

### Quantification of the effect of legume proportion in the sward on yield advantage and options to keep stable legume proportions (over climatic zones relevant for livestock production)

Andreas Lüscher, Matthias Suter, John Finn, Rosemary Collins, Francois Gastal

#### ▶ To cite this version:

Andreas Lüscher, Matthias Suter, John Finn, Rosemary Collins, Francois Gastal. Quantification of the effect of legume proportion in the sward on yield advantage and options to keep stable legume proportions (over climatic zones relevant for livestock production). [Contract] 7.3, 2014, 35 p. hal-01611404

HAL Id: hal-01611404

https://hal.science/hal-01611404

Submitted on 5 Oct 2017

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



### ANIMALCHANGE

#### SEVENTH FRAMEWORK PROGRAMME

### THEME 2: FOOD, AGRICULTURE AND FISHERIES, AND BIOTECHNOLOGIES



Grant agreement number: FP7 - 266018

#### **DELIVERABLE 7.3**

Quantification of the effect of legume proportion in the sward on yield advantage and options to keep stable legume proportions (over climatic zones relevant for livestock production)

**Abstract**: Grassland systems using mixed grass-legume swards have higher productivity and less environmental impact than their respective monocultures. The positive effect of legumes on yield is most pronounced with a mixture legume proportion of 30-60%.

Due date of deliverable: M36 Actual submission date: M36

Start date of the project: March 1<sup>st</sup>, 2011 Duration: 48 months

Organisation name of lead contractor: WBF-Agroscope

Contributors (Authors): Andreas Lüscher, Matthias Suter (WBF-Agroscope, John Finn (Teagasc), Rosemary Collins (AU-IBERS), François Gastal (INRA)

Revision: First submission

Dissemination level: PU



### **Table of Contents**

1.	SUMMARY	4
2.	LEGUME-GRASS SWARDS: A KEY TO SUSTAINABLE INTENSIFICATION OF AGRICULTURE	ON 4
3.	OVERYIELDING AND TRANSGRESSIVE OVERYIELDING IN GRASLEGUME MIXTURES	SS- 6
3.1	Overyielding	6
3.2	2 Transgressive overyielding	7
4.	POSITIVE LEGUME EFFECTS OPERATE OVER A WIDE RANGE LEGUME PROPORTIONS IN MIXTURES	OF 10
5.	OPTIONS FOR MAINTAINING LEGUME ABUNDANCE IN SWARDS	11
5.1	FERTILISER APPLICATION	13
5.2	P DEFOLIATION FREQUENCY	14
5.3	MORE COMPLEX MIXTURES	14
5	5.3.1 Number of species	14
5	5.3.2 Genetic diversity within species	15
5.4	SELECTION OF DIFFERENT LEGUME SPECIES AND BREEDING OF ADAPTED CULTIVARS	16
6.	LEGUMES ARE RESPONSIBLE FOR LARGE N INPUTS INTO THE SO VEGETATION SYSTEM BY SYMBIOTIC N <sub>2</sub> FIXATION	)IL- 17
7.	LEGUME-BASED FORAGE SYSTEMS CAN REDUCE NEGATIVE IMPAC	;TS
		24



7.1 FORAGE LEGUMES CAN REDUCE NITRATE LEACHING	21				
7.2 FORAGE LEGUMES CAN CONTRIBUTE TO REDUCED GREENHOUSE GAS EMISSIONS	22				
7.2.1 Methane	22				
7.2.2 Nitrous oxide	22				
7.2.3 Carbon dioxide	23				
7.3 FORAGE LEGUMES REDUCE CONSUMPTION OF NON-RENEWABLE ENERGY	24				
8. REFERENCES					
ANNEX 1: SUPPORTING INFORMATION	34				



#### 1. Summary

Grassland-based livestock production faces global challenges to meet the growing demand for meat and milk through increased production. This goal, however, has to be achieved using fewer resources in a more sustainable way than so far. Legumes offer great potential for coping with such challenges. Legumes have access to atmospheric nitrogen through symbiotic  $N_2$  fixation, and they exhibit numerous features that impact positively on the soil-plant-animal-atmosphere system. Their positive contribution to grassland systems is most pronounced in mixed swards with a legume proportion of 30-60%. The resulting benefits include higher productivity and increased protein self-sufficiency, lower costs of production, a reduced dependency on both fossil energy and inorganic N fertilizer, and lower quantities of harmful emissions to the environment. Due to increased productivity and reduced emissions, both per area cropped, these benefits are evident not only at the functional unit of managed land area but also at the unit of the final product. This makes legumes to a key option for sustainable intensification of agriculture.

However, legumes suffer from some limitations, one of which is a reduction in their proportion in mixed swards over time. In temperate regions of Europe, stable legume proportions in swards can be achieved by the following management practices: sowing more complex mixtures that include several grass and legume species and/or adapted cultivars, reducing N fertilizer input, and adjusting cutting and grazing rates. These practices need not necessarily reduce production levels. Growth restrictions of legumes due to low winter temperatures and/or limited water availability should be counteracted by selecting legume species and cultivars adapted to these environmental conditions. In conclusion, the promotion and development of legume-based grassland systems undoubtedly constitutes a highly relevant support for more environmentally sustainable and commercially competitive grassland-livestock systems.

# 2. Legume-grass swards: a key to sustainable intensification of agriculture

Under fertile agricultural conditions, monocultures of selected, highly productive grass species give high forage yields that are dependent on high inputs of fertilizer nitrogen (N) (Frame 1991; Daepp et al. 2001). Although there is strong demand to increase global food production, maximizing high forage yield in livestock systems through the use of industriallyproduced fertilizers has recently been challenged through several lines of evidence. First, substantial N losses in highly fertilized grass monocultures due to nitrate (NO<sub>3</sub>) leaching (Thomas 1992; Hooda et al. 1998; Ledgard et al. 2009; Peyraud et al. 2009) pose a threat to human health as leachates can reach groundwater resources used for drinking water (Squillace et al. 2002). Second, greenhouse gas emissions of 3.3 and 8.6 kg CO<sub>2</sub>equivalents are calculated per kg of urea-N and ammonium nitrate-N, respectively, for production and transport to the regional storehouse (global warming potential of over 100 years according to Christensen et al. 2007; database ecoinvent data v2.2 2010). Third, each kg of inorganic N produced in the industrial Haber-Bosch process consumes large amounts of energy (Kaltschmitt and Reinhardt 1997; Kitani et al. 1999). It is axiomatic that current levels of crop and livestock production have to be maintained or even to be increased; however, the manifold deleterious side-effects of intensive N fertilizer application to natural ecosystems call for more sustainable production than so far (Guckert and Hay 2001;



Rockström *et al.* 2009). The need to foster sustainable, resource-efficient yet productive agricultural systems poses a new challenge not only to European countries but also to other regions worldwide.

The use of mixtures instead of monocultures in grassland systems can be expected to be a promising strategy to increase yield. Established theory suggests that plant communities with higher species number (richness) are expected to (i) utilize available resources better due to species niche complementarity, (ii) have a higher probability of showing positive interspecific interactions, and (iii) may contain highly productive species that dominate the community (selection effect) (Tilman 1999; Loreau and Hector 2001; Loreau et al. 2001). Indeed, many experiments in nutrient-poor grasslands have shown that biomass production was enhanced in species-rich communities, compared to the average yield of the respective monocultures (Spehn et al. 2002; Hille Ris Lambers et al. 2004; Hooper and Dukes 2004; Hooper et al. 2005; Roscher et al. 2005; Marquard et al. 2009; Mommer et al. 2010). The presence of legumes in mixtures had generally a more pronounced positive effect on biomass yield than the presences of other functional groups (e.g. forbs, grasses) or species richness itself (Lambers et al. 2004; Thompson et al. 2005; Marquard et al. 2009). In a meta-analysis of 44 biodiversity experiments in which plant species richness was manipulated, Cardinale et al. (2007) found that mixtures, on average, outperformed monocultures by +77%. Yet, compared to the most productive monoculture, these mixtures showed a yield disadvantage of -12%. Transgressive overyielding (where mixtures outperform the best monoculture) (Trenbath 1974; Schmid et al. 2008) occurred in only 12% of cases and it took about five years to become evident. In an agronomic context, however, mixtures with transgressive overyielding are clearly preferred, as stakeholders can select the highest yielding species for monoculture cultivation, and any mixture performance has to compete against this high benchmark.

The combination of grasses with legumes in agricultural systems offers great potential for maintaining not only high levels of production, but doing so in an environmentally sustainable manner. Legumes exhibit numerous features that impact positively on the soil-plant-animal-atmosphere system (Castle *et al.* 1983; Hooda *et al.* 1998; Dewhurst *et al.* 2009; Ledgard *et al.* 2009; Peyraud *et al.* 2009; Soussana *et al.* 2010; Jensen *et al.* 2012) and comprehensive research has established their positive contribution to grassland systems in terms of biomass yield (Kirwan *et al.* 2007; Nyfeler *et al.* 2009; Finn *et al.* 2013) and harvest quality (Frehner *et al.* 1997; Nyfeler *et al.* 2011). Regarding quality, forage from grass-legume mixtures leads to an improved balance of the protein-energy ratio and increased digestibility of animal diet compared to grass or legumes alone (Sleugh *et al.* 2000; Marshall *et al.* 2004; Baumont *et al.* 2008; Dewhurst *et al.* 2009). The resulting benefits are higher productivity and increased protein self-sufficiency, lower costs of production, a reduced dependency on fossil energy and mineral N fertilizers, and lower quantities of harmful emissions to the environment (greenhouse gases and nitrate).

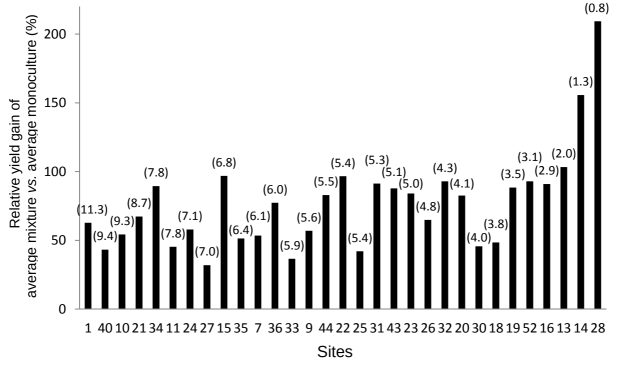


# 3. Overyielding and transgressive overyielding in grass-legume mixtures

#### 3.1 Overyielding

One of the most comprehensive agrodiversity experiments ever conducted involved 31 sites in 17 countries under the auspices of the COST Action 852, 'Quality legume-based forage systems for contrasting environments' (www.cost.eu/domains\_actions/fa/Actions/852). The geographical range of sites spanned a gradient of climate from Atlantic to continental, and from temperate to arctic (see Annex 1 for a list of included sites). Under conditions typical of cutting regimes, it was tested whether higher yields can be achieved with grass-legume mixtures containing four species as compared to the respective monocultures (Kirwan et al. 2007; Nyfeler et al. 2009). The four species represented four functional groups, which were chosen so as to maximize beneficial interspecific interactions: a fast-establishing grass, a fast-establishing legume, a temporarily-persistent grass, and a temporarily-persistent legume. The grass-legume combinations were intended to extend the method of nitrogen acquisition, as legumes have access to the unlimited resource of atmospheric nitrogen through symbiotic N<sub>2</sub> fixation (Hartwig 1998; Soussana et al. 2002), while fast/persistent combinations were intended to maximize sward cover by varying patterns of species development. The legume species examined were Trifolium pratense L. (red clover, 29 sites), Trifolium repens L. (white clover, 26), Medicago sativa L. (lucerne, 3), Medicago polymorpha L. (burr medic, 2), and Trifolium ambiguum M. Bieb. (Caucasian clover, 2) (Finn et al. 2013), meaning that legumes other than the widely investigated white and red clover were also included. Stands were established in two sowing densities and with greatly varying relative abundances of species, and received N fertilisers according to the general productivity level of sites. Across the three years of experimental duration, the yield of sown species (total yield excluding weed biomass) in mixtures was higher than the average monoculture in 99.7% of comparisons, with mixtures having, on average, 77% more yield than the average of monocultures (Fig. 1). Overyielding in harvest yield of mixtures appeared already in the first year (being 62%) and was maintained for at least three years (82% and 116% in second and third years, respectively) (Finn et al. 2013). It was most remarkable that overvielding occurred at all sites, which spanned a latitudinal range from 40°44'N (Sardinia, Italy) to 69°40'N (Tromsø, Norway), indicating that the relative benefit of yield was not restricted to highly productive sites (Fig. 1, to the left), but was equally apparent to the same relative degree at low yielding sites (Fig. 1, to the right), e.g. arctic or Mediterranean sites (Sturludóttir et al. 2013).





**Fig. 1.** Relative benefit of average mixture yield of sown species *versus* average yield of the respective monocultures over the whole experimental duration in %. Sites are arranged in order of decreasing means of monoculture yields (values above bars in t ha<sup>-1</sup> year<sup>-1</sup>), indicating that the mixture overyielding was not restricted to high yielding sites (to the left). Site names to numbers and further site information are given in Annex 1.

#### 3.2 Transgressive overyielding

While overyielding (the degree to which mixture yield exceeds the *average* yield of monocultures) was prevalent in the pan-European COST 852 experiment, it was most remarkable that transgressive overyielding (TO) was evident at the majority of sites (mixtures outperform the best monoculture, Finn *et al.* 2013). Across all experimental years, the total annual yield (sown species plus weed biomass) of mixtures exceeded that of the best monoculture in about 60% of sites, with the benefit of mixtures being 7% (Fig. 2). However, if yield of sown species (excluding weed biomass) was considered, TO was significant at about 70% of sites, with a mixture benefit of 18%. This is a highly valuable result because it demonstrates that mixtures of agronomically relevant grass-legume species outperform even the best monoculture that is usually selected to achieve maximal yields (often a grass species in highly fertilized monoculture systems).



