Can we manipulate root system architecture to control soil erosion?

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Abstract

Soil erosion is a major threat to soil functioning. The use of vegetation to control erosion has long been a topic for research. Much of this research has focused on the above ground properties of plants, demonstrating the important role that canopy structure and cover plays in the reduction of water erosion processes. Less attention has been paid to plant roots. Plant roots are a crucial yet under-researched factor for reducing water erosion through their ability to alter soil properties, such as aggregate stability, hydraulic function and shear strength. However, there have been few attempts to specifically manipulate plant root system properties to reduce soil erosion. Therefore, this review aims to explore the effects that plant roots have on soil erosion and hydrological processes, and how plant root architecture might be manipulated to enhance its erosion control properties. We demonstrate the importance of root system architecture for the control of soil erosion. We also show that some plant species respond to nutrient enriched patches by increasing lateral root proliferation. The erosional response to root proliferation will depend upon its location: at the soil surface dense mats of roots may reduce soil erodibility but block soil pores thereby limiting infiltration, enhancing runoff. Additionally, in nutrient deprived regions, root hair development may be stimulated
and larger amounts of root exudates released, thereby improving aggregate stability and decreasing erodibility. Utilising nutrient placement at specific depths may represent a potentially new, easily implemented, management strategy on nutrient poor agricultural land or constructed slopes to control erosion, and further research in this area is needed.

1. Introduction

Soil erosion by water is a serious environmental problem representing a much greater threat to agricultural production than soil erosion by wind (Lal, 2010). Approximately 430 million ha of arable land, or 30% of the globally available land for agricultural production, has been degraded by water erosion and a further 56% is at risk (Jankauskas et al., 2008). The effect of vegetation on reducing soil erosion is well known. Plants intercept rainfall, some is stored in the canopy with the remainder evaporating or reaching the soil surface either directly as throughfall, or indirectly through stem flow or leaf drainage (Stocking and Elwell, 1976; Puigdefàbregas, 2005). Water reaching the soil surface may be stored in small depressions on the soil surface or infiltrate into the soil. The remainder may move downhill as overland flow, entraining soil particles and transporting them, and may also concentrate to form rills and gullies. There are a number of factors controlling erosion, namely the erosivity of the eroding agent, slope angle, the nature of the plant cover and the erodibility of the soil (Morgan, 1986).

Erosion may result in on-site soil degradation reducing the soil’s productivity and water storage capacity, due to changes in soil structure, soil depth and losses of nutrients and organic matter (OM). This may lead to significant losses of crop yield (Lal, 2001; Pimentel, 2006; Jankauskas et al., 2008), posing a threat to food security or limiting the production of renewable biotic resources (Pimentel, 2006; Jankauskas et al., 2008). Off-site problems associated with downstream sedimentation include flooding, pollution, negative effects on biodiversity, as well as the loss of reservoir storage (Pimentel and Kounang, 1998; Lal, 2001; Pimentel, 2006). In the UK, the annual costs of both on-site and off-site problems were
recently estimated to range between £248 and £469 million (Posthumus et al., 2013). Hence, there is a need to stabilize soils and preserve this crucial resource. The importance of vegetation cover to control erosion has been demonstrated by many studies. Both linear and exponential decreases in erosion rates associated with vegetation cover have been observed, as the nature of the relationship varies for canopy and ground covers, or inter-rill and rill erosion (Boer and Puigdefàbregas, 2005). In Montana and Utah, for example, overland flow erosion rates increased ca. 200 times in response to a decrease in ground cover from 100% to less than 1% (Trimble and Mendel, 1995). A minimum of 10% plant cover was required to significantly reduce erosion rates on a semi-arid hillslope in Arizona (Abrahams et al., 1988), however, the magnitude of the response ultimately depends on vegetation type and species (Bochet et al., 2006). Positive effects of the aerial biomass in controlling erosion are generally attributed to reductions in the kinetic energy of raindrops and reduced overland flow velocities (Stocking and Elwell, 1976; Puigdefàbregas, 2005). Furthermore, vegetation covers modify intrinsic soil properties determining a soil’s erodibility (Gyssels et al., 2005) creating a microclimate and supplying OM, that affect the activity of microorganisms, and therefore nutrient availability, resulting in a positive feedback on plant productivity. Both additional OM as well as microbial and fungal secretions enhance the formation of stable aggregates (Puigdefàbregas, 2005, Vásquez-Méndez et al., 2010), which affect hydraulic conductivity and, hence, water storage capacity (Puigdefàbregas, 2005) and may increase soil shear strength (Tengbeh, 1993; De Baets et al, 2008).

