previously by Bouma et al (2007) and Temmerman et al. (2007). Specifically, flow divergence associated with the additional vegetative-drag results in flow accelerations at the patch edge, which can lead to erosion at the patch edge. This phenomenon is similar to the scour patterns observed in the field around...
individual flow obstructions, such as bridge piers or abutments.

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References


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**Figure 1:** Plan view of the Riparian Basin of the Outdoor Stream Lab indicating measurement locations.
Figure 2: Diagram of the stream cross section with the vegetation added to the area of the sand bar emergent at baseflow. The dimensions provided are the design dimensions and do vary somewhat due to the mobility of the bed.
Figure 3: The width of the point bar developed over time, beginning with the first flood event on July 10, 2008, and also with the addition of vegetation to the point bar in Bend #2 on August 5, 2008. These widths were derived from survey data points taken at the apex of meander 2. The pre-vegetation mean width is $148 \pm 11$ cm. The vegetated mean width is $99 \pm 2$ cm.
Figure 4: Plan view schematic of the vegetative array on the point bar in meander #2. The length, $d_{\text{plant}}$, refers to the effective width of the plant (averaged over its height), which is the combined width of all of the stems from a single plug projected onto the streamflow. The spacing, $s_{\text{plant}}$, refers to the average distance between the centers of two plugs. The thumbnail photos show the plant plugs against a 5 cm grid.
FIGURE 5: Velocity Measurements at Apex 2 on July 16, 2008. (a) Depth averaged
downstream velocity, $<u>$, and (b) velocity components in the lateral and vertical directions, $v$
and $w$, showing the secondary circulation in the $n$-$z$ plane. The cross-sectional outline shows the
measured bed profile, measured by hand from the stream surface.
FIGURE 6: Velocity Measurements at Apex 2 with the vegetated sand bar on August 26, 2008. (a) Depth averaged downstream velocity, $\langle u \rangle$, and (b) velocity components in the lateral and vertical directions, $v$ and $w$, showing the secondary circulation in the $n$-$z$ plane. Note the lateral outflow present over the entire stream depth near the right bank. The cross-sectional outline shows the measured bed profile, measured by hand from the stream surface.
FIGURE 7: The depth averaged centrifugal force and the cross-sectionally averaged centrifugal force at the apex of Meander # 2 from before and after the vegetation was added to the system.
FIGURE 8: (a) The ratio of the Shields Parameters $\psi_{\text{veg}}/\psi_{\text{unveg}}$ showing the tendency of the system towards either deposition ($<1$) or erosion ($>1$) and (b) the change in the bed height following the addition of vegetation to the system.
FIGURE 9: Residence time distributions of dye passing through the open channel and vegetated regions in (a) Plane C and (b) Plane D.
Figure 10: A continuous injection of Rhodamine WT (black in the exaggerated contrast image). Note that the tracer cannot spread across the width of the vegetation because the observed diffusion is offset by an outward advection from the bank toward the open channel.