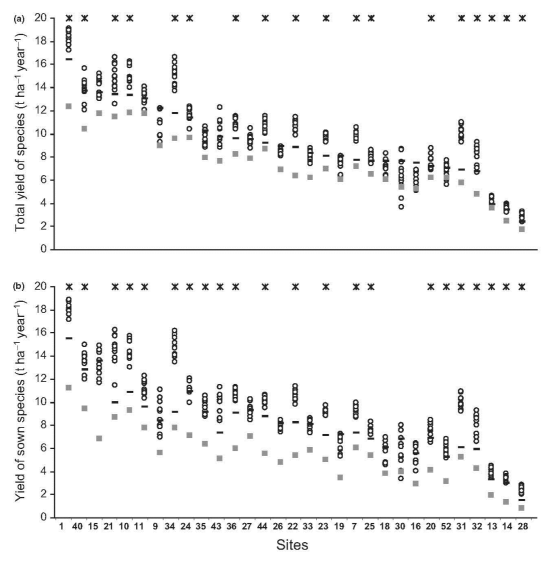


Fig. 2. Average annual yield (dry matter) over the whole experimental duration of (a) total yield and (b) yield of sown agronomic species only (excludes weeds) at each of 31 sites. For all communities, these data are averaged across seed density and across years per site. Sites are arranged in order of decreasing total yield of the best-performing monoculture. Open circles represent each of the 11 mixture communities that differed in their relative abundance at sowing; horizontal bars represent the yield of the best-performing monoculture; squares represent the mean monoculture performance. Significant transgressive overyielding is indicated by an asterisk (ж) over a site at the top of each panel. Note that not all of these sites had 3 years of yield data (see Table A1, Annex 1).

This figure is provided (with permission) from Finn et al. (2013). Figure number changed and legend shortened.

For example, at the Swiss site of the experiment, grass-clover mixtures fertilized with 50 or 150 kg of N ha<sup>-1</sup> year<sup>-1</sup> had maximal yields of 15.2 t and 16.1 t ha<sup>-1</sup> year<sup>-1</sup>, respectively, and thereby exceeded significantly the yields of the heavily fertilized (450 kg of N ha<sup>-1</sup> year<sup>-1</sup>) monocultures of the productive grass species *Lolium perenne* L. (perennial ryegrass, 12.0 t dry matter ha<sup>-1</sup> year<sup>-1</sup>) and *Dactylis glomerata* L. (cocksfoot, 13.4 t dry matter ha<sup>-1</sup> year<sup>-1</sup>, Nyfeler *et al.* 2009). With respect to all European sites, the high degree of TO appeared in the first year, persisted for at least three years and was not restricted to high yielding sites (Fig. 2) (Finn *et al.* 2013). Thus, similar to overyielding, TO was prevalent also at the low



yielding sites, which became directly apparent when computing the ratio of mixture yield to the yield of the best performing monoculture (Fig. 3).

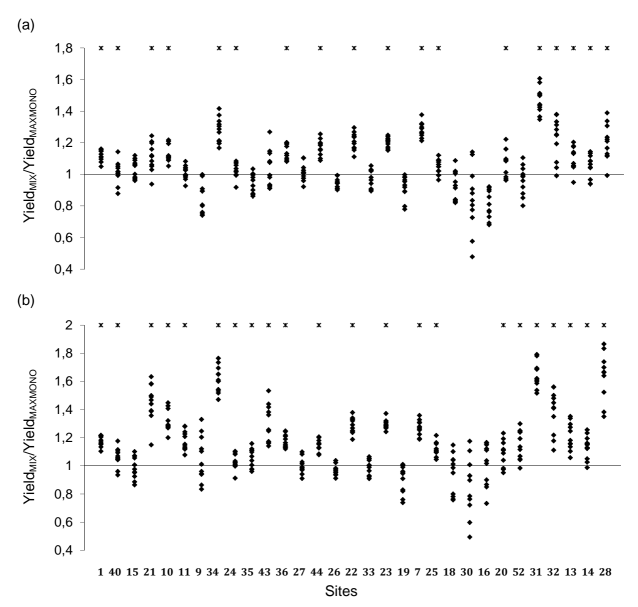


Fig. 3. Ratio of yield of each mixture community to yield of the best-performing monoculture (across years) at each site based on a) total yield and b) yield of sown species. Each point represents the sum of aboveground biomass over multiple years. Significant transgressive overyielding is indicated by an asterisk (x) over a site at the top of each panel. Sites arranged in order of decreasing total yield of the best-performing monoculture (as in Fig. 2).

This figure is provided (with permission) from Finn et al. (2013). Figure number changed and site numbers added.

The benefits in harvest yield of agronomically-relevant grass-legume mixtures can be attributed to several reasons. First and most importantly, the mixture swards have access to symbiotically-fixed  $N_2$  from the atmosphere through the legume component. Detailed analyses at one individual site of the pan-European COST 852 experiment (Nyfeler *et al.* 2011) demonstrated that symbiotic  $N_2$  fixation of legumes is responsible for large quantities



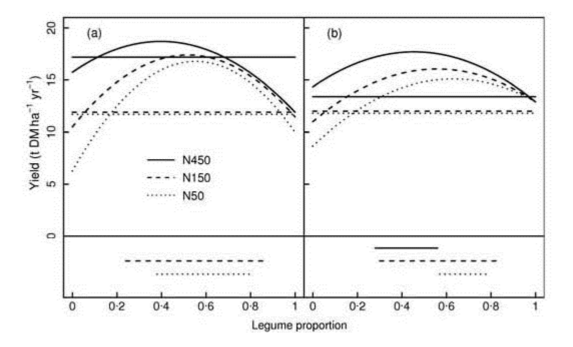
of N input into the system (see also chapter 6 of this deliverable), in full agreement with data from other, comparable systems (Boller and Nösberger 1987; Zanetti et al. 1997). Symbiotic N<sub>2</sub> fixation can only play an important role for overyielding if the legumes' growth and N<sub>2</sub> fixation perform well, and if nitrogen is a main limiting resource (Hebeisen et al. 1997). In particular N limitation is common in agronomic systems that target high forage yield and quality. For example, the maximum levels of 150 kg N ha<sup>-1</sup> year<sup>-1</sup> applied at the productive sites of the pan-European experiment (Finn et al. 2013) have been shown to limit the performance of agronomically relevant grass species when grown in monoculture (Daepp et al. 2001; Nyfeler et al. 2009). Second, trait interactions other than between N2 fixing legumes and non-fixing grasses have played a relevant role, because symbiotic N2 fixation cannot explain the highly significant yield advantage of mixtures over legume monocultures (detailed results in Nyfeler et al. 2009). Rooting depth between some of the grasses and legumes in mixtures differs and enhanced nutrient uptake from a larger soil pool has been suggested to contribute to overyielding in mixtures (van Ruijven and Berendse 2003; de Kroon 2007; Roscher et al. 2008; de Kroon et al. 2012; Gastal et al. 2012). Third, agronomic species as used in the COST 852 experiment have been selected for high quantity and quality of yield. This contributes to a reduction in the magnitude of yield differences across monocultures of the different species, and reduces the expected negative effect (in comparison to the bestperforming species) on mixture yield due to combining higher-yielding with lower-yielding mixtures. Fourth, mixtures as set-up in COST 852 strongly benefit from the combination of fast-establishing with more slow-growing, but temporally-persistent species (Kirwan et al. 2007; Finn et al. 2013); again, such a response is well recognised in other types of grassland (McKane et al. 1990). Fifth, characteristic within-season growth patterns that favour the grasses in spring during reproductive growth (Daepp et al. 2001) and the legumes in summer when temperatures are high (Vinther and Jensen 2000; Lüscher et al. 2005) could lead to temporal niches within the growing season that contribute to overyielding. In summary, these findings and underlying causes suggest that grass-legume mixtures offer great potential for increased production even at relatively low levels of species richness.

# 4. Positive legume effects operate over a wide range of legume proportions in mixtures

Studies that quantify the effects of species relative abundance on harvest yield are scarce, as most experiments assessing diversity effects on biomass production were set up with equal species proportions (e.g. "BioDepth" Hector et al. 1999; the "Jena experiment" Roscher et al. 2004). Again, the pan-European grassland experiment of COST 852 allows an assessment of the effects of legume proportion on mixture yield, as mixture stands were established in greatly varying species proportions. It appeared that increasing evenness (the degree of equal species proportion) had a significant positive effect on harvest biomass yield (Kirwan et al. 2007; Finn et al. 2013), indicating that mixtures with 50:50% of grasses and legumes were, on average, amongst the best performing. More importantly, the positive mixture effect was persistent over a large range of legume proportion, as e.g. the yield of communities dominated by one species (70% sown proportion of one species) was comparable to that of the most even community (Finn et al. 2013), and this feature persisted over three years of experimental duration. Detailed analyses at the Swiss site demonstrated that the maximum yield of the mixtures was reached with a clover percentage in the sward between 36 and 70%, depending on the year and the N fertilizer treatment. Averaged over three years, clover proportions in mixtures ranging between 30 and 80% produced mixture yields higher than the best performing monoculture (Fig. 4, stands fertilised with 150 kg of N



ha<sup>-1</sup> year<sup>-1</sup>) (Nyfeler *et al.* 2009). Thus, it did not matter whether the grass-clover mixtures were compared with the clover or grass monocultures: they were better than any of the monocultures. This result has high relevance as it highlights both the temporal stability of the mixture effect and its robustness over species proportions. This stability of high mixture yield in the face of compositional change is a most important feature for land managers and farmers because the benefit of grass-legume mixtures can be achieved despite a considerable change in relative proportions of the involved species (Connolly *et al.* 2013; Finn *et al.* 2013).



**Fig. 4.** Predicted mixture yield for increasing legume proportions and the three N fertilisation levels in the second year (a) and over all 3 years (b). Predictions are based on regression analysis, and are displayed for mixtures with equal proportions of the two legume species and equal proportions of the two grass species. Highest monoculture yield is indicated by horizontal lines<sup>1</sup> (above the x-axis). Horizontal lines below the x-axis indicate the range of significant transgressive overyielding ( $P \le 0.05$ ), that is, mixture yield being significantly higher than the highest monoculture yield.

<sup>1</sup>In the second year, *T. repens* at N50 and N150 and *D. glomerata* at N450; over all 3 years *T. pratense* at N50 and N150 and *D. glomerata* at N450

This figure is provided (with permission) from Nyfeler et al. (2009). Figure number changed and legend shortened.

### 5. Options for maintaining legume abundance in swards

Legumes have a distinct competitive advantage in N-limited systems (Hartwig 1998). Grasses supply their demand for N solely by uptake of mineral N in the soil; if soil N levels are low, then the growth rate of grasses will be slow. In contrast, legumes avoid N deprivation by supplementing mineral-N uptake with symbiotic  $N_2$  fixation, thereby retaining a relatively high growth rate even in an environment with low levels of soil N (Woledge 1988; Schwinning and Parsons 1996a). However, where mineral N is abundant,  $N_2$  fixation is

energetically costly and  $N_2$  fixers tend to be outcompeted by non-fixing species (Soussana and Tallec 2010).

The process of N<sub>2</sub> fixation carried out by legumes adds variable amounts of N to the soil, depending mainly on the amount of legume present but also on environmental constraints (Wu and MeGechan 1999). Variability in the processes of N<sub>2</sub> fixation by legumes, transfer via microbial processes, and uptake by associated grasses result in a degree of uncertainty in predicting the soil N status in mixed swards. Schwinning and Parsons (1996a, 1996b) described an intrinsic oscillatory component of variation in grass-legume systems, which, together with external stimulants of legume yield variation, interacts to produce a random element to species balance. The key to understanding the source of yield variability in grasslegume mixtures is the difference in these functional groups' response to N in the soil. The contrasting responses of grasses and legumes to the availability of soil N provides one explanation for their sustainable coexistence: in conditions of low soil N legumes have a growth advantage and should exclude grasses; in conditions of higher soil N grasses have a competitive advantage and should exclude legumes (Soussana and Tallec 2010). Thus, in conditions of stable soil N, Schwinning and Parsons (1996a) argued that these species could not coexist. However, soil N level is a dynamic variable that not only affects plants, but is also affected by them. In conditions of low soil N the dominance of legumes causes N to accumulate in the soil, which eventually promotes dominance of the grass component (the so-called 'paradox of enrichment'). Schwinning and Parsons (1996a) stated that this negative feedback control of soil N status maintains the competitive balance between grasses and legumes in mixed swards. However, the control of soil N is a relatively slow process, and models have shown that grasses and legumes can oscillate in dominance for many years before equilibrium is reached (Schwinning and Parsons 1996b). Moreover, much of the evidence concerning fluctuations in grass-legume abundance is derived from ryegrass-clover mixtures. Evidence from other systems is sparse, and mixtures such as grass-lucerne could be more unstable and less flexible.

For farmers, variability in legume abundance translates into uncertainties in herbage production and forage quality, so for this reason it is vital to understand the complexities of grass-legume interactions. Control of the synchrony of N supply and demand in grasslegume swards appears to be the key to achieving sustainable grassland agriculture. The issue may be approached through improving plant uptake of N from the soil, and by manipulating soil N supply. The former can be approached through manipulation of the species content of grassland mixtures via the use of perennial forage species with different spatial and temporal nutrient uptake patterns (Daepp et al. 2001; Lüscher et al. 2005). Achieving the latter objective is more problematic, due to large gaps in current knowledge of plant-driven N fluxes (Høgh-Jensen 2006). For example, legume species differ substantially in the amount of N they add to grassland systems through N2 fixation, and in the amount of inorganic soil N they take up. For example, the widely-grown temperate forage legume derives a smaller proportion of N from atmospheric nitrogen than other forage legumes due to the high uptake of soil inorganic N by its deep and extensive root system (Blumenthal and Russelle 1996; Rasmussen et al. 2012). Furthermore, lucerne seems to transfer a lower proportion of fixed N to the rhizosphere and companion grass species than other legumes (Brophy and Heichel 1989; Lory et al. 1992).

Probably due to the challenging control of the synchrony of N supply and demand, legume proportions in mixed swards have been observed repeatedly to decrease over time (Frame 1986; Elgersma and Schlepers 1997; Guckert and Hay 2001), the decline in relative legume abundance being considerably accelerated under higher levels of N fertilisation (Hebeisen *et al.* 1997; Aydin and Uzun 2005; Nyfeler 2009). This matches well the remarkable decrease in legume proportion of mixture swards throughout the three years of experimental duration in



the pan-European COST 852 experiment (Finn *et al.* 2013). The big challenge for legume-based grassland-livestock systems is therefore to maintain the proportion of legumes within an optimal range. Four of the most promising strategies are outlined below.