Compared to studies on above ground plant properties and soil erosion, the effects of below ground biomass have received relatively little attention. However, it is clear that the reduction in soil loss due to vegetation results from the combined effects of the above and below ground biomass (Gyssels and Poesen, 2003). Concentrated flow erosion rates were reduced to almost zero within the 0-10% soil cross-sectional occupation by grass roots, but decreased only by
25-50% for the same increase in aerial cover (De Baets et al., 2006). Furthermore, a rainfall simulation experiment conducted 27 weeks after ryegrass (*Lolium perenne*) was sown in erosion pans demonstrated (by removal of the above ground material) that the below ground material of this species accounted for 90% of the decrease in soil detachment (Zhou and Shangguan, 2007).

In this paper we take an interdisciplinary view of the plant root and its interaction with the soil, in line with the interdisciplinary nature of this journal (Brevik et al., 2015), and review the literature concerning the effects of roots on soil properties determining the resistance of soil to water erosion (concentrated and sheet flow) with emphasis on root system architecture (RSA). We then explore the factors influencing RSA especially lateral root (LR) growth, and finally the potential of fertilizer applications to stimulate LR proliferation, and thus form the basis for a soil erosion control method, will be discussed.

### 2. The effects of roots on soil erodibility

Erodibility is the result of various mechanical, adhesive and cohesive bonding forces acting within the soil matrix (Knapen et al., 2007) and should be regarded as a ‘summation of a highly complex response pattern, strongly influenced by intrinsic soil characteristics and extrinsic variables‘ (Bryan, 2000). The erosion response may be influenced by any soil property, but will be dominated by shear strength, aggregate stability and hydraulic function (Bryan, 2000). Roots alter the erodibility of a soil through their effect on these soil properties in a number of ways (Figure 1). Firstly, roots have a high tensile strength (Gyssels et al., 2005; De Baets et al., 2008) providing up to 100% of the cohesive strength of a root permeated soil (Hales et al., 2009), thus increasing soil shear strength. The additional tensile strength associated with roots is responsible for the transfer of shear stresses via tensile resistance or interface friction within the soil-root matrix (Prosser et al., 1995, De Baets et al.,
There was a positive linear relationship between the cross-sectional area of barley (Hordeum vulgare) roots at the shear plane and, subsequent increases in soil shear strength in a silty clay loam soil (Waldron, 1977). In ryegrass, increasing root densities from 0.20 g cm$^{-3}$ to 1.80 g cm$^{-3}$ also increased strength from 1 kPa to 5 kPa in a sandy clay loam (Tengbeh, 1993). Hence, the presence of roots considerably increases shear strength.

Secondly, roots prevent sediment transport by retaining soil particles (De Baets et al. 2006) and increasing the number of stable aggregates, due to their enmeshing effect (Haynes and Beare, 1997), which is also true for fungal hyphae (Haynes and Beare, 1997; Degens, 1997; Moreno-Espíndola et al. 2007), and the release of various organic and inorganic binding agents (Amezketa, 1999). Root exudates such as mucilage are considered the main mechanism by which plant roots enhance aggregate stability (Morel et al., 1991; Moreno-Espíndola et al. 2007). Root mucilage expands under wet conditions covering soil particles, while subsequent drying leads to the contraction of mucilage, pulling these soil particles tightly together to form a rhizosheath, and thereby reorienting clay particles parallel to the axis of the root (Oades, 1984; Watt et al., 1994; Amezketa, 1999; McCully, 1999). Moreover, root exudates are associated with the release of polyvalent cations that form strong bonds between organic molecules and clays (Oades, 1984; Pojasok and Kay, 1990; Amezketa, 1999). Increasing amounts of ions in solution after exudates were added limit clay dispersion and significantly increase aggregate stability (Pojasok and Kay, 1990). Decaying roots and fungal hyphae also represent a source of OM (Oades, 1984; Haynes and Beare, 1997; Amezketa, 1999) that contributes to aggregate stability. Thus aggregate stability is enhanced by biological activity (both plant and fungal) within the soil.

Root exudates represent 5 to 21% of a plant’s photosynthetically fixed carbon (C) (Walker et al., 2003) serving as a crucial energy source for microorganisms (Haynes and Beare, 1997; Amezketa, 1999). Both fungal and bacterial extracellular polymeric substances contribute to
the cementation of aggregates, and, thus enhance aggregate persistence (Oades, 1984; Morel et al., 1991; Watt et al., 1994; Haynes and Beare, 1997; Bryan, 2000). However, the effect of roots on aggregate stability is species specific (Haynes and Beare, 1997; Amezketa, 1999) due to differences in root morphology (Pohl et al., 2009), turnover rates (Stokes et al., 2009), the quantity and chemical composition of the root exudates and the influence of differences in root exudates on microbial activity (Haynes and Beare, 1997, Bertin et al., 2003). Barley roots increased aggregate stability (Haynes and Beare, 1997), whilst there is contradicting evidence about the effects of roots of maize (Zea mays). Root mucilage of maize reportedly increased aggregation (Morel et al., 1991), but the release of chelating agents, such as organic acids (which destroy iron and aluminium bonds with OM) by maize decreased aggregate stability (Oades, 1984). These contrasting results may be attributed to genotypic variation in the chemistry of the root exudates. For example, within different barley genotypes, three structural variations of mugineic acid (an organic ligand) have been found (Bertin et al., 2003). In addition, some organic compounds released by roots and microorganisms may also induce water repellency depending on soil matric potential and the number of wetting and drying cycles undergone (Hallett et al., 2003). However, OM and aggregate stability are crucial in the formation of macropores, which reduce bulk density and facilitate gaseous diffusion and improve infiltration rates (Gyssels and Poesen, 2003). Thus, the effects of root exudates vary between species and depend on their composition.

Roots may also provide soils with additional surface roughness enhancing infiltration rates (De Baets et al. 2006), while water uptake through roots (Macleod et al., 2007) and the creation of continuous pore spaces increase the infiltration capacity, reducing surface runoff (Bryan, 2000; Bronick and Lal, 2005; Macleod et al., 2007). Quinton et al. (2002) found a positive correlation between infiltration rates and root density for needle grass (Stipa tenacissima) and, to a lesser extent, for downy plantain (Plantago albicans), while unclear
results were obtained for albaida (*Anthyllis cytisoides*) and white wormwood (*Artemisia herba-alba*). Under alfalfa (*Medicago sativa*), 60% of all flow paths are associated with the roots or the channels of decomposed roots allowing water to rapidly percolate through the soil matrix (Stokes *et al.*, 2009). Shallow, fibrous root systems and rhizomatous mats of ryegrass and common bent grass (*Agrostis capillaris*), respectively, resulted in a low saturated hydraulic conductivity (Morgan *et al.*, 1995). Several authors (Mitchell *et al.*, 1995; Meek *et al.*, 1989; Archer *et al.*, 2002; Devitt and Smith, 2002) suggest that living roots occupy available macropores for flow pathways, and enhance infiltration rates over time, as a result of channels created by decaying roots. Infiltration rates in a sandy loam, for instance, tripled within three years of alfalfa cultivation (Meek *et al.*, 1989) and significantly increased infiltration rates were recorded for the last half of the growing season, and after harvest, of a maize crop (Archer *et al.*, 2002). This is also supported by observations that the root effect of ryegrass on increasing infiltration rates (thus decreasing runoff) increased with time (Zhou and Shangguan, 2007). Moreover, the effects associated with cyclic shrinking-expanding of roots in accordance with water availability also created flow pathways over time (Archer *et al.*, 2002). However, the effect of roots on flow pathways, and thus infiltration rates, differed under a number of Mediterranean species, due to differences in root morphology (Quinton *et al.*, 2002), and turnover rates, as a result of differences in root diameter and chemical composition (Stokes *et al.*, 2009). Thus, not only the presence of roots, but their physical arrangement within the soil, alters soil hydraulic properties.