#### 5.1 Fertiliser application

Sward management strategies that reduce N fertilizer input can stabilise the proportion of white clover in permanent grassland (Schwank et al. 1986; Ledgard et al. 2001). The effectiveness of reduced N fertilisation in regulating the proportion of white clover is, for example, evident from the Swiss Free-Air CO<sub>2</sub> Enrichment (FACE) experiment (Hebeisen et al. 1997; Zanetti et al. 1997): averaged over the first three years, the contribution of white clover was 57% at low levels of N fertilisation (100 and 140 kg N ha<sup>-1</sup> year<sup>-1</sup>, combined with frequent defoliation), whereas it was only 14% with high N fertilisation (420 and 560 kg N ha<sup>-1</sup> year<sup>1</sup>, combined with infrequent defoliation). Further evidence comes from the Swiss site of the earlier mentioned COST 852 project, where three different levels of mineral-N (50, 150, 450 kg N ha<sup>-1</sup> year<sup>-1</sup>) were applied to the grass-clover mixtures (two grass and two clover species) managed for three consecutive years. Here, N fertilisation strongly affected clover proportion: averaged over all mixtures receiving 50 kg N ha<sup>-1</sup> year<sup>-1</sup>, the two clover species (sum of both) achieved 42, 56, and 24% proportional biomass in years 1, 2, 3, respectively; however, fertilized with 450 kg N ha<sup>-1</sup> year<sup>-1</sup>, clover proportions were only 32, 24, and 5%. Similar results were found by Nassiri and Elgersma (2002), where in only one growing season the application of 150 kg N ha<sup>-1</sup> year<sup>-1</sup> reduced the clover content in dry matter harvest of ryegrass-white clover mixtures to 12% compared to 43% without N application. In a similar system with respect to species and N application, the average difference in the mixture clover proportion between the N treatments was 25% (59% vs. 34% with zero N and 165 kg N ha<sup>-1</sup> year<sup>-1</sup>, respectively; Elgersma et al. 2000). Most recently, Oberson et al. (2013), assessed the dynamics of N<sub>2</sub> fixation and N transfer in grass-clover leys under different cropping systems (organic, conventional). Irrespective of the cropping system, clover proportion was greater under reduced N fertilisation and reached 47% in the unfertilised control, while it decreased to 30% in a reference treatment that received 120 kg of N ha<sup>-1</sup> year<sup>-1</sup> (averaged over two years). It is most relevant to note that high productivity levels can still be maintained at moderate to intermediate levels of N fertilisation. For example, Nyfeler et al. (2009) found yields of grass-clover mixtures fertilized with 50 or 150 kg of N ha<sup>-1</sup> year<sup>-1</sup> to be as high (Fig. 4a) or even higher (Fig. 4b) than the yields of the heavily fertilized (450 kg of N ha<sup>-1</sup> year<sup>-1</sup>) monocultures of the grasses Lolium perenne and Dactvlis alomerata.

Regarding nutrients other than N, in particular phosphorus (P), potassium (K), and sulphur (S), their availability is crucial for growth and  $N_2$  fixation of legume species (Brown *et al.* 2000; Tallec *et al.* 2009; Batterman *et al.* 2013). For example, both  $N_2$  fixation and plant growth have been shown to be constrained by P in native, phosphorus-poor soils of the tropics (Batterman *et al.* 2013) and in nutrient solution (Almeida *et al.* 2000). In a similar way, limitations in the availability of soil S significantly restricted the growth of clover in experimental ryegrass-white clover mixtures receiving 180 kg N ha<sup>-1</sup> year<sup>-1</sup> (Tallec *et al.* 2009).

In conclusion, the available evidence suggests that, in temperate grassland systems, legume abundance in mixtures can be balanced through adjusting N fertilizer application. It is most promising that even substantial reduction in N fertilizer application is not expected to impact negatively on productivity, because a potential N deficiency due to reduced fertilisation is



counteracted by increased symbiotic  $N_2$  fixation of legumes in mixture swards (see chapter 6).

#### **5.2** Defoliation frequency

In well-managed grassland systems, defoliation frequency is closely related to fertilisation rates. Nevertheless, adapting the frequency of defoliation provides an additional management strategy to adjust the legume proportion in swards. Most evidence to this option comes from white clover, the most promising legume for mixture swards in temperate, northern and upland Europe (Guckert and Hay 2001). White clover is resistant to frequent defoliation because it has stolons that grow closely at the ground surface and escape the harvest of petioles and leaves. Therefore, more frequent cutting will reduce competition for light from companion grass species and thereby promote growth of clover. The earlier-cited FACE experiment manipulated not only CO<sub>2</sub> partial pressure and N fertilisation, but also defoliation frequency. It became evident that, throughout all treatments and years, increased defoliation raised the clover proportion in mixed swards (Hebeisen et al. 1997). Further studies confirm the relationship between cutting and relative abundance of white clover (Schwank et al. 1986; Elgersma and Schlepers 1997). For example, Elgersma and Schlepers (1997) examined grass-clover mixtures that were cut in two frequencies over three years: the more frequently harvested sward had finally clover proportions of 50%, while it was only 42% with less frequent cutting. Similarly, Wen and Jiang (2005) found a tendency for higher proportions of white clover in mixed swards with increased cutting frequency. Their experiment ran for two years and tested a ryegrass-white clover pasture under four different cutting frequencies (8 cuts year<sup>-1</sup>, 4, 2, 1): the final proportion of white clover was ca. 20% in the most frequently cut plots, while it was less than 10% with one cut per year. Regarding pasture legumes other than white clover, only few information is available to effects of defoliation frequency on mixture legume proportion (but see Peterson et al. 1994; Giambalvo et al. 2011).

Although the influence of defoliation frequency on clover proportion in mixed swards appears to be smaller than the effects of N application (Schwank *et al.* 1986; Hebeisen *et al.* 1997), adjustment of cutting and grazing rates is one of several options to keep clover proportions in swards at stable levels.

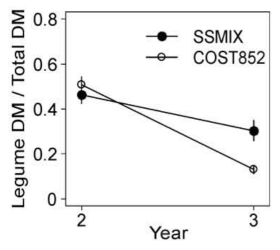
#### 5.3 More complex mixtures

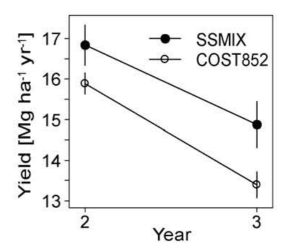
#### 5.3.1 Number of species

A further option to stabilise legume abundance in mixtures is an optimised species composition of seed mixtures that prevents competitive dominance. Improved seed mixtures will require decisions on how many and which species (or variety) to include, and which proportions of the species to choose. The potential of such improved mixtures is, for example, evident from the Swiss site of the COST 852 project, where, besides the experimental four-species mixtures (Kirwan *et al.* 2007; Nyfeler *et al.* 2009), Swiss Standard Mixtures (SSMIX, Suter *et al.* 2012) were also examined. These mixtures contain more species (up to eight) and their composition (relative and absolute abundances of species)



has been continuously improved over several decades to maintain stable species composition based on experiments and observations on farms. Over the three years of experimental duration, the decline in clover abundance in the SSMIX was much smaller than that in the four-species mixtures of the pan-European experiment (Fig. 5a) and, at the same time, SSMIX were more productive throughout (Fig. 5b).





**Fig. 5a.** Legume proportion of total spring-DM (dry matter) yield of COST852 and SSMIX mixtures (bars = 1 SE).

**Fig. 5b.** Annual yield of COST852 and SSMIX mixtures for the second and third experimental year (bars = 1 SE).

Figures are provided (with permission) from Suter et al. (2010). Figure numbers changed.

Current concepts suggest that combining species with high "combining ability" in more complex mixtures would result in better resource exploitation through niche differentiation (Aarssen 1983; Hill 1990), and empirical evidence supports this idea (Hooper *et al.* 2005; Roscher *et al.* 2005; Nyfeler *et al.* 2009; Finn *et al.* 2013). In comparison, combining species or genotypes in mixtures that have comparable "competitive abilities" would result in more balanced and stable mixtures (Evans *et al.* 1985, 1989; Turkington 1989a; Lüscher *et al.* 1992). Both existing theory and experimental evidence on combining species in more complex mixtures is intriguing and call for extended research on how such mixtures can stabilise legume proportion and increase total yield in agricultural grassland systems.

#### 5.3.2 Genetic diversity within species

Not only do species differ in their competitive abilities, so also do cultivars within species. For example, Suter *et al.* (2007) found that the realized species composition of the established sward differed tremendously, depending on which cultivars were chosen for the seed mixture. In that context, the presence of a large degree of genetic diversity for morphophysiological traits in natural plant populations might suggest that the use of blends of several cultivars and thus of composite populations with a high degree of heterogeneity would also be advantageous under agricultural conditions. However, there is relatively limited evidence to support this idea. It has been demonstrated that greater heterogeneity in genotype morphology in white clover populations failed to translate into higher dry matter production (Annicchiarico and Piano 1997), comparable to results obtained with cultivar mixtures on perennial ryegrass (Surault *et al.* 2010). A similar result was obtained by Williams *et al.* (2003), also for white clover, although there was some evidence that yield stability over the duration of the experiment was increased in the mixture containing a blend



of cultivars. The impact of increased heterogeneity in legume populations on their performance in grass-legume mixtures was also investigated by Collins et al. (2004). It was hypothesised that 'wide genetic base' (WGB) composite populations of red and white clover, comprising mixtures of seed of many varieties, would provide a platform for sustained positive effects on harvest yield in grass-legume mixtures; however, no yield advantage was observed in mixtures with WGB populations (Collins et al. 2004). A detailed study of the influence of the WGB legumes on sward dry matter productivity and unsown species invasion in complex mixtures was carried out at the COST 852 site in southern Sweden (Frankow-Lindberg et al. 2009). It was found that the impact of the WGB populations on both these ecosystem properties was small compared with the diversity effect brought about by the mixing of species. This could be due to the fact that traits are more likely to be similar within species (even in populations with deliberately enhanced heterogeneity) than between different species. Another explanation for the weak effect of the WGB composites might be a reduction in intra-specific genetic diversity caused by natural selection over time of the most adapted individuals for the site in challenging environments such as the Swedish COST site. The phenomenon of genetic shift was observed in a genome-wide analysis of population genetic differentiation in the WGB and single-cultivar germplasm in Northern European COST 852 sites (Collins et al. 2012). Thus, the use in diverse species mixtures of WGB populations containing site-adapted germplasm may produce more positive results in terms of sward yield.

In summary, these results demonstrate that the species composition of multi-species seed mixtures offers a multi-factorial opportunity for optimization, but that the use of blends of cultivars might provide only limited benefits. Further research is needed to clarify the role of blends of cultivars in mixed grass-legume systems before such on option can generally be recommended.

# 5.4 Selection of different legume species and breeding of adapted cultivars

Under climatic conditions of low winter temperatures, limited water availability or severe drought, the long-term growth of red and white clover appears to be restricted (Sturludóttir et al. 2013; unpublished results). Here, selecting different legume species adapted to harsh environmental conditions and/or specific breeding programs for suitable cultivars can significantly contribute to higher persistence of legumes in grassland, and thus to more stable legume abundance in swards (Serraj et al. 1999; Patto et al. 2006; Carbonero et al. 2011). For example, evaluation of two white clover cultivars at twelve sites over a European gradient revealed significant differences in clover mixture proportion following overwintering, much of the difference being due to the cultivars' varying response to temperature (daily means) (Wachendorf et al. 2001). Frankow-Lindberg (2001) tested nine genetically different populations of white clover for their physiological response to frequent sub-lethal frost in northern climates. She found not only relevant differences among populations, but concluded that there would be sufficient genetic variation in at least one population for further trait selection in the field. The research of Collins et al. (2012) on genetic change in legume germplasm (commercial cultivars and experimental populations) further supported this hypothesis; the authors found that measurable genetic shift occurred in both red and white clover over a relatively short time period when grown in cold sites in Northern Europe. The degree of genetic change that occurred depended on how well adapted the populations initially were to their growing environment, as well as on the climatic characteristics of the



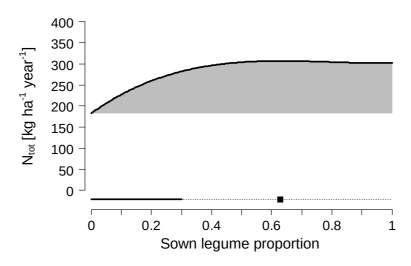
site itself. One practical implication of these results is that it would probably be beneficial for legume breeding programmes aimed at producing cultivars for challenging environments to consider collecting germplasm from survivor populations in these areas in order to reinforce existing breeding lines.

Newly selected legume species and cultivars should be evaluated in combined growth with grasses, either in single species combination or in more complex mixtures as outlined above. Current European initiatives Legume Plus (<a href="www.legumeplus.eu">www.legumeplus.eu</a>) and MultiSward (<a href="www.multisward.eu">www.multisward.eu</a>), all within the seventh frame framework program, deal with such requirements.