A frequently used root parameter, that sufficiently describes the effectiveness of a species to control concentrated flow erosion rates (Burylo *et al.*, 2012), is root length density (RLD). RLD is the total length of the roots divided by the volume of the root permeated soil sample (De Baets *et al.*, 2006), thus providing information about the occupation of soil by roots (Bauhus and Messier, 1999; De Baets *et al.*, 2006, De Baets *et al.*, 2007). Under cereal and
grass plants in the loess belt of Belgium, concentrated flow erosion rates exponentially
decreased with a linear increase in RLD (Gyssels and Poesen, 2003). Similar results were
obtained by Mamo and Bubenzer (2001a, b) between erosion rates of a silt loam soil
subjected to concentrated flow and RLD in maize and soybean (*Glycine max*), which reduced
detachment rates by 50% compared to fallow soils. A similar relation exists between RLD of
barley and detachment rates (De Baets *et al*., 2006). However, increasing RLD either affects
soil shear strength positively (Ghidey and Alberts, 1997; Mamo and Bubenzer, 2001a, b), or
not at all, depending on species and soil type (Pierret *et al*., 2007). Furthermore, higher RLDs
enhance soil aggregation in crop species (Haynes and Beare, 1997) and when comparing
grass, forb and legume species from a natural grassland ecosystem (Rilling *et al*., 2002).
However, the ability of roots to reinforce a soil is not only determined by root characteristics
such as RLD, but also their distribution within the soil.

3. Root system architecture and its manipulation

RSA is the three-dimensional deployment of roots (Linkohr *et al*., 2002; Basu *et al*., 2011). In
eudicots, RSA is mainly determined by the extent and direction of root tip growth and the
positioning and frequency of LRs (Jones and Ljung, 2012). Therefore, in this review
‘manipulating RSA‘ refers to alterations being made to the spatial configuration of the roots
of an individual plant in respect of LR growth (i.e. initiation, elongation, diameter, branching)
by stimulating proliferation of these roots at certain depths. RSA is constitutively influenced
by genetic factors (Smucker, 1993; Malamy, 2005; Stokes *et al*., 2009), as demonstrated by
mutants of maize lacking different root classes (Malamy, 2005). Mature cereal root systems
consist mostly of coarse (seminal – seed derived, or nodal – shoot derived) roots and fine
(lateral) roots (Ø <1mm) (LRs), which often develop root hairs (McCully, 1999) from the
trichoblasts in the meristematic zone (Gilroy & Jones, 2000). Although LRs are the major
sites of water and nutrient uptake, coarse roots are mainly responsible for anchorage, the
storage of nutrients, water and C, as well as the penetration of hard soils and deep horizons (Guerrero-Campo et al. 2006). LRs tend to have greater growth rates than other roots (Guerrero-Campo et al., 2006; Stokes et al., 2009) and account for ca. 90-98% of the total root length of a root system (Engineer and Kranz, 2007; Pohl et al., 2009). Hence, total root length is mainly determined by the density of LRs (Guerrero-Campo et al., 2006; De Baets et al., 2007; Pierret et al., 2007). Thus RLD significantly increased with the percentage of LRs (Burylo et al., 2012), and decreased with increasing root diameters (Pohl et al., 2009).

Moreover, different types of root system are often distinguished based on the presence of a primary root representing a single dominant axis with varying degrees of LR development, which is characteristic for primary or tap root systems (Fitter, 1987), whereas root systems lacking a primary root are referred to as a fibrous root system (De Beats et al., 2007). Only a few studies have compared the effect of these different root systems on soil erosion rates, with fibrous root systems being more effective in reinforcing soils against concentrated flow than tap root systems with similar root mass (De Baets et al., 2007). In contrast, tap rooting species increase saturated hydraulic conductivity, which tends to be low under fibrous root systems (Archer et al., 2002; Quinton et al., 2002). These observations can be explained by the larger number of LRs and greater root length densities (RLD) associated with fibrous root systems compared to tap root systems (De Baets et al., 2007).

Macleod et al. (2007) propose specific genetic selection and breeding for root traits, such as rooting depth, to improve soil hydraulic function. However, these root traits are under complex genetic control (Zhu et al., 2005; Macleod et al., 2007) even though alleles determining, for example, LR primordia development in thale cress (Arabidopsis thaliana) (Roycewicz and Malamy, 2014), as well as LR branching or length in maize (Zhu et al., 2005), have been identified. Limited knowledge of the root phenome, the ‘chaotic’ nature of the interactions among phenes and environmental factors, ultimately determining crop
performance and function (Lynch and Brown, 2012), hinder the development of new crop
varieties with altered root traits. Although RSA is genetically constrained, it is highly plastic
in response to environmental cues. This means that genetically identical plants may differ in
root density, growth rate, placement and growth direction (Müller and Schmidt, 2004;
Malamy, 2005), as a result of adaptive responses (López-Bucio et al., 2003; Jones and Ljung,
2012) to the prevailing environmental conditions. The heterogeneous distribution of nutrients
can increase LR length and initiation, as well as coordinate proliferation of LRs in soil regions
where these resources are present (e.g. Fitter, 1987; Smucker, 1993; Stokes et al. 2009).
Hence, the localized application of nutrients at prescribed depths and associated LR
proliferation offers an opportunity to manipulate LR distribution within the soil profile,
thereby promoting RSAs that reduce soil erosion.