# 6. Legumes are responsible for large N inputs into the soil-vegetation system by symbiotic N<sub>2</sub> fixation

The major benefit of legumes to more sustainable but still productive grassland systems comes through their ability to incorporate large amounts of N into the system. In temperate and Northern grassland, amounts of symbiotically fixed N<sub>2</sub> (N<sub>sym</sub>) by legumes can range from 100 to 380 kg of N ha<sup>-1</sup> year<sup>-1</sup>, but values as high as 500 kg of N<sub>sym</sub> ha<sup>-1</sup> year<sup>-1</sup> have also been reported (Boller and Nösberger 1987; Ledgard and Steele 1992; Carlsson and Huss-Danell 2003; Lüscher et al. 2013; Oberson et al. 2013). Amounts of N<sub>sym</sub> by crop legumes are smaller and range between 30 and 150 kg of N ha<sup>-1</sup> year<sup>-1</sup>, with maximal values around 200 kg of N<sub>svm</sub> ha<sup>-1</sup> year<sup>-1</sup> (Peoples et al. 2009). In agricultural grass-legume mixtures, legumes can acquire more than 80% of their own N demand through symbiosis (Boller and Nösberger 1987; Zanetti et al. 1997; Nyfeler et al. 2011; Rasmussen et al. 2012; Oberson et al. 2013), and consequently, the relative availability of soil N increases for grasses due to 'N-sparing'. Moreover, in mixed grass-legume systems, amounts of 10-75 kg of N ha<sup>-1</sup> year<sup>-1</sup> are transferred from legumes to grasses (Høgh-Jensen and Schjoerring 1997; Zanetti et al. 1997; Nyfeler et al. 2011), while the amounts of transferred N depend on the donor and the receiver plant species (Pirhofer-Walzl et al. 2012). In a recent analysis of the amount of symbiotic N<sub>2</sub> fixation within the pan-European COST 852 project (Suter et al. 2013; unpublished results), the average amount of N<sub>sym</sub> in 50:50% grass-legume mixtures across all evaluated sites was 106 kg ha<sup>-1</sup> year<sup>-1</sup>; however, maximal N<sub>sym</sub> in mixtures at individual sites and years was as high as 325 kg ha<sup>-1</sup> year<sup>-1</sup>. Further, across sites, mixtures with one third of legume proportion attained amounts of N<sub>sym</sub> that were not significantly different from maximal amounts in mixtures with much higher legume proportion (including legume pure stands), suggesting that grassland with largely varying legume proportions can achieve substantial gain from the process of symbiotic N<sub>2</sub> fixation (Fig. 6).





**Fig. 6.** Total nitrogen yield ( $N_{tot}$ ) as affected by sown legume proportion at the first year of a multisite grassland experiment across Europe. The curved bold line displays the predicted  $N_{tot}$  across 15 sites, which spanned a gradient of climate from Atlantic to continental, and from temperate to arctic. The grey shaded area indicates N derived from symbiotic  $N_2$  fixation of legumes across all sites based on the N difference method (Ledgard and Steele 1992). The horizontal bold line at bottom indicates the range of legume proportion for which  $N_{tot}$  was significantly smaller ( $P \le 0.05$ ) than at maximum ( $\blacksquare$ ); consequently, the dotted line displays the non-significant range.

In mixtures containing ryegrass, cocksfoot, red and white clover and being fertilised with 150 kg N ha<sup>-1</sup> year<sup>-1</sup>, Nyfeler et al. (2011) found maximum N<sub>sym</sub> values of 295 kg N ha<sup>-1</sup> year<sup>-1</sup> (average over two years, Fig. 7). More importantly, total harvested dry matter yield (Nyfeler et al. 2009) and total yield of harvested N (Nyfeler et al. 2011) were maximal at intermediate levels of clover proportion of 40-60% in mixture. Interestingly, Nyfeler et al. (2011) found stimulatory effects of the accompanying grasses on the symbiotic N<sub>2</sub> fixation activity of clover. This effect was so strong that (i) the amount of N from symbiosis was maximized not in pure clover stands but in mixtures with 60-80% of clovers, and (ii) a clover proportion of 40-60% in the mixture was sufficient to attain the same amount of N from symbiosis as in pure clover stands. The activity of symbiotic N<sub>2</sub> fixation of clover plants was very high in grass-dominated swards, where the availability of mineral N to clovers was limited because most of the N from mineral sources was taken up by the competitive grass component (Nyfeler et al. 2011). However, in clover-dominated swards (> 60% of clovers), the activity of symbiotic N<sub>2</sub> fixation was down-regulated. This was due to the clovers' adequate access to mineral N sources because of low grass abundance in addition to a significantly reduced N demand of the whole sward in clover-dominated stands, as these were less productive than well balanced mixtures (Nyfeler et al. 2009).



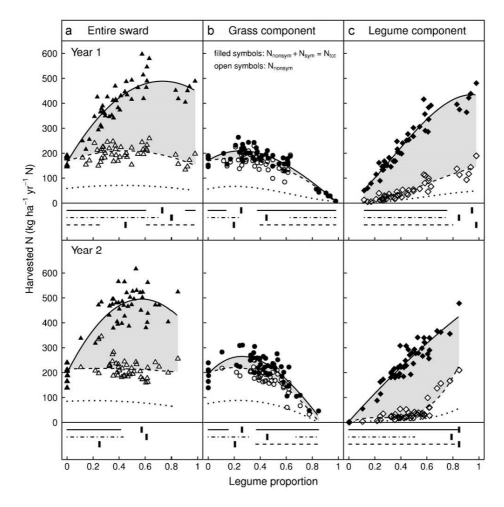


Fig. 7. Nitrogen yield and its fractions harvested in the entire sward (a) or in either its grass (b) or legume (c) component as affected by the legume proportion in moderately fertilised swards (150 kg N ha⁻¹ year⁻¹) for the first and second experimental year. Curves indicate N from fertilizer (N<sub>fert</sub>, · · ·) and N from non-symbiotic sources (N<sub>nonsym</sub>, - · -). N<sub>nonsym</sub> is the sum of N<sub>fert</sub> plus N derived from soil organic matter (N<sub>soil</sub>). The shaded area is the difference of N<sub>nonsym</sub> to total harvested N (N<sub>tot</sub>, —) and represents N from symbiotic sources (N<sub>sym</sub>), which comprises of N from symbiotic N₂ fixation in the legume component (N<sub>symfix</sub>), N<sub>symfix</sub> transferred to the grass component (N<sub>symtrans</sub>), and the sum of both in the entire sward. Horizontal lines below the curves indicate the range of legume proportions for which N<sub>nonsym</sub> (- - -), N<sub>sym</sub> (·-·-·-) and total harvested N<sub>tot</sub> (—) are significantly smaller ( $P \le 0.05$ ) than at maximum (▮).

This figure is provided (with permission) from Nyfeler et al. (2011). Figure number changed and legend shortened.

It has been suggested that the degree of symbiotic  $N_2$  fixation of legumes is controlled by a series of eco-physiological triggers and N feedback mechanisms from the individual plant to the ecosystem level (Schwinning and Parsons 1996b; Hartwig 1998; Soussana and Tallec 2010), with  $N_2$  fixation of legumes being largely regulated by the level of site productivity, i.e. the N sink strength (N-demand) of the whole system. Following this concept, legumes would account for the gap between N-demand (sink) and N-availability (source) from non-symbiotic N-sources by adjusting the degree of symbiotic  $N_2$  fixation (Hartwig 1998; Soussana and Tallec 2010; Lüscher *et al.* 2011). Based on experimental results, mechanistic modeling could tightly couple symbiotic  $N_2$  fixation in legume-based grassland systems to the N



demand at scales ranging from the individual plant to the grassland ecosystem (Soussana and Tallec 2010). For example, it was modeled that N uptake rates of legumes and grasses under varying soil N availability at the patch scale could lead to stable grass-legume coexistence at the community level (Soussana and Lafarge 1998).

The growth and ability of legumes to fix nitrogen may be hampered in climatic conditions with low winter temperatures and/or scarce precipitation (Nesheim and Boller 1991; Serraj et al. 1999). It is known from controlled experiments that low temperatures and severe water stress negatively affect the process of nitrogen fixation-nodulation (e.g. Roughley and Dart 1970) and nitrogenase activity (e.g. Cralle and Heichel 1982; Durand et al. 1987). However, only a few field studies have been published on symbiotic N2 fixation under marginal conditions such as high altitude (e.g. Bowman et al. 1996; Jacot et al. 2000a) or high latitude (e.g. Henry and Svoboda 1986; Sparrow et al. 1995). Jacot et al. (2000a, 2000b) studied symbiotic N<sub>2</sub> fixation of naturally occurring legumes and the N balance of the whole grassland ecosystem in species-rich semi-natural pastures in the Swiss Alps along a gradient from 900 m a.s.l. to the altitudinal limit of legume occurrence at 2600 m a.s.l. Investigated legumes were: Lotus alpinus (DC.) Schleicher, Lotus corniculatus L., Trifolium alpinum L., Trifolium badium Schreber, Trifolium nivale Sieber, Trifolium pratense L., Trifolium repens L., Trifolium thalii Vill., Vicia sativa L. Absolute amounts of N<sub>sym</sub> decreased significantly with increasing altitude (from 18 to 1 kg N ha-1 year-1) in conjunction with a reduction of the total productivity of the community and with a decrease in sward legume proportion from 15 to 4%. These features went along with a decline in summer precipitation, mean temperatures, and a decrease in soil pH. In the context of regulation of N<sub>2</sub> fixation, results of Jacot et al. (2000a) are in agreement with Hartwig (1998) and Soussana and Tallec (2010) that predict lower symbiotic N<sub>2</sub> fixation at low productivity levels. However, although absolute levels of N<sub>svm</sub> were closely related to productivity and pedo-climatic factors, the percentage of N<sub>sym</sub> in legumes was not (Jacot et al. 2000a): % N<sub>sym</sub> in legume plants (59 to 90%) was not diminished over the altitudinal range, most probably due to the high abundance of non-legumes (grasses, forbs) that competed strongly for mineral N in soil. Results of Jacot et al. (2000a) indicate a substantial contribution of symbiotic N<sub>2</sub> fixation to the legumes' N requirements over varying environmental conditions, suggesting that - within restrictions of low temperatures (Cralle and Heichel 1982) and severe water stress (Durand et al. 1987; Serraj et al. 1999) - the physiological process of symbiotic N<sub>2</sub> fixation in naturallyoccurring species is adapted to pedo-climatic conditions. Interestingly, results of N<sub>2</sub> symbiosis in the COST 852 project agree with such conclusion (Suter et al. 2013; unpublished results). Here, the proportion of N<sub>sym</sub> to the total N yield of sites was, on average, not correlated with site productivity (within the conditions of nitrogen application across sites, ranging from 0 to 150 kg N ha<sup>-1</sup> year<sup>-1</sup>), but negatively affected by low annual precipitation and minimal site temperature. This provides evidence from a European gradient of sites that agricultural grassland systems of largely differing productivity levels can benefit from N<sub>2</sub> fixation, given that legumes can be grown in mixtures and that symbiotic N<sub>2</sub> fixation remains active in legume plants.

These findings and the sink/source model of regulation of symbiotic  $N_2$  fixation at the ecosystem level have several critical implications for the exploitation of atmospheric  $N_2$  in grassland systems.

1. Grass-legume mixtures can potentially fix more N<sub>2</sub> from the atmosphere than legume monocultures.

There is no trade-off between high productivity and high gains of nitrogen from symbiotic  $N_2$  fixation because they are positively linked through N demand (sink). Consequently, sufficient availability of other nutrients, such as P, K, and S, is crucial for stimulating demand of N from



symbiosis (Sangakkara et al. 1996; Hartwig 1998; Brown et al. 2000; Tallec et al. 2009; Batterman et al. 2013).

2. Even though high inputs of N into the ecosystem from symbiosis can cause a risk for N losses into the environment (Loiseau et al. 2001; Scherer-Lorenzen et al. 2003), such N losses can be prevented by sufficiently great abundance of grasses in the sward to ensure a highly competitive uptake of mineral N from the soil (no less than 40% of sward biomass occupied by grasses) (Nyfeler 2009; Nyfeler et al. 2011; Lüscher et al. 2013) (see also Chapter 7).

# 7. Legume-based forage systems can reduce negative impacts of livestock production on the environment

The great opportunities of legumes for environmentally friendly yet productive grassland-livestock systems derive from different features: (i) increased yield, (ii) replacement of mineral-N fertilizer by symbiotically-fixed  $N_2$ , (iii) higher nutritive value and voluntary intake of forage, and (iv) greater livestock performance. Taken together, all of these effects create important environmental advantages of legume-based grassland husbandry systems. The advantages are evident not only at the sward level but also at the whole-farm level. In addition, the benefits apply also to the functional unit of managed land area and to the unit of the final product. This makes legumes to a key option for sustainable intensification of agriculture.

#### 7.1 Forage legumes can reduce nitrate leaching

In comparison to fertilized pure ryegrass stands (300 kg N ha<sup>-1</sup> year<sup>-1</sup>), the presence of legumes in mixed swards can reduce or stabilise nitrate (NO<sub>3</sub>) leaching to groundwater while, at the same time, reduce the need for mineral N fertilizers (Eriksen *et al.* 2004). For example, Vertès *et al.* (1997) found a 5 to 10% reduction of NO<sub>3</sub> leaching under grass-clover receiving no fertilizer compared to fertilized (250 kg N ha<sup>-1</sup> year<sup>-1</sup>) pure ryegrass pastures; both systems had comparable biomass yields (8.7 t ha<sup>-1</sup> year<sup>-1</sup> for ryegrass-white clover, 9.5 t ha<sup>-1</sup> year<sup>-1</sup> for pure grass). At similar stocking rate (3.3 cows ha<sup>-1</sup>) and milk yield per hectare, Ledgard *et al.* (2009) reported identical levels of N leaching (30 kg N ha<sup>-1</sup> year<sup>-1</sup>) for grass-clover pastures receiving no mineral N fertilizer as for pure ryegrass pastures fertilised with 160 kg N ha<sup>-1</sup> year<sup>-1</sup>. However, losses of nitrate under grazed grass-clover swards can increase with increased clover proportion (Schils 1994), supplemental N fertilisation (Ledgard *et al.* 1999; Ledgard *et al.* 2009) or high stocking rates (Simon *et al.* 1997; Eriksen *et al.* 2010). Regarding legumes other than clover, NO<sub>3</sub> concentrations in ground water were found to be lower under lucerne than under annual crops (Russelle *et al.* 2001; Russelle *et al.* 2007) and grass-clover mixtures (Ledgard *et al.* 2009).

However, very high inputs of N from symbiosis into grassland systems can increase the risk of nitrate leaching (Hooper and Vitousek 1998; Loiseau *et al.* 2001; Scherer-Lorenzen *et al.* 2003). Loiseau *et al.* (2001) reported higher annual leaching losses of N from lysimeters when swards were sown with pure white clover (28 to 140 kg N ha<sup>-1</sup>) than with pure perennial ryegrass (1 to 10 kg N ha<sup>-1</sup>); yet, much higher values were reported for bare soils (84 to 149



kg N ha<sup>-1</sup>). Nevertheless, it can be expected that N losses by leaching will be prevented as long as the percentage of grass in mixed grass-legume swards is well balanced to take up mineral N from the soil (no less than 40% of sward biomass occupied by grasses). For example, under a mowing regime and 150 kg N fertilizer input ha<sup>-1</sup> year<sup>-1</sup>, Nyfeler (2009) found an increased risk of nitrate leaching only if the percentage of legumes in the mixture exceeded 60-80%. Only few studies have assessed changes in soil nitrate content under mixtures containing legumes for periods of more than a few years. In the long-term, levels of soil nitrate might rise due to the mineralization of nitrogen-rich legume residues. During a five-year measurement period in an unfertilized experimental grassland, the mere presence of legumes resulted in increased soil NO<sub>3</sub>-N content but the legume effect on nitrate in leaching water did not change over time, suggesting no increased risk of N leaching (Oelmann *et al.* 2011).

# 7.2 Forage legumes can contribute to reduced greenhouse gas emissions

#### 7.2.1 Methane

Methane produced in the rumen is a large contributor to greenhouse gas (GHG) emissions by livestock systems (Tamminga et al. 2007; Rowlinson et al. 2008; Waghorn and Hegarty 2011). Legumes can contribute to reduced ruminal methane production per unit of intake. Ruminants fed legume forages generally emitted less methane than grass-fed animals, per unit of feed intake (McCaughey et al. 1999), although not in all cases (van Dorland et al. 2007; Hammond et al. 2013). Inconsistency of results between experiments can arise from difference in forage composition (forage species, stage of maturity, presence of condensed tannins) and animal genotypes. Condensed tannins (CT) may also be useful for reducing greenhouse gases (Kingston-Smith et al. 2010), as several studies have shown that CT reduced methane production in vitro. A recent meta-analysis revealed a general antimethanogenic effect of CT above 20 g kg<sup>-1</sup> dry matter in feeds (Jayanegara et al. 2012). Some effects of CT to lessening methane emissions were also reported from in vivo studies with Onobrychis viciifolia Scop. (sainfoin, Waghorn 2008), Lotus corniculatus L. (birdsfoot trefoil, Woodward et al. 2004) and Hedysarum coronarium L. (sulla, Woodward et al. 2002). Both sainfoin and sulla have higher CT contents than the widely used red and white clover and lucerne, which are low in CT (Häring et al. 2007; Häring et al. 2008). Therefore, CT related effects on methane emissions can only be achieved by cropping the above mentioned CT containing legume species, which, up to now, are only rarely cropped in Europe.

#### 7.2.2 Nitrous oxide

Each kg of N as ammonium nitrate produced in the industrial Haber-Bosch process emits significant amounts of greenhouse gases (ecoinvent Centre 2010), of which 19 g consists of  $N_2O$ , a greenhouse gas that is *ca.* 300 times more active than  $CO_2$  (Kingston-Smith *et al.* 2010). In addition, the IPCC (2006) suggested that for each of 100 kg of N fertilizer added to the soil, on average 1.0 kg of N is emitted as  $N_2O$ . The process of denitrification is the most important source of  $N_2O$  from pasture systems (Soussana *et al.* 2010) and large peaks of  $N_2O$  emissions are measured in grassland immediately after N-fertilizer applications (Ineson *et al.* 1998; Klumpp *et al.* 2011). There are four reasons why  $N_2O$  emissions (as other N



losses like NO<sub>3</sub> leaching) from legume-based grassland systems should be lower than from fertilized grass systems: (i) nitrogen is fixed symbiotically within the legume nodules and thus is not freely available in the soil in a reactive form, (ii) symbiotic N<sub>2</sub> fixation activity is down-regulated if the sink of N for plant growth is small, (iii) in balanced grass-legume mixtures the grass roots take up N derived from legume roots and from mineralization of soil organic matter, and (iv) pure grass swards are generally fertilized with inorganic N, and application of N fertiliser in itself leads to N<sub>2</sub>O losses. A recent review by Jensen *et al.* (2012) showed that annual N<sub>2</sub>O emissions were largest in N fertilized grass swards (19 site-years; 4.49 kg N<sub>2</sub>O-N ha<sup>-1</sup>) followed by pure legume stands (17 site-years; 0.79 and 1.99 kg N<sub>2</sub>O-N ha<sup>-1</sup> for white clover and lucerne, respectively) and mixed grass-clover swards (8 site-years; 0.54 kg N<sub>2</sub>O-N ha<sup>-1</sup>). Within the revised greenhouse gas guidelines (IPCC 2006), symbiotic N<sub>2</sub> fixation has actually been removed as a direct source of N<sub>2</sub>O because of a lack of evidence of significant emissions arising from the fixation process itself (Rochette and Janzen 2005). Emissions of N<sub>2</sub>O from legumes can occur as a result of the decomposition of residues from leguminous plants but the magnitude of such emissions remains uncertain (Baggs *et al.* 2000).

At the level of the whole livestock system, Ledgard *et al.* (2009) and Basset-Mens *et al.* (2009) used life-cycle analysis to show that greenhouse gas emissions decreased by 1.15 to 1.00 kg eq-CO<sub>2</sub> kg<sup>-1</sup> milk with mixed grass-clover pastures compared to pure grass pastures. Basset-Mens *et al.* (2005) have compared greenhouse gas emissions from dairy farm systems in Sweden, Southern Germany and New Zealand using life-cycle analysis and emission coefficients. The New Zealand system essentially relies on permanent grass-white clover pastures, which are grazed all year round with an annual N fertilizer input of 100 kg ha<sup>-1</sup> and less than 10% of the feed requirement of cows is provided by feed supplements. The authors showed that the total emission per kg milk is 30 to 80% lower for the New Zealand system than for intensive European dairy farms based on predominantly grass pastures. Schils *et al.* (2005) compared the total emissions from dairy systems in the Netherlands, which were either fertilized ryegrass or grass-clover pastures (inorganic-N inputs of 275 and 69 kg ha<sup>-1</sup> year<sup>-1</sup>, respectively): greenhouse gas emissions per kg of milk were 10% lower for grass-clover pasture-based systems than for the ryegrass monocultures.

#### 7.2.3 Carbon dioxide

Industrial production of each kg of inorganic N emits 2.25 kg of  $CO_2$  (ecoinvent Centre 2010). In this respect, legumes offer a big advantage because the entire carbon (C) needed for symbiotic  $N_2$  fixation comes directly from the atmosphere via photosynthesis and this process is 'greenhouse-gas neutral'.

A further aspect mitigating climate change is C sequestration into the soil (Soussana *et al.* 2010). Carbon can only be introduced into the soil via plant photosynthesis and the C:N ratio of soil organic matter is fairly constant in almost all soils (Kirkby *et al.* 2011). Consequently, C sequestration into soil organic matter ultimately indicates sequestration of N into soil organic matter (80 kg N t<sup>-1</sup> of C). Current evidence suggests that humus formation is particularly limited by the availability of N (Christopher and Lal 2007). Legumes and their symbiotic N<sub>2</sub> fixation thus offer the option to couple C and N cycles and deliver the N needed to sequester C into soil organic matter (Zanetti *et al.* 1997; Lüscher *et al.* 2004). Data from a large survey of soil organic matter in France (Arrouays *et al.* 2001) and models (Soussana *et al.* 2004) show that the conversion of short-term N-fertilized grass leys into grass-legume mixtures could sequester C into soil organic matter (300-500 kg C ha<sup>-1</sup> year<sup>-1</sup> over a time span of 20 years). Indeed, several studies found higher soil organic matter contents under mixed grass-legume swards than under pure grass swards (Mortenson *et al.* 2004; Fornara and Tilman 2008).



### 7.3 Forage legumes reduce consumption of nonrenewable energy

The introduction of legumes reduces non-renewable energy consumption in livestock farming since legume-based grassland systems access atmospheric N and no direct financial or energetic cost is linked to such N input. In comparison, each kg of inorganic N produced in the industrial Haber-Bosch process consumes large amounts of energy. The estimations are highly variable and range from 44 MJ (Kaltschmitt and Reinhardt 1997) to 78 MJ (Kitani *et al.* 1999). Further, it is estimated that under French conditions 0.17 MJ of energy are required to produce 1 MJ of net energy with ryegrass fertilized at 150 kg N ha<sup>-1</sup>, but only 0.06 MJ with ryegrass-white clover mixtures, and 0.13 MJ for maize silage planted after wheat (Besnard *et al.* 2006). Similarly, energy consumption decreased from 5.0 MJ kg<sup>-1</sup> milk for intensive dairy farms in the Netherlands to 4.0 MJ kg<sup>-1</sup> milk for French farms using maize silage and fertilized grasses and to 3.1 and 1.4 MJ kg<sup>-1</sup> for systems based on grazing in Ireland and New Zealand, respectively (Le Gall *et al.* 2009; Peyraud *et al.* 2009). The higher energy consumption in Irish grassland based systems appears to be linked to the utilization of higher amounts of N fertilizer on pure ryegrass pastures, in comparison with the lower use of N in New Zealand systems.

#### 8. References

- Aarssen, L. W. (1983) Ecological combining ability and competitive combining ability in plants: towards a general evolutionary theory of coexistence in systems of competition. *American Naturalist* 122:707-731.
- Almeida, J. P. F., Hartwig, U. A., Frehner, M., Nösberger, J., Lüscher, A. (2000) Evidence that P deficiency induces N feedback regulation of symbiotic N<sub>2</sub> fixation in white clover (*Trifolium repens* L.). *Journal of Experimental Botany* 51:1289-1297.
- Annicchiarico, P., Piano, E. (1997) Response of white clover genotypes to intergenotypic and interspecific interference. *Journal of Agricultural Science* 128:431-437.
- Arrouays, D., Deslais, W., Badeau, V. (2001) The carbon content of topsoil and its geographical distribution in France. *Soil Use and Management* 17:7-11.
- Aydin, I., Uzun, F. (2005) Nitrogen and phosphorus fertilization of rangelands affects yield, forage quality and the botanical composition. *European Journal of Agronomy* 23:8-14.
- Baggs, E. M., Rees, R. M., Smith, K. A., Vinten, A. J. A. (2000) Nitrous oxide emission from soils after incorporating crop residues. *Soil Use and Management* 16:82-87.
- Basset-Mens, C., Ledgard, S., Carran, A. (2005) First life cycle assessment of milk production from New Zealand dairy farm systems. AgResearch Limited, Ruakura Research Centre, New Zealand. <a href="https://www.anzsee.org/anzsee2005papers/Basset-Mens\_LCA\_NZ\_milk\_production.pdf">www.anzsee.org/anzsee2005papers/Basset-Mens\_LCA\_NZ\_milk\_production.pdf</a> (accessed 24 January 2014).
- Basset-Mens, C., Ledgard, S., Boyes, M. (2009) Eco-efficiency of intensification scenarios for milk production in New Zealand. *Ecological Economics* 68:1615-1625.



- Batterman, S. A., Wurzburger, N., Hedin, L. O. (2013) Nitrogen and phosphorus interact to control tropical symbiotic N<sub>2</sub> fixation: a test in Inga punctata. *Journal of Ecology* 101:1400-1408.
- Baumont, R., Aufrere, J., Niderkorn, V., Andueza, D., Surault, F., Peccatte, J. R., Delaby, L., Pelletier, P. (2008) Specific diversity in forages: its consequences on the feeding value. *Fourrages*:189-206.
- Besnard, A., Montarges-Lellahi, A., Hardy, A. (2006) Système de culture et nutrition azotée. Effets sur les émissions de GES et le bilan énergétique. *Fourrages* 187:311-320.
- Blumenthal, J. M., Russelle, M. P. (1996) Subsoil nitrate uptake and symbiotic dinitrogen fixation by alfalfa. *Agronomy Journal* 88:909-915.
- Boller, B. C., Nösberger, J. (1987) Symbiotically fixed nitrogen from field-grown white and red clover mixed with ryegrasses at low levels of <sup>15</sup>N-fertilization. *Plant and Soil* 104:219-226.
- Bowman, W. D., Schardt, J. C., Schmidt, S. K. (1996) Symbiotic N<sub>2</sub>-fixation in alpine tundra: ecosystem input and variation in fixation rates among communities. *Oecologia* 108:345-350.
- Brophy, L. S., Heichel, G. H. (1989) Nitrogen release from roots of alfalfa and soybean grown in sand culture. *Plant and Soil* 116:77-84.
- Brown, L., Scholefield, D., Jewkes, E. C., Preedy, N., Wadge, K., Butler, M. (2000) The effect of sulphur application on the efficiency of nitrogen use in two contrasting grassland soils. *Journal of Agricultural Science* 135:131-138.
- Carbonero, C. H., Mueller-Harvey, I., Brown, T. A., Smith, L. (2011) Sainfoin (*Onobrychis viciifolia*): a beneficial forage legume. *Plant Genetic Resources: Characterization and Utilization* 9:70-85.
- Cardinale, B. J., Wrigh, J. P., Cadotte, M. W., Carroll, I. T., Hector, A., Srivastava, D. S., Loreau, M., Weis, J. J. (2007) Impacts of plant diversity on biomass production increase through time because of species complementarity. *Proceedings of the National Academy of Sciences of the United States of America* 104:18123-18128.
- Carlsson, G., Huss-Danell, K. (2003) Nitrogen fixation in perennial forage legumes in the field. *Plant and Soil* 253:353-372.
- Castle, M. E., Reid, D., Watson, J. N. (1983) Silage and milk-production studies with diets containing withe clover silage. *Grass and Forage Science* 38:193-200.
- Christensen, J. H., Hewitson, B., Busuioc, A., Chen, A., Gao, X., Held, R., Jones, R., Kolli, R. K., Kwon, W. K., Laprise, R., Magana Rueda, V., Mearns, L., Menendez, C. G., Räisänen, J., Rinke, A., Sarr, A., Whetton, P., Arritt, R., Benestad, R., Beniston, M., Bromwich, D., Caya, D., Comiso, J., de Elia, R., Dethloff, K. (2007) Regional climate projections. pp. 847–940 *In* S. Solomon, D. Qin, M. Manning, Z. Chen, M. Marquis, K. B. Averyt, M. Tignor, H. L. Miller (eds.). *Climate Change 2007: The Physical Science Basis. Contribution of Working group I to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change.* University Press, Cambridge, UK.
- Christopher, S. F., Lal, R. (2007) Nitrogen management affects carbon sequestration in North American cropland soils. *Critical Reviews in Plant Sciences* 26:45-64.
- Collins, R. P., Connolly, J., Porqueddu, C. (2004) Effects of legume genetic diversity on the productivity of legume/grass mixtures COST Action 852. *Grassland Science in Europe* 9:486-488.
- Collins, R. P., Helgadottir, A., Frankow-Lindberg, B. E., Skot, L., Jones, C., Skot, K. P. (2012) Temporal changes in population genetic diversity and structure in red and white clover grown in three contrasting environments in northern Europe. *Annals of Botany* 110:1341-1350.
- Connolly, J., Bell, T., Bolger, T., Brophy, C., Carnus, T., Finn, J. A., Kirwan, L., Isbell, F., Levine, J., Lüscher, A., Picasso, V., Roscher, C., Teresa Sebastia, M., Suter, M., Weigelt, A. (2013) An improved model to predict the effects of changing biodiversity levels on ecosystem function. *Journal of Ecology* 101:344-355.