4. Lateral root proliferation in response to local nutrient availability

The local proliferation of LRs allows the optimal exploitation of locally available nutrient
reserves (Smucker, 1993; Robinson, 1994; Hodge, 2006). Lack of nitrogen (N), as well as
phosphorus (P), often limits growth (Linkohr et al., 2002; López-Bucio et al., 2003). Nitrate
(NO$_3^-$) is the major source of N for most plants and is, as well as ammonium (NH$_4^+$), readily
available to plants (Engineer and Kranz, 2007). A series of studies by Drew and co-workers
investigated the response of barley roots to local nutrient enrichment. Although a
heterogeneous distribution of potassium (K) did not affect RSA (Drew, 1975; Robinson,
1994), LR initiation and extension increased within a 4 cm long segment of a seminal root
exposed to 100 times greater concentrations of NO$_3^-$ and NH$_4^+$ than the remainder of the root
system, where nutrient-limited conditions inhibited LR development (Drew et al., 1973;
Drew, 1975; Drew and Staker 1975) (Figure 2). The same is true for wheat (Triticum
aestivum) (Drew, 1975; Forde and Lorenzo, 2001) and maize grown in a flowing nutrient
culture whereby two nodal roots were exposed to different NO$_3^-$ concentrations revealed
similar results (Granato and Raper, 1989). Furthermore, a concentration of 1 mM NO$_3^-$ stimulated LR branching and increased LR diameters in both maize and barley (Drew, 1975; Granato and Raper, 1989), possibly due to increases in stele diameter and the numbers of cells in the root cortex and stele (Forde and Lorenzo, 2001). When thale cress was grown on vertical agar plates with a NO$_3^-$-enriched band halfway down the plate, LR lengths increased 2-3 fold, although there was no evidence for enhanced LR initiation (Zhang and Forde, 1998). However, nutrient solutions containing NH$_4^+$ as primary N source lead to higher root densities and increased root elongation in maize compared to solutions containing NO$_3^-$ (Jing et al., 2010). The localized availability of P often also results in increased LR extension (Drew, 1975; Robinson, 1994) and sometimes increased LR initiation within this zone (Drew, 1975). However, Drew (1975) highlights the importance of the presence of both nutrients, N and P, in stimulating LR proliferation. In addition, LR proliferation has been reported in response to organic and inorganic N and P, although the response to the former is determined by the extent and rate in which mineralization has occurred (Robinson, 1994). Thus LR proliferation in response to localised nutrient patches is well documented in many species grown in different artificial (not soil) media.

Root proliferation is believed to be triggered partly by shoot-to-root signalling, as the developmental response to localized increased NO$_3^-$ was most pronounced with plants of low N status (Forde and Lorenzo, 2001; Forde, 2002). High NO$_3^-$ levels result in increased auxin concentration in the LR tip stimulating LR growth, while high NH$_4^+$ levels enhance higher order root branching. The presence of both forms of N results in a complementary response (Forde, 2002; Jones & Ljung, 2012). However, if NO$_3^-$ concentrations are too high, LR growth tends to be inhibited (Zhang and Forde, 1998; Linkohr et al., 2002), which is mediated by abscisic acid (López-Bucio et al. 2003). A detailed discussion of the molecular
mechanisms responsible for LR proliferation in nutrient-rich zones is beyond the scope of this paper, but the topic has been recently reviewed by Jones and Ljung (2012).

However, the growth media used in these studies (agar, hydroponics) do not necessarily reflect natural conditions and it is important to determine whether such responses occur in soil. Paradoxically, when wheat plants were grown in soil-filled rhizotrons (1.4 m deep) with nutrients (both N and P) banded at 0-40 cm or 60-100 cm below the soil surface and the soil allowed to dry (simulating a terminal drought stress), the greatest RLD detected at 1.4 m occurred with the surface fertiliser application (Jin et al., 2015). However, fertiliser banding (stratification) and local placement (patches) can elicit different responses. LRs of thale cress in a NO$_3^-$ enriched patch in a heterogeneous soil were only marginally longer than in the corresponding patch under homogeneous conditions (Linkohr et al. 2002). Nevertheless, LR proliferation has been observed in the field (Sattelmacher et al., 1993; Robinson, 1994). A recent study in North China showed that banding NH$_4^+$ and P at 10 cm depth resulted in a 23-30% increase in total root length and higher RLD at a depth of 0-15 cm in maize compared to broadcast application, or banding of P only while NH$_4^+$ was broadcast. Moreover, banding of both nutrients also increased RLD at a depth of 15-30 cm (Jing et al., 2010). However, the responsiveness of plants to localized nutrient availability varies between different species and genotypes (Robinson, 1994).