- Cralle, H. T., Heichel, G. H. (1982) Temperature and chilling sensitivity of nodule nitrogenase activity of unhardened alfalfa. *Crop Science* 22:300-304.
- Daepp, M., Nösberger, J., Lüscher, A. (2001) Nitrogen fertilization and developmental stage alter the response of *Lolium perenne* to elevated CO<sub>2</sub>. *New Phytologist* 150:347-358.
- de Kroon, H. (2007) How do roots interact? Science 318:1562-1563.
- de Kroon, H., Hendriks, M., van Ruijven, J., Ravenek, J., Padilla, F. M., Jongejans, E., Visser, E. J. W., Mommer, L. (2012) Root responses to nutrients and soil biota: drivers of species coexistence and ecosystem productivity. *Journal of Ecology* 100:6-15.
- Dewhurst, R. J., Delaby, L., Moloney, A., Boland, T., Lewis, E. (2009) Nutritive value of forage legumes used for grazing and silage. *Irish Journal of Agricultural and Food Research* 48:167-187.
- Durand, J.-L., Sheehy, J. E., Minchin, F. R. (1987) Nitrogenase activity, photosynthesis and nodule water potential in soybean plants experiencing water deprivation. *Journal of Experimental Botany* 38:311-321.
- ecoinvent Centre (2010) The life cycle inventory data, v2.2. Swiss Centre for Life Cycle Inventories, Dübendorf. ISBN 3-905594-38-2. <a href="www.ecoinvent.org">www.ecoinvent.org</a> (accessed 24 January 2014).
- Elgersma, A., Schlepers, H. (1997) Performance of white clover perennial ryegrass mixtures under cutting. *Grass and Forage Science* 52:134-146.
- Elgersma, A., Schlepers, H., Nassiri, M. (2000) Interactions between perennial ryegrass (*Lolium perenne* L.) and white clover (*Trifolium repens* L.) under contrasting nitrogen availability: productivity, seasonal patterns of species composition, N<sub>2</sub> fixation, N transfer and N recovery. *Plant and Soil* 221:281-299.
- Eriksen, J., Vinther, F. P., Soegaard, K. (2004) Nitrate leaching and N<sub>2</sub>-fixation in grasslands of different composition, age and management. *Journal of Agricultural Science* 142:141-151.
- Eriksen, J., Ledgard, S., Luo, J., Schils, R., Rasmussen, J. (2010) Environmental impacts of grazed pastures. *Grassland Science in Europe* 15:880-890.
- Evans, D. R., Hill, J., Williams, T. A., Rhodes, I. (1985) Effects of coexistence on the performance of white clover-perennial ryegrass mixtures. *Oecologia* 66:536-539.
- Evans, D. R., Hill, J., Williams, T. A., Rhodes, I. (1989) Coexistence and the productivity of white clover-perennial ryegrass mixtures. *Theoretical and Applied Genetics* 77:65-70.
- Finn, J. A., Kirwan, L., Connolly, J., Sebastià, M. T., Helgadottir, A., Baadshaug, O. H., Bélanger, G., Black, A., Brophy, C., Collins, R. P., Čop, J., Dalmannsdóttir, S., Delgado, I., Elgersma, A., Fothergill, M., Frankow-Lindberg, B. E., Ghesquiere, A., Golinska, B., Golinski, P., Grieu, P., Gustavsson, A.-M., Höglind, M., Huguenin-Elie, O., Jørgensen, M., Kadziuliene, Z., Kurki, P., Llurba, R., Lunnan, T., Porqueddu, C., Suter, M., Thumm, U., Lüscher, A. (2013) Ecosystem function enhanced by combining four functional types of plant species in intensively managed grassland mixtures: a 3-year continental-scale field experiment. *Journal of Applied Ecology* 50:365-375.
- Fornara, D. A., Tilman, D. (2008) Plant functional composition influences rates of soil carbon and nitrogen accumulation. *Journal of Ecology* 96:314-322.
- Frame, J. (1986) The production and quality potential of four forage legumes sown alone and combined in various associations. *Crop Research* 25:103-122.
- Frame, J. (1991) Herbage production and quality of a range of secondary grass species at 5 rates of fertilizer nitrogen application. *Grass and Forage Science* 46:139-151.
- Frankow-Lindberg, B. E. (2001) Adaptation to winter stress in nine white clover populations: Changes in non-structural carbohydrates during exposure to simulated winter conditions and 'spring' regrowth potential. *Annals of Botany* 88:745-751.



- Frankow-Lindberg, B. E., Brophy, C., Collins, R. P., Connolly, J. (2009) Biodiversity effects on yield and unsown species invasion in a temperate forage ecosystem. *Annals of Botany* 103:913-921.
- Frehner, M., Lüscher, A., Hebeisen, T., Zanetti, S., Schubiger, F., Scalet, M. (1997) Effects of elevated partial pressure of carbon dioxide and season of the year on forage quality and cyanide concentration of *Trifolium repens* L. from a FACE experiment. *Acta Oecologica* 18:297-304.
- Gastal, F., Julier, B., Surault, F., Litrico, I., Durand, J.-L., Denoue, D., Ghesquière, M., Sampoux, J.-P. (2012) Intérêt des prairies cultivées multiespèces dans le contexte des systèmes de polyculture-élevage. *Innovations Agronomiques* 22:169-183.
- Giambalvo, D., Ruisi, P., di Miceli, G., Frenda, A. S., Amato, G. (2011) Forage production, N uptake, N<sub>2</sub> fixation, and N recovery of berseem clover grown in pure stand and in mixture with annual ryegrass under different managements. *Plant and Soil* 342:379-391.
- Guckert, A., Hay, R. K. M. (2001) The overwintering, spring growth, and yield in mixed species swards, of white clover in Europe Preface. *Annals of Botany* 88:667-668.
- Hammond, K. J., Burke, J. L., Koolaard, J. P., Muetzel, S., Pinares-Patino, C. S., Waghorn, G. C. (2013) Effects of feed intake on enteric methane emissions from sheep fed fresh white clover (*Trifolium repens*) and perennial ryegrass (*Lolium perenne*) forages. *Animal Feed Science and Technology* 179:121-132.
- Häring, D. A., Suter, D., Amrhein, N., Lüscher, A. (2007) Biomass allocation is an important determinant of the tannin concentration in growing plants. *Annals of Botany* 99:111-120.
- Häring, D. A., Scharenberg, A., Heckendorn, F., Dohme, F., Lüscher, A., Maurer, V., Suter, D., Hertzberg, H. (2008) Tanniferous forage plants: Agronomic performance, palatability and efficacy against parasitic nematodes in sheep. *Renewable Agriculture and Food Systems* 23:19-29.
- Hartwig, U. A. (1998) The regulation of symbiotic  $N_2$  fixation: A conceptual model of N feedback from the ecosystem to the gene expression level. *Perspectives in Plant Ecology Evolution and Systematics* 1:92-120.
- Hebeisen, T., Lüscher, A., Zanetti, S., Fischer, B. U., Hartwig, U. A., Frehner, M., Hendrey, G. R., Blum, H., Nösberger, J. (1997) Growth response of *Trifolium repens* L. and *Lolium perenne* L. as monocultures and bi-species mixture to free air CO<sub>2</sub> enrichment and management. *Global Change Biology* 3:149-160.
- Hector, A., Schmid, B., Beierkuhnlein, C., Caldeira, M. C., Diemer, M., Dimitrakopoulos, P. G., Finn, J. A., Freitas, H., Giller, P. S., Good, J., Harris, R., Hogberg, P., Huss-Danell, K., Joshi, J., Jumpponen, A., Körner, C., Leadley, P. W., Loreau, M., Minns, A., Mulder, C. P., O Donovan, G., Otway, S. J. (1999) Plant diversity and productivity experiments in European grasslands. Science 286:1123-1127.
- Henry, G. H. R., Svoboda, J. (1986) Dinitrogen fixation (acetylene-reduction) in high arctic sedge meadow communities. *Arctic and Alpine Research* 18:181-187.
- Hill, J. (1990) The 3 C's competition, coexistence and coevolution and their impact on the breeding of forage crop mixtures. *Theoretical and Applied Genetics* 79:168-176.
- Hille Ris Lambers, J., Harpole, W. S., Tilman, D., Knops, J., Reich, P. B. (2004) Mechanisms responsible for the positive diversity-productivity relationship in Minnesota grasslands. *Ecology Letters* 7:661-668.
- Høgh-Jensen, H. (2006) The nitrogen transfer between plants: An important but difficult flux to quantify. *Plant and Soil* 282:1-5.
- Høgh-Jensen, H., Schjoerring, J. K. (1997) Interactions between white clover and ryegrass under contrasting nitrogen availability: N<sub>2</sub> fixation, N fertilizer recovery, N transfer and water use efficiency. *Plant and Soil* 197:187-199.



- Hooda, P. S., Moynagh, M., Svoboda, I. F., Anderson, H. A. (1998) A comparative study of nitrate leaching from intensively managed monoculture grass and grass-clover pastures. *Journal of Agricultural Science* 131:267-275.
- Hooper, D. U., Vitousek, P. M. (1998) Effects of plant composition and diversity on nutrient cycling. *Ecological Monographs* 68:121-149.
- Hooper, D. U., Dukes, J. S. (2004) Overyielding among plant functional groups in a long-term experiment. *Ecology Letters* 7:95-105.
- Hooper, D. U., Chapin, E. S., III, Ewel, J. J., Hector, A., Inchausti, P., Lavorel, S., Lawton, J. H., Lodge, D. M., Loreau, M., Naeem, S., Schmid, B., Setala, H., Symstad, A. J., Vandermeer, J., Wardle, D. A. (2005) Effects of biodiversity on ecosystem functioning: A consensus of current knowledge. *Ecological Monographs* 75:3-35.
- Ineson, P., Coward, P. A., Hartwig, U. A. (1998) Soil gas fluxes of N<sub>2</sub>O, CH<sub>4</sub> and CO<sub>2</sub> beneath *Lolium perenne* under elevated CO<sub>2</sub>: The Swiss free air carbon dioxide enrichment experiment. *Plant and Soil* 198:89-95.
- IPCC (2006) *Guidelines for National Greenhouse Gas Inventories*. Intergovernmental Panel of Climate Change, Tokyo, Japan. <a href="www.ipcc-nggip.iges.or.jp/public/index.html">www.ipcc-nggip.iges.or.jp/public/index.html</a> (accessed 24 January 2014).
- Jacot, K. A., Lüscher, A., Nösberger, J., Hartwig, U. A. (2000a) Symbiotic N<sub>2</sub> fixation of various legume species along an altitudinal gradient in the Swiss Alps. *Soil Biology & Biochemistry* 32:1043-1052.
- Jacot, K. A., Lüscher, A., Nösberger, J., Hartwig, U. A. (2000b) The relative contribution of symbiotic N<sub>2</sub> fixation and other nitrogen sources to grassland ecosystems along an altitudinal gradient in the Alps. *Plant and Soil* 225:201-211.
- Jayanegara, A., Leiber, F., Kreuzer, M. (2012) Meta-analysis of the relationship between dietary tannin level and methane formation in ruminants from *in vivo* and *in vitro* experiments. *Journal of Animal Physiology and Animal Nutrition* 96:365-375.
- Jensen, E. S., Peoples, M. B., Boddey, R. M., Gresshoff, P. M., Hauggaard-Nielsen, H., Alves, B. J. R., Morrison, M. J. (2012) Legumes for mitigation of climate change and the provision of feedstock for biofuels and biorefineries. A review. *Agronomy for Sustainable Development* 32:329-364.
- Kaltschmitt, M., Reinhardt, G. A. (1997) *Nachwachsende Energieträger Grundlagen, Verfahren, ökologische Bilanzierung.* Vieweg Verlag, Braunschweig/Wiesbaden, Germany.
- Kingston-Smith, A. H., Edwards, J. E., Huws, S. A., Kim, E. J., Abberton, M. (2010) Plant-based strategies towards minimising 'livestock's long shadow'. *Proceedings of the Nutrition Society* 69:613-620.
- Kirkby, C. A., Kirkegaard, J. A., Richardson, A. E., Wade, L. J., Blanchard, C., Batten, G. (2011) Stable soil organic matter: A comparison of C:N:P:S ratios in Australian and other world soils. *Geoderma* 163:197-208.
- Kirwan, L., Lüscher, A., Sebastia, M. T., Finn, J. A., Collins, R. P., Porqueddu, C., Helgadottir, A., Baadshaug, O. H., Brophy, C., Coran, C., Dalmannsdottir, S., Delgado, I., Elgersma, A., Fothergill, M., Frankow-Lindberg, B. E., Golinski, P., Grieu, P., Gustavsson, A. M., Hoglind, M., Huguenin-Elie, O., Iliadis, C., Jorgensen, M., Kadziuliene, Z., Karyotis, T., Lunnan, T., Malengier, M., Maltoni, S., Meyer, V., Nyfeler, D., Nykanen-Kurki, P., Parente, J., Smit, H. J., Thumm, U., Connolly, J. (2007) Evenness drives consistent diversity effects in intensive grassland systems across 28 European sites. *Journal of Ecology* 95:530-539.
- Kitani, O., Jungbluth, T., Peath, R. M., Ramdani, A. (1999) *CIGR handbook of agricultural engineering. Volume V: Energy & Biomass Engineering.* American Society of Agricultural Engineers, St Joseph, USA.
- Klumpp, K., Bloor, J. M. G., Ambus, P., Soussana, J.-F. (2011) Effects of clover density on N<sub>2</sub>O emissions and plant-soil N transfers in a fertilised upland pasture. *Plant and Soil* 343:97-107.



- Lambers, J. H. R., Harpole, W. S., Tilman, D., Knops, J., Reich, P. B. (2004) Mechanisms responsible for the positive diversity-productivity relationship in Minnesota grasslands. *Ecology Letters* 7:661-668.
- Le Gall, A., Beguin, E., Dollé, J. B., Manneville, V., Pflimlin, A. (2009) Nouveaux compromis techniques pour concilier les impératifs d'efficacité économique et environnementale en élevage herbivore. *Fourrages* 198:131-152.
- Ledgard, S., Schils, R., Eriksen, J., Luo, J. (2009) Environmental impacts of grazed clover/grass pastures. *Irish Journal of Agricultural and Food Research* 48:209-226.
- Ledgard, S. F., Steele, K. W. (1992) Biological nitrogen fixation in mixed legume-grass pastures. *Plant and Soil* 141:137-153.
- Ledgard, S. F., Penno, J. W., Sprosen, M. S. (1999) Nitrogen inputs and losses from clover/grass pastures grazed by dairy cows, as affected by nitrogen fertilizer application. *Journal of Agricultural Science* 132:215-225.
- Ledgard, S. F., Sprosen, M. S., Penno, J. W., Rajendram, G. S. (2001) Nitrogen fixation by white clover in pastures grazed by dairy cows: Temporal variation and effects of nitrogen fertilization. *Plant and Soil* 229:177-187.
- Loiseau, P., Carrere, P., Lafarge, M., Delpy, R., Dublanchet, J. (2001) Effect of soil-N and urine-N on nitrate leaching under pure grass, pure clover and mixed grass/clover swards. *European Journal of Agronomy* 14:113-121.
- Loreau, M., Hector, A. (2001) Partitioning selection and complementarity in biodiversity experiments. *Nature* 412:72-76.
- Loreau, M., Naeem, S., Inchausti, P., Bengtsson, J., Grime, J. P., Hector, A., Hooper, D. U., Huston, M. A., Raffaelli, D., Schmid, B., Tilman, D., Wardle, D. A. (2001) Biodiversity and ecosystem functioning: Current knowledge and future challenges. *Science* 294:804-808.
- Lory, J. A., Russelle, M. P., Heichel, G. H. (1992) Quantification of symbiotically fixed nitrogen in soil surrounding alfalfa roots and nodules. *Agronomy Journal* 84:1033-1040.
- Lüscher, A., Connolly, J., Jacquard, P. (1992) Neighbour specificity between *Lolium perenne* and *Trifolium repens* from a natural pasture. *Oecologia* 91:404-409.
- Lüscher, A., Fuhrer, J., Newton, P. C. D. (2005) Global atmospheric change and its effect on managed grassland systems. pp. 251-264 *In* D. A. McGilloway (ed.) *Grassland: A Global Resource*. Wageningen Academic Publishers, Wageningen, The Netherlands.
- Lüscher, A., Soussana, J. F., Huguenin-Elie, O. (2011) Role and impacts of legumes in grasslands for high productivity and N gain from symbiotic N<sub>2</sub> fixation. pp. 101-107 *In* G. Lemaire, H. J., C. A. (eds.). *Grassland Productivity and Ecosystem Services*. CABI, Oxfordshire, UK.
- Lüscher, A., Daepp, M., Blum, H., Hartwig, U. A., Nösberger, J. (2004) Fertile temperate grassland under elevated atmospheric CO<sub>2</sub> role of feed-back mechanisms and availability of growth resources. *European Journal of Agronomy* 21:379-398.
- Lüscher, A., Mueller-Harvey, I., Soussana, J. F., Rees, R. M., Peyraud, J. L. (2013) Potential of legume-based grassland-livestock systems in Europe. *Grassland Science in Europe* 18:3-29.
- Marquard, E., Weigelt, A., Temperton, V. M., Roscher, C., Schumacher, J., Buchmann, N., Fischer, M., Weisser, W. W., Schmid, B. (2009) Plant species richness and functional composition drive overyielding in a six-year grassland experiment. *Ecology* 90:3290-3302.
- Marshall, A. H., Williams, T. A., Abberton, M. T., Michaelson-Yeates, T. P. T., Olyott, P., Powell, H. G. (2004) Forage quality of white clover (*Trifolium repens* L.) x Caucasian clover (*T. ambiguum* M. Bieb.) hybrids and their grass companion when grown over three harvest years. *Grass and Forage Science* 59:91-99.
- McCaughey, W. P., Wittenberg, K., Corrigan, D. (1999) Impact of pasture type on methane production by lactating beef cows. *Canadian Journal of Animal Science* 79:221-226.



- McKane, R. B., Grigal, D. F., Russelle, M. P. (1990) Spatiotemporal differences in <sup>15</sup>N uptake and the organization of an old-field plant community. *Ecology* 71:1126-1132.
- Mommer, L., van Ruijven, J., de Caluwe, H., Smit-Tiekstra, A. E., Wagemaker, C. A. M., Ouborg, N. J., Bogemann, G. M., van der Weerden, G. M., Berendse, F., de Kroon, H. (2010) Unveiling below-ground species abundance in a biodiversity experiment: a test of vertical niche differentiation among grassland species. *Journal of Ecology* 98:1117-1127.
- Mortenson, M. C., Schuman, G. E., Ingram, L. J. (2004) Carbon sequestration in rangelands interseeded with yellow-flowering alfalfa (*Medicago sativa* ssp *falcata*). *Environmental Management* 33:S475-S481.
- Nassiri, M., Elgersma, A. (2002) Effects of nitrogen on leaves, dry matter allocation and regrowth dynamics in *Trifolium repens* L. and *Lolium perenne* L. in pure and mixed swards. *Plant and Soil* 246:107-121.
- Nesheim, L., Boller, B. C. (1991) Nitrogen-fixation by white clover when competing with grasses at moderately low temperatures. *Plant and Soil* 133:47-56.
- Nyfeler, D. (2009) Productivity and nitrogen utilization in productive grassland: effect of species combinations, species proportions and nitrogen fertilization. PhD thesis. ETH Zürich, Zürich.
- Nyfeler, D., Huguenin-Elie, O., Suter, M., Frossard, E., Lüscher, A. (2011) Grass-legume mixtures can yield more nitrogen than legume pure stands due to mutual stimulation of nitrogen uptake from symbiotic and non-symbiotic sources. *Agriculture Ecosystems & Environment* 140:155-163.
- Nyfeler, D., Huguenin-Elie, O., Suter, M., Frossard, E., Connolly, J., Lüscher, A. (2009) Strong mixture effects among four species in fertilized agricultural grassland led to persistent and consistent transgressive overyielding. *Journal of Applied Ecology* 46:683-691.
- Oberson, A., Frossard, E., Bühlmann, C., Mayer, J., Mäder, P., Lüscher, A. (2013) Nitrogen fixation and transfer in grass-clover leys under organic and conventional cropping systems. *Plant and Soil* 371:237-255.
- Oelmann, Y., Buchmann, N., Gleixner, G., Habekost, M., Roscher, C., Rosenkranz, S., Schulze, E.-D., Steinbeiss, S., Temperton, V. M., Weigelt, A., Weisser, W. W., Wilcke, W. (2011) Plant diversity effects on aboveground and belowground N pools in temperate grassland ecosystems: Development in the first 5 years after establishment. *Global Biogeochemical Cycles* 25:Gb2014. DOI:10.1029/2010gb003869.
- Patto, M. C. V., Skiba, B., Pang, E. C. K., Ochatt, S. J., Lambein, F., Rubiales, D. (2006) *Lathyrus* improvement for resistance against biotic and abiotic stresses: From classical breeding to marker assisted selection. *Euphytica* 147:133-147.
- Peoples, M. B., Brockwell, J., Herridge, D. F., Rochester, I. J., Alves, B. J. R., Urquiaga, S., Boddey, R. M., Dakora, F. D., Bhattarai, S., Maskey, S. L., Sampet, C., Rerkasem, B., Khan, D. F., Hauggaard-Nielsen, H., Jensen, E. S. (2009) The contributions of nitrogen-fixing crop legumes to the productivity of agricultural systems. *Symbiosis* 48:1-17.
- Peterson, P. R., Sheaffer, C. C., Jordan, R. M., Christians, C. J. (1994) Responses of kura clover to sheep grazing and clipping. 1. Yield and forage quality. *Agronomy Journal* 86:655-660.
- Peyraud, J. L., Le Gall, A., Lüscher, A. (2009) Potential food production from forage legume-based-systems in Europe: an overview. *Irish Journal of Agricultural and Food Research* 48:115-135.
- Pirhofer-Walzl, K., Rasmussen, J., Høgh-Jensen, H., Eriksen, J., Soegaard, K., Rasmussen, J. (2012) Nitrogen transfer from forage legumes to nine neighbouring plants in a multi-species grassland. *Plant and Soil* 350:71-84.
- Rasmussen, J., Soegaard, K., Pirhofer-Walzl, K., Eriksen, J. (2012) N<sub>2</sub>-fixation and residual N effect of four legume species and four companion grass species. *European Journal of Agronomy* 36:66-74.
- Rochette, P., Janzen, H. H. (2005) Towards a revised coefficient for estimating N<sub>2</sub>O emissions from legumes. *Nutrient Cycling in Agroecosystems* 73:171-179.



- Rockström, J., Steffen, W., Noone, K., Persson, A., Chapin, F. S., Lambin, E. F., Lenton, T. M., Scheffer, M., Folke, C., Schellnhuber, H. J., Nykvist, B., de Wit, C. A., Hughes, T., van der Leeuw, S., Rodhe, H., Sorlin, S., Snyder, P. K., Costanza, R., Svedin, U., Falkenmark, M., Karlberg, L., Corell, R. W., Fabry, V. J., Hansen, J., Walker, B., Liverman, D., Richardson, K., Crutzen, P., Foley, J. A. (2009) A safe operating space for humanity. *Nature* 461:472-475.
- Roscher, C., Thein, S., Schmid, B., Scherer-Lorenzen, M. (2008) Complementary nitrogen use among potentially dominant species in a biodiversity experiment varies between two years. *Journal of Ecology* 96:477-488.
- Roscher, C., Schumacher, J., Baade, J., Wilcke, W., Gleixner, G., Weisser, W.-W., Schmid, B., Schulze, E.-D. (2004) The role of biodiversity for element cycling and trophic interactions: an experimental approach in a grassland community. *Basic and Applied Ecology* 5:107-121.
- Roscher, C., Temperton, V. M., Scherer-Lorenzen, M., Schmitz, M., Schumacher, J., Schmid, B., Buchmann, N., Weisser, W. W., Schulze, E.-D. (2005) Overyielding in experimental grassland communities irrespective of species pool or spatial scale. *Ecology Letters* 8:419-429.
- Roughley, R. J., Dart, P. J. (1970) Growth of *Trifolium subterraneum* L. selected for sparse and abundant nodulation as affected by root temperature and rhizobium strain. *Journal of Experimental Botany* 21:776-786.
- Rowlinson, P., Steele, M., Nefzaoui, A. (2008) *Livestock and Global Climate Change*. Cambridge University Press, Cambridge.
- Russelle, M. P., Lamb, J. F. S., Turyk, N. B., Shaw, B. H., Pearson, B. (2007) Managing nitrogen contaminated soils: benefits of N<sub>2</sub>-fixing alfalfa. *Agronomy Journal* 99:738-746.
- Russelle, M. P., Lamb, J. F. S., Montgomery, B. R., Elsenheimer, D. W., Miller, B. S., Vance, C. P. (2001) Alfalfa rapidly remediates excess inorganic nitrogen at a fertilizer spill site. *Journal of Environmental Quality* 30:30-36.
- Sangakkara, U. R., Hartwig, U. A., Nösberger, J. (1996) Soil moisture and potassium affect the performance of symbiotic nitrogen fixation in faba bean and common bean. *Plant and Soil* 184:123-130.
- Scherer-Lorenzen, M., Palmborg, C., Prinz, A., Schulze, E. D. (2003) The role of plant diversity and composition for nitrate leaching in grasslands. *Ecology* 84:1539-1552.
- Schils, R. L. M. (1994) Nitrate losses from grazed grass and grass/clover pastures on clay soils. *Meststoffen* 1994:78-84.
- Schils, R. L. M., Verhagen, A., Aarts, H. F. M., Sebek, L. B. J. (2005) A farm level approach to define successful mitigation strategies for GHG emissions from ruminant livestock systems. *Nutrient Cycling in Agroecosystems* 71:163-175.
- Schmid, B., Hector, A., Saha, P., Loreau, M. (2008) Biodiversity effects and transgressive overyielding. *Journal of Plant Ecology* 1:95-102.
- Schwank, O., Blum, H., Nösberger, J. (1986) The influence of irradiance distribution on the growth of white clover (*Trifolium repens* L) in differently managed canopies of permanent grassland. *Annals of Botany* 57:273-281.
- Schwinning, S., Parsons, A. J. (1996a) Analysis of the coexistence mechanisms from grasses and legumes in grazing systems. *Journal of Ecology* 84:799-813.
- Schwinning, S., Parsons, A. J. (1996b) A spatially explicit population model of stoloniferous N-fixing legumes in mixed pasture with grass. *Journal of Ecology* 84:815-826.
- Serraj, R., Sinclair, T. R., Purcell, L. C. (1999) Symbiotic N<sub>2</sub> fixation response to drought. *Journal of Experimental Botany* 50:143-155.
- Simon, J. C., Vertès, F., Decau, M. L., Le Corre, L. (1997) Nitrogen flows in grazed pastures. 1-Balances on the farm scale and nitrate leaching under pastures. *Fourrages* 151:249-262.