5. Manipulating root system architecture with nutrient placement to control erosion

Beyond the selection of different plant species, little consideration has been given to the potential for manipulating RSA for soil erosion control, despite it being known that the distribution of LRs within the top 50 cm of the soil is crucial in determining the effect of roots on the resistance of soils to concentrated flow erosion (Li et al., 1991). Dense mats of LRs are
commonly found at the soil surface (Pierret et al., 2007) and the abundance of roots gradually declines with increasing soil depth (Quinton et al., 2002; De Baets et al., 2008). However, it has been argued higher RLDs at deeper soil depths would be particularly beneficial, as this increases possible water uptake and evapotranspiration rates, thus infiltration rates (Stokes et al., 2009) and would provide reinforcement to deeper soil layers in situations where concentrated flow erosion is problematic. In addition, longer vertical roots, that increase RLD at depth, provide conduits to deeper soil layers (Archer et al., 2002; Devitt and Smith, 2002; Quinton et al., 2002) and directly affect water storage capacity of the effective soil depth (Macleod et al., 2007; Stokes et al., 2009). Improvements in aggregate stability tend to be particularly pronounced in, if not spatially limited to, the rhizosphere (Oades, 1994; Foster, 1988; Morel et al., 1991; Haynes and Beare, 1997), so the extent of the root system determines its effect on soil aggregation (Haynes and Beare, 1997). The effects of roots of the grasses Stipa bungeana and Bothriochlon ischaemum on the properties of a Chinese loess soil (SOM and aggregate stability, especially in large diameter classes) decreased with increasing depth, as RLD deminished (Li and Li, 2011). The higher sediment load derived from deep soil horizons associated with the higher susceptibility of these soil horizons to erosional forces was further enhanced by increasing runoff discharge and slope gradient (Li and Li, 2011) and was particularly critical during concentrated flow erosion. Furthermore, soil cohesion provided by roots is also limited to the rooting depth of the vegetation, consequently soil reinforcement decreases with increasing depth (Gyssels et al., 2005; De Baets et al., 2008). Although Mediterranean shrubs typically only stabilize the top 50 cm of gully walls and hillslopes, failures may occur at deeper depths and increases in soil strength depend on the number of LRs that cross the potential shear plane (Reubens et al., 2007; De Baets et al., 2008).
Plants may also respond to heterogeneous nutrient supply with compensatory root growth (Drew, 1975; Robinson, 1994). Compensatory growth means that proliferation within the N or P enriched zone compensates, and may be proportional to, the inhibitory effect of the limited availability of these nutrients on LR development outside this zone (Robinson, 1994).

In this case, soil properties at different soil layers may change accordingly to the proportion of LRs present within these layers. Reduced LR growth on the soil surface, when nutrients are only available at deeper depths, could reduce resistance to surface erosion, but could potentially increase the water storage capacity of a soil, as the number of blocked pores by LRs at the soil surface would be reduced. Furthermore, although the diameter of LR increases in nutrient enriched zones (Drew, 1975; Granato and Raper, 1989; Forde and Lorenzo, 2001), LR proliferation may still increase shear strength locally (De Baets et al., 2008; Loades et al., 2010) and could help to reinforce soils at the shear plane. In addition, aggregate stability may be improved within the nutrient poor patch, as larger quantities of mucilage may be released within this zone to mobilise scarcely available P (Bertin et al., 2003). In contrast, Paterson et al. (2006) showed that LR proliferation in response to localized availability of NO₃⁻ resulted only in a locally increased release of root exudates in barley. Compensatory growth may also enhance root hair development in the nutrient poor zone, and root hairs contribute to soil aggregation in the rhizosphere and are crucial for rhizosheath development (Hochholdinger et al., 2004; Moreno-Espindola et al., 2007). Root hair length increased from 0.1-0.2 to 0.7 mm and density by a factor of 2-4 when spinach (Spinacia oleracea), tomato (Solanum lycopersicum) and rapeseed (Brassica napus) were grown in nutrient solutions with concentrations of 2 μM P compared to concentrations of 1000 μM P (Foehse and Junk, 1983). Split-root system experiments demonstrated root hair development of spinach in high P concentrations is not affected if the remainder of the root system is exposed to low P concentrations. In contrast, the magnitude of the positive effect on root hair development of limited NO₃⁻ varied between species. However, further research is needed into the effects of
differences in LR growth dynamics in response to localized fertilizer application on soil properties and subsequently erosion rates.