- Sleugh, B., Moore, K. J., George, J. R., Brummer, E. C. (2000) Binary legume-grass mixtures improve forage yield, quality, and seasonal distribution. *Agronomy Journal* 92:24-29.
- Soussana, J.-F., Tallec, T. (2010) Can we understand and predict the regulation of biological N<sub>2</sub> fixation in grassland ecosystems? *Nutrient Cycling in Agroecosystems* 88:197-213.
- Soussana, J. F., Lafarge, M. (1998) Competition for resources between neighbouring species and patch scale vegetation dynamics in temperate grasslands. *Annales de Zootechnie* 47:371-382.
- Soussana, J. F., Tallec, T., Blanfort, V. (2010) Mitigating the greenhouse gas balance of ruminant production systems through carbon sequestration in grasslands. *Animal* 4:334-350.
- Soussana, J. F., Minchin, F. R., Macduff, J. H., Raistrick, N., Abberton, M. T., Michaelson, Y. T. (2002) A simple model of feedback regulation for nitrate uptake and N<sub>2</sub> fixation in contrasting phenotypes of white clover. *Annals of Botany* 90:139-147.
- Soussana, J. F., Loiseau, P., Vuichard, N., Ceschia, E., Balesdent, J., Chevallier, T., Arrouays, D. (2004) Carbon cycling and sequestration opportunities in temperate grasslands. *Soil Use and Management* 20:219-230.
- Sparrow, S. D., Cochran, V. L., Sparrow, E. B. (1995) Dinitrogen fixation by seven legume crops in Alaska. *Agronomy Journal* 87:34-41.
- Spehn, E. M., Scherer, L. M., Schmid, B., Hector, A., Caldeira, M. C., Dimitrakopoulos, P. G., Finn, J. A., Jumpponen, A., O Donnovan, G., Pereira, J. S., Schulze, E. D., Troumbis, A. Y., Körner, C. (2002) The role of legumes as a component of biodiversity in a cross-European study of grassland biomass nitrogen. *Oikos* 98:205-218.
- Squillace, P. J., Scott, J. C., Moran, M. J., Nolan, B. T., Kolpin, D. W. (2002) VOCs, pesticides, nitrate, and their mixtures in groundwater used for drinking water in the United States. *Environmental Science & Technology* 36:1923-1930.
- Sturludóttir, E., Brophy, C., Bélanger, G., Gustavsson, A.-M., Jørgensen, M., Lunnan, T., Helgadóttir, A. (2013) Benefits of mixing grasses and legumes for herbage yield and nutritive value in Northern Europe and Canada. *Grass and Forage Science*. DOI:10.1111/gfs.12037.
- Surault, F., Julier, B., Barre, P. (2010) Agricultural values of perennial ryegrass pastures constituted by one or several cultivars. *Fourrages* 204:255-262.
- Suter, D., Briner, H. U., Lüscher, A. (2007) Effect of *Lolium perenne* L. variety on the botanical composition of grass-clover mixtures. *Grassland Science in Europe* 12:417-420.
- Suter, D., Huguenin-Elie, O., Nyfeler, D., Lüscher, A. (2010) Agronomically improved grass-legume mixtures: higher dry matter yields and more persistent legume proportions. *Grassland Science in Europe* 15:761-763.
- Suter, D., Rosenberg, E., Mosimann, E., Frick, R. (2012) Standardmischungen für den Futterbau Revision 2013-2016. *Agrarforschung Schweiz* 3:1-12.
- Suter, M., Connolly, J., Finn, J. A., Helgadóttir, Á., Golinski, P., Kirwan, L., Loges, R., Kadžiuliene, Ž., Sebastià, M. T., Taube, F., Lüscher, A. (2013) Grass-legume mixtures enhance yield of total nitrogen and uptake from symbiotic N₂ fixation: Evidence from a three-year multisite experiment. *Grassland Science in Europe* 18:76-78.
- Tallec, T., Diquelou, S., Avice, J.-C., Lesuffleur, F., Lemauviel-Lavenant, S., Cliquet, J.-B., Ourry, A. (2009) Availability of N and S affect nutrient acquisition efficiencies differently by *Trifolium repens* and *Lolium perenne* when grown in monoculture or in mixture. *Environmental and Experimental Botany* 66:309-316.
- Tamminga, S., Bannink, A., Dijkstra, J., Zom, R. (2007) Feeding strategies to reduce methane loss in cattle. Animal Science Group, Wageningen University, Wageningen. Report 34. 44p. Available at: <a href="https://www.asg.wur.nl/NR/rdonlyres/F81D8745-6596-4296-A292-8553950E2B98/42707/34.pdf">www.asg.wur.nl/NR/rdonlyres/F81D8745-6596-4296-A292-8553950E2B98/42707/34.pdf</a> (accessed 24 January 2014).
- Thomas, R. J. (1992) The role of the legume in the nitrogen cycle of productive and sustainable pastures. *Grass and Forage Science* 47:133-142.



- Thompson, K., Askew, A. P., Grime, J. P., Dunnett, N. P., Willis, A. J. (2005) Biodiversity, ecosystem function and plant traits in mature and immature plant communities. *Functional Ecology* 19:355-358.
- Tilman, D. (1999) The ecological consequences of changes in biodiversity: A search for general principles. *Ecology* 80:1455-1474.
- Trenbath, B. R. (1974) Biomass productivity of mixtures. pp. 177–210 *In* N. C. Brady (ed.) *Advances in Agronomy*. 26. Academic Press, New York.
- Turkington, R. (1989a) The growth, distribution and neighbor relationships of *Trifolium repens* in a permanent pasture. 5. The coevolution of competitors. *Journal of Ecology* 77:717-733.
- van Dorland, H. A., Wettstein, H. R., Leuenberger, H., Kreuzer, M. (2007) Effect of supplementation of fresh and ensiled clovers to ryegrass on nitrogen loss and methane emission of dairy cows. *Livestock Science* 111:57-69.
- van Ruijven, J., Berendse, F. (2003) Positive effects of plant species diversity on productivity in the absence of legumes. *Ecology Letters* 6:170-175.
- Vertès, F., Simon, J. C., Le Corre, L., Decau, M. L. (1997) Nitrogen flows in grazed pastures. II. Study of flows and their effects on leaching. *Fourrages* 151:263-280.
- Vinther, F. P., Jensen, E. S. (2000) Estimating legume N<sub>2</sub> fixation in grass-clover mixtures of a grazed organic cropping system using two <sup>15</sup>N methods. *Agriculture Ecosystems* & *Environment* 78:139-147.
- Wachendorf, M., Collins, R. P., Elgersma, A., Fothergill, M., Frankow-Lindberg, B. E., Ghesquiere, A., Guckert, A., Guinchard, M. P., Helgadottir, A., Lüscher, A., Nolan, T., Nykänen-Kurki, P., Nösberger, J., Parente, G., Puzio, S., Rhodes, I., Robin, C., Ryan, A., Stäheli, B., Stoffel, S., Taube, F., Connolly, J. (2001) Overwintering and growing season dynamics of *Trifolium repens* L. in mixture with *Lolium perenne* L.: A model approach to plant-environment interactions. *Annals of Botany* 88:683-702.
- Waghorn, G. (2008) Beneficial and detrimental effects of dietary condensed tannins for sustainable sheep and goat production Progress and challenges. *Animal Feed Science and Technology* 147:116-139.
- Waghorn, G. C., Hegarty, R. S. (2011) Lowering ruminant methane emissions through improved feed conversion efficiency. *Animal Feed Science and Technology* 166-67:291-301.
- Wen, Y. Y., Jiang, H. F. (2005) Cutting effects on growth characteristics, yield composition, and population relationships of perennial ryegrass and white clover in mixed pasture. *New Zealand Journal of Agricultural Research* 48:349-358.
- Williams, T. A., Abberton, M. T., Rhodes, I. (2003) Performance of white clover varieties combined in blends and alone when grown with perennial ryegrass under sheep and cattle grazing. *Grass and Forage Science* 58:90-93.
- Woledge, J. (1988) Competition between grass and clover in spring as affected by nitrogen fertilizer. *Annals of Applied Biology* 112:175-186.
- Woodward, S. L., Waghorn, G. C., Laboyrie, P. G. (2004) Condensed tannins in birdsfoot trefoil (*Lotus corniculatus*) reduce methane emissions from dairy cows. *Proceedings of New Zealand Society of Animal Production* 64:160-164.
- Woodward, S. L., Waghorn, G. C., Lassey, K. R., G., L. P. (2002) Does feeding sulla (*Hedysarum coronarium*) reduce methane emissions from dairy cows? *Proceedings of New Zealand Society of Animal Production* 62:227-230.
- Wu, L., MeGechan, M. B. (1999) Simulation of nitrogen uptake, fixation and leaching in a grass white clover mixture. *Grass and Forage Science* 54:30-41.
- Zanetti, S., Hartwig, U. A., van Kessel, C., Lüscher, A., Hebeisen, T., Frehner, M., Fischer, B. U., Hendrey, G. R., Blum, H., Nösberger, J. (1997) Does nitrogen nutrition restrict the CO<sub>2</sub> response of fertile grassland lacking legumes? *Oecologia* 112:17-25.



### **Annex 1: Supporting information**

**Table A1.** Information on sites, and basic management and weather information. Sites are distinguished by their identifier number in the database from which these data are derived, and are arranged in order of declining total yield of the best-performing monoculture over the three years (as in Fig. 2).

This table is provided (with permission) from Finn et al. (2013). Table number changed and header shortened.

Site # in database	Country	Site	Latitude	Longitude	Altitude (m a.s.l.)	Nitrogen fertilizer (kg ha <sup>-1</sup> per annum)	Harvests per annum	Annual rainfall (mm)	Annual mean temp (°C)	No. of years
							(year 1,2,3)			
1	Belgium	Merelbeke	50°59'N	3°49E	11	150	4,3,4	780	9.9	3
40	Slovenia	Ljubljana	46°03'N	14°28'E	300	120	4	1147	10.7	2
15	Ireland	Wexford	52°16'N	6°30'W	54	150	5	1033	10.1	3
21	Netherlands	Wageningen	51°58'N	5°40'E	7	0,108,108	5	760	9.6	3
10	Germany	Renningen	48°46'N	9°11'E	460	150	4,5,5	693	8.2	3
11	Germany	St. Johann	48°28'N	9°18E	700	150	4	1046	7.4	2
9	France	Auzeville Tolosane	43°05'N	1°43'E	162	120	3,2,3	680	13.0	3
34	Switzerland	Zurich- Reckenholz**	47°26'N	8°32'E	491	150	5	1031	9.4	3
24	Norway	Ås	59°40'N	10°51'E	95	135	3	785	5.3	3
35	Wales	Aberystwyth	52°26'N	4°01'W	30	90	4	1038	9.7	3
43	Ireland	Athenry	53°17'N	8°44'W	40	75	7	885	10.4	2
36	Wales	Bronydd Mawr	51°57'N	3°37'W	323	93	4,3,4	1500	8.2	3
27	Poland	Brody	52°26'N	16°18'E	91.4	90	4,3	587	8.0	2
44	Ireland	Moorepark	52°08'N	8°16'W	48	100	7	1207	9.5	2
26	Poland	Brody	52°26'N	16°18'E	94.2	120	4,4,3	587	8.0	3
22	Norway	Saerheim	58°46'N	5°39'E	90	0	3	1180	7.1	3
33	Sweden	Öjebyn (Piteå)	65°19'N	21°24′E	5	60	2,3,2	539	2.1	3
23	Norway	Tromsø	69°40'N	18°56'E	15	60	2	1031	3.1	3



19	Lithuania (b)	Dotnuva	55°24'N	23°50'E	71	120	3,2,2	650	6.1	3
7	Finland	Mikkeli***	61°40'N	27°13′E	107	60	3	643	3.1	2
25	Norway	Løken	61°07'N	9°04′E	435	80	2	576	1.6	3
18	Lithuania (a)	Dotnuva	55°24'N	23°50'E	71	120	3	650	6.1	3
30	Spain	Gosol	42°13'N	1°39'E	1410	0	2	948	7.9	1
16	Italy	Ottava	40°44'N	8°32′E	80	31, 57, 61	4,5,5	547	16.2	3
20	Lithuania (c)	Dotnuva	55°24'N	23°50'E	71	120	3,3,2	650	6.1	3
52	Canada	Lévis	46°46'N	71°12′W	43	60	2	1175	5.26	3
31	Sweden (a)	Svalöv	55°55'N	13°07'E	55	0	3	700	7.7	3
32	Sweden (b)	Svalöv	55°55'N	13°07'E	55	0	3	700	7.7	3
13	Iceland (a)	Korpa	64°09'N	21°45'W	35	40	2	900	4.5	3
14	Iceland (b)	Korpa	64°09'N	21°45'W	35	80	2	900	4.5	3
28	Spain	Zaragoza**	41°44'N	2°53′E	225	61	2,3,1	409	14.3	3

<sup>\*</sup>Species codes as follows. Non-fixing grasses:  $Dg = Dactylis \ glomerata$ ,  $Fa = Festuca \ arundinaceum$ ,  $Lp = Lolium \ perenne$ ,  $Lr = Lolium \ rigidum$ ,  $Pp = Phleum \ pratense$ ,  $Poa = Poa \ pratensis$ .  $N_2$ -fixing legumes:  $Mp = Medicago \ polymorpha$ ,  $Ms = Medicago \ sativa$ ,  $Ta = Trifolium \ armbiguum$ ,  $Tp = Trifolium \ pratense$ ,  $Tr = Trifolium \ repens$ .



<sup>\*\*</sup>The plots at Switzerland were weeded for the first two harvests in the establishment year and the plots at Spain-Zaragoza were weeded only in year 1 of harvesting.

<sup>\*\*\*</sup>Barley was used on all plots as a nurse crop at establishment and harvested in the establishment year, as per conventional practice at this site.