6. Effects of LR proliferation on aerial growth

Potential effects of LR proliferation on above ground biomass production are crucial to achieve high agricultural production, but may also be important in soil erosion control as mentioned before. Recent estimates suggest root proliferation is a relative low cost strategy adapted by plants to effectively absorb available ions and water in heterogeneous environments, requiring only an additional 0.2% of the plant’s daily carbon gain (Hodge, 2006). Only roots with small diameters proliferate (Drew, 1975; Granato and Raper, 1989; Smucker, 1993; McCully, 1999), as fine roots represent less resistance to the flow of ions and water during the uptake of these resources (Smucker, 1993). However, net C costs will be determined by factors such as rooting volume, photosynthetic supply and may be even zero if compensatory growth occurs (Robinson, 2001). This should buffer potential negative effects of limited nutrient availability within some soil areas on above ground biomass, although biomass partitioning ultimately depends on shoot nutrient status (Drew, 1975; Drew and Saker, 1975; Robinson, 1994). LR proliferation either increases or has no effect on root to shoot ratios (Robinson, 1994). Shoot development may be slightly retarded, which is reflected in lower shoot biomass of barley grown under heterogeneous nutrient availability compared to the controls (Drew and Saker, 1975). This shoot growth inhibition was attributed to initial nutrient deficiencies until the roots reached the nutrient-rich zone (Drew and Saker, 1975). In addition, placement of N fertilizer (calcium ammonium nitrate) at 10 cm depth compared to surface mixing significantly increased dry matter production and grain yield at maturity of wheat grown in a loamy sand soil (Sharma and Chaudhary, 1983). However, there is a large variability between species (Robinson, 1994; Hodge, 2006) and genotypes (Sattelmacher et al., 1993; Forde and Lorenzo, 2001) in their shoot responsiveness, which depends on
differences in nutrient uptake and utilization efficiency (Sattelmacher et al., 1993). Nevertheless, LR proliferation may occur without limiting above ground growth (and yield).

7. Conclusion

In conclusion, plant roots are crucial in reducing soil erosion rates due to overland flow, as their presence affects soil properties such as aggregate stability, hydraulic function and shear strength. However, the effect of roots on soil erodibility is determined by RLD, and, thus, LR density and length, and the distribution of roots within the soil. It has been argued that a higher RLD would be particularly beneficial in reducing soil erosion rates and that fertilizer placement may be one way of achieving this. In nutrient poor soils, the local availability of high N or P may result in LR proliferation, although the response will depend on plant species, and may include a general increase in rooting depth or compensatory growth.

However, research is needed to determine whether the localized application of N and P at specific soil depths may represent a potential management technique to control erosion of degraded agricultural soils with low nutrient levels. Investigations should therefore aim to explore the effects of LR distribution at varying depths on soil properties, and, hence, soil erodibility. Trade offs between reduced erodibility due to LR proliferation and potential reductions in soil hydraulic conductivity need to be explored. In addition, responsiveness of different crops, grasses and shrubs (and different genotypes within a species) to heterogeneous nutrient availability should be determined. Moreover, suitable depth(s) for deep fertilizer placement should be established (with reference to practical implementation) and the effects of varying environmental conditions, such as soil type, slope and flow characteristics, on the erosion reducing effect of this potential management technique needs to be assessed.
8. References


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Figure 1. Key mechanisms by which plant roots influence soil physical properties
Figure 2. Effect of a localised supply of phosphate, nitrate, ammonium, and potassium on root form. Control plants received the complete nutrient solution to all parts of the root system. The other roots received the complete nutrient solution only in the middle zone (shaded), the top and bottom being supplied with a solution deficient in the specified nutrient. Adapted and reproduced with permission from Drew (1975) © John Wiley and Sons